

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00009-3381-02	J3490			01/01/2002	11/21/2018	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE (PREMIX) 300 MG/50 ML	50	ML	PC	IV	ML	1	EA	1	01/01/2002	11/21/2018						
50242-0041-63	J2997			01/18/2007	12/20/2018	INJECTION, ALTEPLASE RECOMBINANT, 1 MG	CATHFLO ACTIVASE (INNER) 2 MG	1	EA	VL	IV	EA	1	MG	2	01/18/2007	12/20/2018						
17478-0114-30	J3260			12/23/2015	12/17/2018	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (MDV;USP,LATEX-FREE) 40 MG/1 ML	30	ML	VL	IJ	ML	80	MG	0.5	12/23/2015	12/17/2018						
17478-0114-02	J3260			12/23/2015	12/17/2018	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (MDV;USP,LATEX-FREE) 40 MG/1 ML	2	ML	VL	IJ	ML	80	MG	0.5	12/23/2015	12/17/2018						
00409-3815-12	J2270			06/28/2005	12/31/2014	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (5X10ML,LATEX-FREE) 1 MG/ML	10	ML	VL	IJ	ML	10	MG	0.1	06/28/2005	12/31/2014						
51552-0106-09	J2001			01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	09/16/2015	99/99/9999	01/01/2004	11/06/2013	100			
00591-2224-55	J7502			12/23/2008	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (1X50ML,MODIFIED) 100 MG/ML	50	ML	VL	PO	ML	100	MG	1	10/28/2015	99/99/9999	12/23/2008	04/07/2014	1			
64679-0962-01	Q0144			02/11/2008	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM COATED) 600 MG	30	EA	BO	PO	EA	1	GM	0.6	09/11/2015	99/99/9999	02/11/2008	05/31/2014	0.6			
64679-0961-01	Q0144			02/11/2008	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	30	EA	BO	PO	EA	1	GM	0.25	08/10/2015	99/99/9999	02/11/2008	05/31/2014	0.25			
00904-6425-61	J7507			01/09/2015	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	1	EA	BO	PO	EA	1	MG	1	08/08/2016	99/99/9999	01/09/2015	01/10/2015	1			
64679-0961-05	Q0144			02/11/2008	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (3X3,FILM COATED) 250 MG	18	EA	DP	PO	EA	1	GM	0.25	08/10/2015	99/99/9999	02/11/2008	05/31/2014	0.25			
64679-0964-05	Q0144			02/11/2008	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (3X3,FILM COATED) 500 MG	9	EA	DP	PO	EA	1	GM	0.5	08/10/2015	99/99/9999	02/11/2008	05/31/2014	0.5			
64679-0964-01	Q0144			02/11/2008	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM COATED) 500 MG	30	EA	BO	PO	EA	1	GM	0.5	08/10/2015	99/99/9999	02/11/2008	05/31/2014	0.5			
64679-0961-04	Q0144			02/14/2008	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	6	EA	BO	PO	EA	1	GM	0.25	08/10/2015	99/99/9999	02/14/2008	05/31/2014	0.25			
68084-0229-01	J7500			03/14/2008	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE 50 MG	100	EA	BO	PO	EA	50	MG	1	08/26/2014	99/99/9999	03/14/2008	05/06/2014	1			
51079-0818-20	J7507			11/01/2010	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (10X10,HARD GELATIN) 1 MG	100	EA	BO	PO	EA	1	MG	1	08/06/2013	99/99/9999	11/01/2010	07/13/2012	1			
49452-1776-02	J1955			06/01/2015	99/99/9999	INJECTION, LEVOCARNITINE, PER 1 GM	L-CARNITINE HYDROCHLORIDE	100	GM	BO	NA	GM	1	GM	1	09/01/2018	99/99/9999	06/01/2015	10/17/2016	1			
49452-1776-01	J1955			06/01/2015	99/99/9999	INJECTION, LEVOCARNITINE, PER 1 GM	L-CARNITINE HYDROCHLORIDE	25	GM	BO	NA	GM	1	GM	1	09/01/2018	99/99/9999	06/01/2015	10/17/2016	1			
00409-4778-86	J0744			08/29/2006	99/99/9999	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	CIPROFLOXACIN (SINGLE-DOSE,USP) 10 MG/ML	40	ML	VL	IV	ML	200	MG	0.05	01/01/2017	99/99/9999	08/29/2006	11/01/2015	0.05			
00409-2689-01	J0295			10/09/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (SDV,ADVANTAGE) 1 GM-0.5 GM	1	EA	VL	IV	EA	1.5	GM	1	07/31/2017	99/99/9999	10/09/2006	10/01/2013	1			
00409-2987-03	J0295			10/09/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (SDV,ADVANTAGE) 2 GM-1 GM	1	EA	VL	IV	EA	1.5	GM	2	01/01/2018	99/99/9999	10/09/2006	10/01/2013	2			
49452-2697-03	J0600			06/01/2015	99/99/9999	INJECTION, EDETATE CALCIUM DISODIUM, UP TO 1000 MG	EDETATE CALCIUM DISODIUM (U.S.P.)	2500	GM	BO	NA	GM	1000	MG	1	04/01/2018	99/99/9999	06/01/2015	10/17/2016	1			
49452-2697-02	J0600			09/01/2015	99/99/9999	INJECTION, EDETATE CALCIUM DISODIUM, UP TO 1000 MG	EDETATE CALCIUM DISODIUM (U.S.P.)	500	GM	BO	NA	GM	1000	MG	1	04/01/2018	99/99/9999	09/01/2015	10/17/2016	1			
49452-2697-01	J0600			09/01/2015	99/99/9999	INJECTION, EDETATE CALCIUM DISODIUM, UP TO 1000 MG	EDETATE CALCIUM DISODIUM (U.S.P.)	125	GM	BO	NA	GM	1000	MG	1	04/01/2018	99/99/9999	09/01/2015	10/17/2016	1			
55289-0373-01	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	100	EA	BO	PO	EA	1	MG	5	11/22/2016	99/99/9999	01/01/2016	02/03/2016	5			
00310-0482-30	J8565			01/01/2005	99/99/9999	GEFITINIB, ORAL, 250 MG	IRESSA 250 MG	30	EA	BO	PO	EA	250	MG	1	07/14/2015	99/99/9999	01/01/2005	01/01/2012	1			
00517-0020-10	J0706			09/10/2007	99/99/9999	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATE (USP,10X3ML,SINGLE-DOSE) 20 MG/ML	3	ML	VL	IV	ML	5	MG	4	08/19/2015	99/99/9999	09/10/2007	03/31/2014	4			
68382-0860-02	J0515			06/01/2015	99/99/9999	INJECTION, BENZTROPINE MESYLATE, PER 1 MG	BENZTROPINE MESYLATE 1 MG/ML	2	ML	VL	IJ	ML	1	MG	1	05/18/2018	99/99/9999	06/01/2015	03/31/2017	1			
68382-0860-10	J0515			06/01/2015	99/99/9999	INJECTION, BENZTROPINE MESYLATE, PER 1 MG	BENZTROPINE MESYLATE 1 MG/ML	2	ML	VL	IJ	ML	1	MG	1	05/18/2018	99/99/9999	06/01/2015	03/31/2017	1			
47426-0201-01	J0185			01/01/2019	99/99/9999	INJECTION, APREPITANT, 1 MG	CINVANTI 130 MG/18 ML	18	ML	VL	IV	ML	1	MG	7.22222	01/01/2019	99/99/9999						
69794-0102-01	J0584			01/01/2019	99/99/9999	INJECTION, BUROSUMAB-TWZA 1 MG	CRYSVITA (PF) 10 MG/1 ML	1	ML	VL	SC	ML	1	MG	10	01/01/2019	99/99/9999						
69794-0203-01	J0584			01/01/2019	99/99/9999	INJECTION, BUROSUMAB-TWZA 1 MG	CRYSVITA (PF) 20 MG/1 ML	1	ML	VL	SC	ML	1	MG	20	01/01/2019	99/99/9999						
69794-0304-01	J0584			01/01/2019	99/99/9999	INJECTION, BUROSUMAB-TWZA 1 MG	CRYSVITA (PF) 30 MG/1 ML	1	ML	VL	SC	ML	1	MG	30	01/01/2019	99/99/9999						
66621-0790-02	J3490			10/30/2018	12/31/2018	UNCLASSIFIED DRUGS	ANAVIP (LYOPHILIZED) (10ML VL) 24 MG/1 ML	1	EA	VL	IV	EA	1	MG	1	10/30/2018	12/31/2018						
00527-1450-06	Q0167			10/30/2018	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (SOFT GEL) 2.5 MG	60	EA	BO	PO	EA	2.5	MG	1	10/30/2018	99/99/9999						
00527-1451-06	Q0167			10/30/2018	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (SOFT GEL) 5 MG	60	EA	BO	PO	EA	2.5	MG	2	10/30/2018	99/99/9999						
00527-1452-06	Q0167			10/30/2018	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (SOFT GEL) 10 MG	60	EA	BO	PO	EA	2.5	MG	4	10/30/2018	99/99/9999						
00781-3421-94	J0637			11/12/2018	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LYOPHILIZED) 50 MG	1	EA	VL	IV	EA	5	MG	10	11/12/2018	99/99/9999						
00781-3423-94	J0637			11/12/2018	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LYOPHILIZED) 70 MG	1	EA	VL	IV	EA	5	MG	14	11/12/2018	99/99/9999						
10885-0003-01	J2062			01/01/2019	99/99/9999	LOXAPINE FOR INHALATION, 1 MG	ADASUVE (INNER PACK) 10 MG	1	EA	PG	IH	EA	1	MG	10	01/01/2019	99/99/9999						
10885-0003-05	J2062			01/01/2019	99/99/9999	LOXAPINE FOR INHALATION, 1 MG	ADASUVE 10 MG	5	EA	PG	IH	EA	1	MG	10	01/01/2019	99/99/9999						
16729-0275-67	J0583			11/01/2018	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (LYOPHILIZED) 250 MG	10	EA	VL	IV	EA	1	MG	250	11/01/2018	99/99/9999						
47335-0177-95	J3245			01/01/2019	99/99/9999	INJECTION, TILDRAKIZUMAB, 1 MG	ILUMYA (PF) 100 MG/1 ML	1	ML	SR	SC	ML	1	MG	100	01/01/2019	99/99/9999						
47426-0201-01	J3490			11/29/2017	12/31/2018	UNCLASSIFIED DRUGS	CINVANTI 130 MG/18 ML	18	ML	VL	IV	ML	1	MG	1	11/29/2017	12/31/2018						
50242-0214-01	J2357			12/03/2018	99/99/9999	INJECTION, OMALIZUMAB, 5 MG	XOLAIR (PF) 75 MG/0.5 ML	0.5	ML	SR	SC	ML	5	MG	30	12/03/2018	99/99/9999						
50242-0215-01	J2357			12/03/2018	99/99/9999	INJECTION, OMALIZUMAB, 5 MG	XOLAIR (PF) 75 MG/0.5 ML	1	ML	SR	SC	ML	5	MG	30	12/03/2018	99/99/9999						
50419-0385-01	J9057			01/01/2019	99/99/9999	INJECTION, COPANLISIB, 1 MG	ALIQOPA (LYOPHILIZED) 60 MG	1	EA	VL	IV	EA	1	MG	60	01/01/2019	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
59353-0002-01		Q5106		01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 2000 U/1 ML	1	ML	VL	IJ	ML	1000	U	2	01/01/2019	99/99/9999						
59353-0002-10		Q5106		01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 2000 U/1 ML	1	ML	VL	IJ	ML	1000	U	2	01/01/2019	99/99/9999						
59353-0003-01		Q5106		01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 3000 U/1 ML	1	ML	VL	IJ	ML	1000	U	3	01/01/2019	99/99/9999						
69639-0102-01	J3490			05/08/2018	12/31/2018	UNCLASSIFIED DRUGS	AKYNZEO (SDV,PF,LYOPHILIZED) 235 MG-	1	EA	VL	IV	EA	1	MG	1	05/08/2018	12/31/2018						
62064-0122-02	J3490			03/06/2018	12/31/2018	UNCLASSIFIED DRUGS	TROGARZO (PF) 150 MG/1 ML	1.33	ML	VL	IV	ML	1	MG	1	03/06/2018	12/31/2018						
10885-0003-01	J3490			11/20/2017	12/31/2018	UNCLASSIFIED DRUGS	ADASUVE (INNER PACK) 10 MG	1	EA	PG	IH	EA	1	MG	1	11/20/2017	12/31/2018						
10885-0003-05	J3490			11/20/2017	12/31/2018	UNCLASSIFIED DRUGS	ADASUVE 10 MG	5	EA	PG	IH	EA	1	MG	1	11/20/2017	12/31/2018						
69656-0102-10	J3490			11/15/2017	12/31/2018	UNCLASSIFIED DRUGS	VARUBI (SDV) 1.8 MG/1 ML	92.5	ML	VL	IV	ML	1	MG	1	11/15/2017	12/31/2018						
70801-0003-01	J3304			01/01/2019	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, PRESERVATIVE-FREE, EXTENDED-RELEASE, MICROSPHERE FORMULATION, 1 MG	ZILRETTA (W/DILUENT) 32 MG	1	EA	VL	IJ	EA	1	MG	32	01/01/2019	99/99/9999						
00002-7140-01	J0130			01/01/2002	12/31/2016	INJECTION ABCIXIMAB, 10 MG	REOPRO (VIAL) 2 MG/ML	5	ML	VL	IV	ML	10	MG	0.2	01/01/2002	12/31/2016						
00002-7501-01	J2941			03/01/2006	99/99/9999	INJECTION, SOMATROPIN, 1 MG	HUMATROPE (WITH STERILE DILUENT) 5 MG	1	EA	VL	SC	EA	1	MG	5	03/01/2006	99/99/9999						
00002-7501-01	J9201			01/01/2002	12/31/2018	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMZAR (VIAL) 200 MG	1	EA	VL	IV	EA	200	MG	1	01/01/2002	12/31/2018						
00002-7502-01	J9201			01/01/2002	12/31/2018	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMZAR (VIAL) 1 MG	1	EA	VL	IV	EA	200	MG	5	01/01/2002	12/31/2018						
00002-7510-01	J1817			01/01/2003	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	HUMALOG (VIAL) 100 U/ML	10	ML	VL	SC	ML	50	U	2	01/01/2003	99/99/9999						
00002-7511-01	J1815			01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG MIX 75/25 (VIAL) 75 U/ML-25 U/ML	10	ML	VL	SC	ML	5	U	20	01/01/2003	99/99/9999						
00002-7512-01	J1815			11/01/2006	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG MIX 50/50 50 U/ML-50 U/ML	10	ML	VL	SC	ML	5	U	2	11/01/2006	99/99/9999						
00002-7516-59	J1815			01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG (CARTRIDGE) 100 U/ML	3	ML	CT	SC	ML	5	U	20	01/01/2003	99/99/9999						
00002-7623-01	J9305			01/01/2005	99/99/9999	INJECTION, PEMETREXED, 10 MG	ALIMTA 500 MG	1	EA	VL	IV	EA	10	MG	50	01/01/2005	99/99/9999						
00002-7640-01	J9305			01/07/2008	99/99/9999	INJECTION, PEMETREXED, 10 MG	ALIMTA (SINGLE-USE) 100 MG	1	EA	VL	IV	EA	10	MG	10	01/07/2008	99/99/9999						
00002-8031-01	J1610			01/01/2002	99/99/9999	INJECTION, GLUCAGON HYDROCHLORIDE, PER 1 MG	GLUCAGON EMERGENCY KIT (HYPORET DISPOSABLE SRN) 1 MG	1	EA	BX	IJ	EA	1	MG	1	01/01/2002	99/99/9999						
00002-8147-01	J2941			08/30/2005	99/99/9999	INJECTION, SOMATROPIN, 1 MG	HUMATROPE (CARTRIDGE W/DILUENT) 6 MG	1	EA	CT	IJ	EA	1	MG	6	08/30/2005	99/99/9999						
00002-8148-01	J2941			08/30/2005	99/99/9999	INJECTION, SOMATROPIN, 1 MG	HUMATROPE (CARTRIDGE W/DILUENT) 12 MG	1	EA	CT	IJ	EA	1	MG	12	08/30/2005	99/99/9999						
00002-8149-01	J2941			08/30/2005	99/99/9999	INJECTION, SOMATROPIN, 1 MG	HUMATROPE (CARTRIDGE W/DILUENT) 24 MG	1	EA	CT	IJ	EA	1	MG	24	08/30/2005	99/99/9999						
00002-8215-01	J1815			01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN R (VIAL) 100 U/ML	10	ML	VL	IJ	ML	5	U	20	01/01/2003	99/99/9999						
00002-8315-01	J1815			01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN N (VIAL) 100 U/ML	10	ML	VL	SC	ML	5	U	20	01/01/2003	99/99/9999						
00002-8501-01	J1817			01/01/2003	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	HUMULIN R U-500 (VIAL, CONCENTRATED) 500 U/ML	20	ML	VL	IJ	ML	50	U	10	01/01/2003	99/99/9999						
00002-8715-01	J1815			01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN 70/30 (VIAL) 70 U/ML-30 U/ML	10	ML	VL	SC	ML	5	U	20	01/01/2003	99/99/9999						
00002-8730-59	J1815			01/01/2003	04/09/2014	INJECTION, INSULIN, PER 5 UNITS	HUMULIN N PEN (PREFILLED DISPOSABLE) 100 U/ML	3	ML	CT	SC	ML	5	U	20	01/01/2003	04/09/2014						
00002-8770-59	J1815			01/01/2003	03/18/2014	INJECTION, INSULIN, PER 5 UNITS	HUMULIN 70/30 PEN (PREFILLED DISPOSABLE) 70 U/ML-30 U/ML	3	ML	CT	SC	ML	5	U	20	01/01/2003	03/18/2014						
00002-8797-59	J1815			12/10/2007	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG MIX75/25 (KWIKPEN,5X3ML) 75 U/ML-25 U/ML	3	ML	SR	SC	ML	5	U	20	12/10/2007	99/99/9999						
00002-8798-59	J1815			12/10/2007	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG MIX 50/50 (KWIKPEN,5X3ML) 50 U/ML-50 U/ML	3	ML	SR	SC	ML	5	U	20	12/10/2007	99/99/9999						
00002-8799-59	J1815			12/10/2007	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG (KWIKPEN,5X3ML) 100 U/ML	3	ML	SR	SC	ML	5	U	20	12/10/2007	99/99/9999						
00003-0494-20	J3301			01/01/2002	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG-10 (VIAL) 10 MG/ML	5	ML	VL	IJ	ML	10	MG	1	01/01/2002	99/99/9999						
00003-0830-50	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDREA 500 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
00003-6335-17	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	DROXIA 200 MG	60	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
00003-6336-17	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	DROXIA 300 MG	60	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
00003-6337-17	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	DROXIA 400 MG	60	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
00004-0038-22	J3490			01/01/2002	99/99/9999	IMMUNOSUPPRESSIVE DRUG, NOT OTHERWISE CLASSIFIED	VALCYTE 450 MG	60	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
00004-0188-09	J1740			01/01/2007	09/30/2012	INJECTION, IBANDRONATE SODIUM, 1 MG	BONIVA 1 MG/ML	3	ML	BX	IV	EA	1	MG	1	01/01/2007	09/30/2012						
00004-0259-01	J7517			01/01/2002	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT 250 MG	100	EA	BO	PO	EA	250	MG	1	01/01/2002	99/99/9999						
00004-0259-05	J7517			01/01/2002	06/30/2015	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT 250 MG	1440	EA	BO	PO	EA	250	MG	1	01/01/2002	06/30/2015						
00004-0259-43	J7517			01/01/2002	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT 250 MG	500	EA	BO	PO	EA	250	MG	1	01/01/2002	99/99/9999						
00004-0260-01	J7517			01/01/2002	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT (CAPLET) 500 MG	100	EA	BO	PO	EA	250	MG	2	01/01/2002	99/99/9999						
00004-0260-43	J7517			01/01/2002	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT (CAPLET) 500 MG	500	EA	BO	PO	EA	250	MG	2	01/01/2002	99/99/9999						
00004-0261-29	J7517			01/01/2002	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT (FRUIT) 200 MG/ML	160	ML	BO	PO	ML	250	MG	0.8	01/01/2002	99/99/9999						
00004-0350-09	J3490			10/16/2002	99/99/9999	UNCLASSIFIED DRUGS	PEGASYS (S.D.V.) 180 MCG/ML	1	ML	VL	MR	EA	1	EA	1	10/16/2002	99/99/9999						
00004-0352-39	J3490			01/19/2004	07/31/2014	UNCLASSIFIED DRUGS	PEGASYS (MONTHLY CONVENIENCE PK) 180 MCG/0.5 ML	2	ML	BX	MR	EA	1	EA	1	01/19/2004	07/31/2014						
00004-0380-39	J1324			01/01/2007	06/30/2013	INJECTION, ENFUVRTIDE, 1 MG	FUZEON (PF) 90 MG	1	EA	PG	SC	EA	1	MG	90	01/01/2007	06/30/2013						
00004-1100-20	None			10/01/2003	99/99/9999	CAPECITABINE, 150 MG, ORAL	XELODA 150 MG	60	EA	BO	PO	EA	150	MG	1	10/01/2003	99/99/9999						
00004-1101-50	None			10/01/2003	99/99/9999	CAPECITABINE, 500 MG, ORAL	XELODA 500 MG	120	EA	BO	PO	EA	500	MG	1	10/01/2003	99/99/9999						
00004-1963-01	J0696	</																					

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00006-0461-02		J8501		01/29/2008	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND (BI-PACK) 80 MG	2 EA	DP	PO	EA		5 MG		16	01/29/2008	99/99/9999						
00006-0461-06		J8501		07/01/2006	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND 80 MG	6 EA	BX	PO	EA		5 MG		16	07/01/2006	99/99/9999						
00006-0462-06		J8501		07/01/2006	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND 125 MG	6 EA	BX	PO	EA		5 MG		25	07/01/2006	99/99/9999						
00006-0464-05		J8501		07/24/2006	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND 40 MG	5 EA	BX	PO	EA		5 MG		8	07/24/2006	99/99/9999						
00006-0464-10		J8501		07/24/2006	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND 40 MG	1 EA	BX	PO	EA		5 MG		8	07/24/2006	99/99/9999						
00006-3514-58		J0743		01/01/2002	05/01/2017	INJECTION, CILASTATIN SODIUM; IMPENEM, PER 250 MG	PRIMAXIN IV (VIAL) 250 MG-250 MG	1 EA	VL	IV	EA		250 MG		1	01/01/2002	05/01/2017						
00006-3516-59		J0743		01/01/2002	99/99/9999	INJECTION, CILASTATIN SODIUM; IMPENEM, PER 250 MG	PRIMAXIN IV (VIAL) 500 MG-500 MG	1 EA	VL	IV	EA		250 MG		2	01/01/2002	99/99/9999						
00006-3551-58		J0743		01/01/2002	05/31/2016	INJECTION, CILASTATIN SODIUM; IMPENEM, PER 250 MG	PRIMAXIN IV (ADD-VANTAGE) 250 MG-250 MG	1 EA	VL	IV	EA		250 MG		1	01/01/2002	05/31/2016						
00006-3552-59		J0743		01/01/2002	05/31/2016	INJECTION, CILASTATIN SODIUM; IMPENEM, PER 250 MG	PRIMAXIN IV (ADD-VANTAGE) 500 MG-500 MG	1 EA	VL	IV	EA		250 MG		2	01/01/2002	05/31/2016						
00006-3822-10		J0637		01/01/2003	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CANCIDAS (VIAL) 50 MG	1 EA	VL	IV	EA		5 MG		10	01/01/2003	99/99/9999						
00006-3823-10		J0637		01/01/2003	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CANCIDAS (VIAL) 70 MG	1 EA	VL	IV	EA		5 MG		14	01/01/2003	99/99/9999						
00006-3843-71		J1335		01/01/2004	99/99/9999	INJECTION, ERTAPENEM SODIUM, 500 MG	INVANZ (S.D.V.) 1 GM	1 EA	VL	IJ	EA		500 MG		2	01/01/2004	99/99/9999						
00006-3862-03		J8501		01/01/2005	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND (COMBO PACK) 1 125mg/ 2 80mg	3 EA	PG	PO	EA		5 MG		19	01/01/2005	99/99/9999						
00006-4981-00		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	RECOMBIVAX HB PEDIATRIC/ADOLESCENT (S.D.V., TAX INCL.PF) 5 MCG/0.5 ML	0.5 ML	VL	IM	ML		1 EA		1	01/01/2002	99/99/9999						
00006-4992-00		J3490		07/09/2002	99/99/9999	UNCLASSIFIED DRUGS	RECOMBIVAX HB (S.D.V., TAX INCL.) 40 MCG/ML	1 ML	VL	IM	ML		1 EA		1	07/09/2002	99/99/9999						
00006-4995-00		J3490		07/09/2002	99/99/9999	UNCLASSIFIED DRUGS	RECOMBIVAX HB (S.D.V., TAX INCL.) 10 MCG/ML	1 ML	VL	IM	ML		1 EA		1	07/09/2002	99/99/9999						
00006-4995-41		J3490		07/16/2002	99/99/9999	UNCLASSIFIED DRUGS	RECOMBIVAX HB (S.D.V., TAX INCL.) 10 MCG/ML	1 ML	VL	IM	ML		1 EA		1	07/16/2002	99/99/9999						
00007-3230-02		J1652		02/06/2006	02/04/2014	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 2.5 MG/0.5 ML	0.5 ML	SR	SC	ML		0.5 MG		10	02/06/2006	02/04/2014						
00007-3230-11		J1652		06/03/2005	05/05/2015	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (SRN.PREFL.27GX1/2",PF) 2.5 MG/0.5 ML	0.5 ML	SR	SC	ML		0.5 MG		10	06/03/2005	05/05/2015						
00007-3232-02		J1652		02/06/2006	04/25/2012	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 5 MG/0.4 ML	0.4 ML	SR	SC	ML		0.5 MG		25	02/06/2006	04/25/2012						
00007-3232-11		J1652		11/16/2004	08/06/2015	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 5 MG/0.4 ML	0.4 ML	SR	SC	ML		0.5 MG		25	11/16/2004	08/06/2015						
00007-3234-02		J1652		02/06/2006	04/25/2012	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 7.5 MG/0.6 ML	0.6 ML	SR	SC	ML		0.5 MG		25	02/06/2006	04/25/2012						
00007-3234-11		J1652		11/16/2004	02/10/2016	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 7.5 MG/0.6 ML	0.6 ML	SR	SC	ML		0.5 MG		25	11/16/2004	02/10/2016						
00007-3236-02		J1652		02/06/2006	08/14/2012	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 10 MG/0.8 ML	0.8 ML	SR	SC	ML		0.5 MG		25	02/06/2006	08/14/2012						
00007-3236-11		J1652		11/16/2004	11/12/2015	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 10 MG/0.8 ML	0.8 ML	SR	SC	ML		0.5 MG		25	11/16/2004	11/12/2015						
00007-4401-01		J9261		04/02/2008	10/10/2016	INJECTION, NELARABINE, 50 MG	ARRANON (LATEX-FREE) 5 MG/ML	50 ML	VL	IV	ML		50 MG		0.1	04/02/2008	10/10/2016						
00008-0923-55		J3490		05/18/2004	99/99/9999	UNCLASSIFIED DRUGS	PROTONIX 40 MG	1 EA	VL	IV	EA		1 EA		1	05/18/2004	99/99/9999						
00008-1030-06		J7520		01/01/2002	99/99/9999	SIROLIMUS, ORAL, 1 MG	RAPAMUNE (M.D. BOTTLE) 1 MG/ML	60 ML	BO	PO	ML		1 MG		1	01/01/2002	99/99/9999						
00008-1041-05		J7520		02/01/2006	99/99/9999	SIROLIMUS, ORAL, 1 MG	RAPAMUNE 1 MG	100 EA	BO	PO	EA		1 MG		1	02/01/2006	99/99/9999						
00008-1041-10		J7520		05/26/2006	99/99/9999	SIROLIMUS, ORAL, 1 MG	RAPAMUNE (REDIPAK,10X10) 1 MG	100 EA	BX	PO	EA		1 MG		1	05/26/2006	99/99/9999						
00008-1042-05		J7520		02/01/2006	99/99/9999	SIROLIMUS, ORAL, 1 MG	RAPAMUNE 2 MG	100 EA	BO	PO	EA		1 MG		2	02/01/2006	99/99/9999						
00009-0022-01		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL 8 MG	25 EA	BO	PO	EA		4 MG		2	01/01/2002	99/99/9999						
00009-0049-02		J7509		01/01/2002	10/09/2013	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL 2 MG	100 EA	BO	PO	EA		4 MG		0.5	01/01/2002	10/09/2013						
00009-0056-02		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL 4 MG	100 EA	BO	PO	EA		4 MG		1	01/01/2002	99/99/9999						
00009-0056-04		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL (UNIT OF USE) 4 MG	21 EA	DP	PO	EA		4 MG		1	01/01/2002	99/99/9999						
00009-0073-01		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL 16 MG	50 EA	BO	PO	EA		4 MG		4	01/01/2002	99/99/9999						
00009-0176-01		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL 32 MG	25 EA	BO	PO	EA		4 MG		8	01/01/2002	99/99/9999						
00009-0233-01		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BACITRACIN 50000 U	1 EA	VL	IM	EA		1 EA		1	01/01/2002	99/99/9999						
00009-0271-01		J1000		01/01/2002	99/99/9999	INJECTION, DEPO-ESTRADIOL CYPIONATE, UP TO 5 MG	DEPO-ESTRADIOL (VIAL) 5 MG/ML	5 ML	VL	IM	ML		5 MG		1	01/01/2002	99/99/9999						
00009-0280-02		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (M.D.V.) 40 MG/ML	5 ML	VL	IJ	ML		40 MG		1	01/01/2002	99/99/9999						
00009-0280-03		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (M.D.V.) 40 MG/ML	10 ML	VL	IJ	ML		40 MG		1	01/01/2002	99/99/9999						
00009-0280-51		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (M.D.V.,5X25ML) 40 MG/ML	5 ML	VL	IJ	ML		40 MG		1	01/01/2002	99/99/9999						
00009-0280-52		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (M.D.V.) 40 MG/ML	10 ML	VL	IJ	ML		40 MG		1	01/01/2002	99/99/9999						
00009-0347-02		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	DEPO-TESTOSTERONE (VIAL) 100 MG/ML	10 ML	VL	IM	ML		100 MG		1	01/01/2002	12/31/2014						
00009-0417-01		J1080		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML	1 ML	VL	IM	ML		200 MG		1	01/01/2002	12/31/2014						
00009-0417-02		J1080		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML	10 ML	VL	IM	ML		200 MG		1	01/01/2002	12/31/2014						
00009-0626-01		J1051		01/01/2003	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE, 50 MG	DEPO-PROVERA (VIAL) 400 MG/ML	2.5 ML	VL	IM	ML		50 MG		8	01/01/2003	12/31/2012						
00009-0698-01		J2930		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	SOLU-MEDROL (VIAL) 1 GM	1 EA	VL	IJ	EA		125 MG		8	01/01/2002	99/99/9999						
00009-0728-09		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE 150 MG/ML	60 ML	VL	IJ	ML		1 EA		1	01/01/2002	99/99/9999						
00009-0746-30		J1055		01/01/2002	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	DEPO-PROVERA CONTRACEPTIVE (VIAL) 150 MG/ML	1 ML	VL	IM	ML		150 MG		1	01/01/2002	12/31/2012						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00009-0746-35	J1055			01/01/2002	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	DEPO-PROVERA CONTRACEPTIVE (VIAL,25X1ML) 150 MG/ML	1	ML	VL	IM	ML	150	MG	1	01/01/2002	12/31/2012						
00009-0758-01	J2930			01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	SOLU-MEDROL (VIAL) 500 MG	1	EA	VL	IJ	EA	125	MG	4	01/01/2002	99/99/9999						
00009-0775-26	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE 150 MG/ML	4	ML	VL	IJ	ML	1	EA	1	01/01/2002	99/99/9999						
00009-0796-01	J2930			01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	SOLU-MEDROL (W/DILUENT) 2 GM	1	EA	VL	IJ	EA	125	MG	16	01/01/2002	99/99/9999						
00009-0825-01	J1720			01/01/2002	99/99/9999	INJECTION, HYDROCORTISONE SODIUM SUCCINATE, UP TO 100 MG	SOLU-CORTEF 100 MG	1	EA	VL	IJ	EA	100	MG	1	01/01/2002	99/99/9999						
00009-0870-26	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE 150 MG/ML	2	ML	VL	IJ	ML	1	EA	1	01/01/2002	99/99/9999						
00009-0902-18	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE 150 MG/ML	6	ML	VL	IJ	ML	1	EA	1	01/01/2002	99/99/9999						
00009-3073-01	J1030			01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (S.D.V.) 40 MG/ML	1	ML	VL	IJ	ML	40	MG	1	01/01/2002	99/99/9999						
00009-3073-03	J1030			01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (S.D.V.,25X1ML) 40 MG/ML	1	ML	VL	IJ	ML	40	MG	1	01/01/2002	99/99/9999						
00009-3124-03	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE (ADD-VANTAGE,25X4ML) 150 MG/ML	4	ML	VL	IJ	ML	1	EA	1	01/01/2002	99/99/9999						
00009-3169-06	J0270			01/01/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	PROSTIN VR PEDIATRIC (AMP,5X1ML) 0.5 MG/ML	1	ML	AM	IV	ML	1.25	MCG	400	01/01/2002	99/99/9999						
00009-3375-02	J3490			01/01/2002	06/05/2018	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE (PREMIX) 600 MG/50 ML	50	ML	PC	IV	ML	1	EA	1	01/01/2002	06/05/2018						
00009-3382-02	J3490			01/01/2002	06/01/2018	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE (PREMIX) 900 MG/50 ML	50	ML	PC	IV	ML	1	EA	1	01/01/2002	06/01/2018						
00009-3447-03	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE (ADD-VANTAGE,25X6ML) 150 MG/ML	6	ML	VL	IJ	ML	1	EA	1	01/01/2002	99/99/9999						
00009-3701-05	J0270			01/01/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	CAVERJECT (VIAL) 20 MCG	1	EA	VL	IC	EA	1.25	MCG	16	01/01/2002	99/99/9999						
00009-3778-05	J0270			01/01/2002	10/17/2016	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	CAVERJECT (VIAL) 10 MCG	1	EA	VL	IC	EA	1.25	MCG	8	01/01/2002	10/17/2016						
00009-3794-01	J1742			01/01/2002	99/99/9999	INJECTION, IBUTILIDE FUMARATE, 1 MG	CORVERT (FLIP-TOP VIAL) 0.1 MG/ML	10	ML	VL	IV	ML	1	MG	0.1	01/01/2002	99/99/9999						
00009-5091-01	J9178			01/01/2004	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	ELLECE (S.D.V.,PF) 2 MG/ML	25	ML	VL	IV	ML	2	MG	1	01/01/2004	99/99/9999						
00009-5093-01	J9178			01/01/2004	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	ELLECE (S.D.V.,PF) 2 MG/ML	100	ML	VL	IV	ML	2	MG	1	01/01/2004	99/99/9999						
00009-5137-01	J2020			01/01/2002	99/99/9999	INJECTION, LINEZOLID, 200MG	ZYVOX (P.C.) 2 MG/ML	100	ML	FC	IV	ML	200	MG	0.01	01/01/2002	99/99/9999						
00009-5140-01	J2020			01/01/2002	99/99/9999	INJECTION, LINEZOLID, 200MG	ZYVOX (P.C.) 2 MG/ML	300	ML	FC	IV	ML	200	MG	0.01	01/01/2002	99/99/9999						
00009-5181-01	J0270			06/25/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	CAVERJECT IMPULSE (SYSTEM) 10 MCG	1	EA	BX	IC	EA	1.25	MCG	8	06/25/2002	99/99/9999						
00009-5182-01	J0270			06/25/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	CAVERJECT IMPULSE (SYSTEM) 20 MCG	1	EA	BX	IC	EA	1.25	MCG	16	06/25/2002	99/99/9999						
00009-7224-02	J7504			01/01/2002	99/99/9999	PARENTERAL, 250 MG	ATGAM (AMP,5X5ML) 50 MG/ML	5	ML	AM	IV	ML	250	MG	0.2	01/01/2002	99/99/9999						
00009-7650-02	J0270			01/01/2002	10/17/2016	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	CAVERJECT (SYSTEM) 0.02 MG/ML	2	ML	AM	IC	ML	1.25	MCG	16	05/03/2002	10/17/2016	01/01/2002	03/26/2002	16			
00009-7663-04	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	AROMASIN 25 MG	30	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
00009-7686-04	J0270			01/01/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	CAVERJECT (VIAL) 40 MCG	1	EA	VL	IC	EA	1.25	MCG	32	01/01/2002	99/99/9999						
00024-5924-10	J1817			01/01/2018	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	ADMELOG 100U/1 ML	10	ML	VL	IJ	ML	50	MG	2	01/01/2018	99/99/9999						
00013-2576-91	J9211			01/01/2002	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDAMYCIN PFS (SDV,PF,CYTOSAFE VIAL,PF) 1 MG/ML	5	ML	VL	IV	ML	5	MG	0.2	01/01/2002	99/99/9999						
00013-2586-91	J9211			01/01/2002	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDAMYCIN PFS (SDV,PF,CYTOSAFE VIAL,PF) 1 MG/ML	10	ML	VL	IV	ML	5	MG	0.2	01/01/2002	99/99/9999						
00013-2596-91	J9211			01/01/2002	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDAMYCIN PFS (SDV,PF,CYTOSAFE VIAL,PF) 1 MG/ML	20	ML	VL	IV	ML	5	MG	0.2	01/01/2002	99/99/9999						
00013-2626-81	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN 5.8 MG	1	EA	CT	SC	EA	1	MG	5.8	01/01/2002	99/99/9999						
00013-2646-81	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN 13.8 MG	1	EA	CT	SC	EA	1	MG	13.8	01/01/2002	99/99/9999						
00013-2649-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PREFILLED,PF) 0.2 MG	1	EA	CT	SC	EA	1	MG	0.2	01/01/2002	99/99/9999						
00013-2650-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PREFILLED,PF) 0.4 MG	1	EA	CT	SC	EA	1	MG	0.4	01/01/2002	99/99/9999						
00013-2651-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PREFILLED,PF) 0.6 MG	1	EA	CT	SC	EA	1	MG	0.6	01/01/2002	99/99/9999						
00013-2652-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PREFILLED,PF) 0.8 MG	1	EA	CT	SC	EA	1	MG	0.8	01/01/2002	99/99/9999						
00013-2653-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PREFILLED,PF) 1 MG	1	EA	CT	SC	EA	1	MG	1	01/01/2002	99/99/9999						
00013-2654-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PF) 1.2 MG	1	EA	CT	SC	EA	1	MG	1.2	01/01/2002	99/99/9999						
00013-2655-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PF) 1.4 MG	1	EA	CT	SC	EA	1	MG	1.4	01/01/2002	99/99/9999						
00013-2656-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PF) 1.6 MG	1	EA	CT	SC	EA	1	MG	1.6	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00013-2657-02		J2941		01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PF) 1.8 MG	1 EA	CT	SC	EA		1 MG		1.8	01/01/2002	99/99/9999						
00013-2658-02		J2941		01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK (SRN,PF) 2 MG	1 EA	CT	SC	EA		1 MG		2	01/01/2002	99/99/9999						
00015-0508-42		J8999		01/01/2002	01/31/2017	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGACE 40 MG/ML	240 ML	BO	PO	ML		1 EA		1	01/01/2002	01/31/2017						
00015-3012-60		J9050		04/07/2008	09/30/2015	INJECTION, CARMUSTINE, 100 MG	BICNU (W/DILUENT) 100 MG	1 EA	VL	IV	EA		100 MG		1	04/07/2008	09/30/2015						
00015-3030-20		J8999		01/01/2002	04/04/2013	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	CEENU 10 MG	20 EA	BO	PO	EA		1 EA		1	01/01/2002	04/04/2013						
00015-3031-20		J8999		01/01/2002	04/04/2013	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	CEENU 40 MG	20 EA	BO	PO	EA		1 EA		1	01/01/2002	04/04/2013						
00015-3032-20		J8999		01/01/2002	04/04/2013	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	CEENU 100 MG	20 EA	BO	PO	EA		1 EA		1	01/01/2002	04/04/2013						
00015-3404-20		J9181		01/01/2002	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPHOS (S.D.V.) 100 MG	1 EA	VL	IV	EA		10 MG		10	01/01/2002	99/99/9999						
00023-1145-01		J0585		01/01/2002	99/99/9999	INJECTION, ONABOTULINUMTOXINA, 1 UNIT	BOTOX 100 U	1 EA	VL	IM	EA		1 U		100	01/01/2002	99/99/9999						
00023-9232-01		J0585		06/07/2002	99/99/9999	INJECTION, ONABOTULINUMTOXINA, 1 UNIT	BOTOX COSMETIC 100 U	1 EA	VL	IM	EA		1 U		100	06/07/2002	99/99/9999						
00024-0222-05		J9217		11/01/2003	09/24/2014	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	ELIGARD (SRN,PREFILLED,W/NDL) 22.5 MG	1 EA	SR	SC	EA		7.5 MG		3	11/01/2003	09/24/2014						
00024-0590-10		J9263		06/08/2005	11/03/2015	INJECTION, OXALIPLATIN, 0.5 MG	ELOXATIN (S.D.V.,PF) 5 MG/ML	10 ML	VL	IV	ML		0.5 MG		10	06/08/2005	11/03/2015						
00024-0591-20		J9263		06/08/2005	11/03/2015	INJECTION, OXALIPLATIN, 0.5 MG	ELOXATIN (S.D.V.,PF) 5 MG/ML	20 ML	VL	IV	ML		0.5 MG		10	06/08/2005	11/03/2015						
00024-0592-40		J9263		08/20/2007	07/25/2013	INJECTION, OXALIPLATIN, 0.5 MG	ELOXATIN (SDV,PF) 5 MG/ML	40 ML	VL	IV	ML		0.5 MG		10	08/20/2007	07/25/2013						
00024-0605-45		J9217		02/18/2005	09/24/2014	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	ELIGARD (SINGLE-USE KIT) 45 MG	1 EA	BX	SC	EA		7.5 MG		6	02/18/2005	09/24/2014						
00024-0610-30		J9217		03/04/2003	09/24/2014	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	ELIGARD (SINGLE-USE) 30 MG	1 EA	BX	SC	EA		7.5 MG		4	03/04/2003	09/24/2014						
00024-0793-75		J9217		07/25/2003	09/24/2014	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	ELIGARD (SRN,PREFILLED,W/NDL) 7.5 MG	1 EA	SR	SC	EA		7.5 MG		1	07/25/2003	09/24/2014						
00024-5150-10		J2783		01/01/2004	99/99/9999	INJECTION, RASBURICASE, 0.5 MG	ELITEK (3 S.D.V. W/DILUENT,PF) 1.5 MG	1 EA	VL	IV	EA		0.5 MG		3	01/01/2004	99/99/9999						
00024-5151-75		J2783		06/27/2006	99/99/9999	INJECTION, RASBURICASE, 0.5 MG	ELITEK (SDV,W/DILUENT) 7.5 MG	1 EA	VL	IV	EA		0.5 MG		15	06/27/2006	99/99/9999						
00026-8196-36		J0365		01/01/2006	01/29/2016	INJECTION, APROTONIN, 10,000 KIU	TRASYLOL 10000 KIU/ML	100 ML	VL	IV	ML		10000 KIU		1	01/01/2006	01/29/2016						
00026-8197-63		J0365		01/01/2006	01/29/2016	INJECTION, APROTONIN, 10,000 KIU	TRASYLOL 10000 KIU/ML	200 ML	VL	IV	ML		10000 KIU		1	01/01/2006	01/29/2016						
00029-6571-26		J3490		01/01/2002	11/17/2014	UNCLASSIFIED DRUGS	TIMENTIN (VIAL) 100 MG-3 GM	1 EA	VL	IV	EA		1 EA		1	01/01/2002	11/17/2014						
00029-6571-31		J3490		01/01/2002	11/21/2014	UNCLASSIFIED DRUGS	TIMENTIN (PREMIX) 100 MG/100 ML-3 GM/100 ML	100 ML	FC	IV	ML		1 EA		1	01/01/2002	11/21/2014						
00029-6571-40		J3490		01/01/2002	08/27/2012	UNCLASSIFIED DRUGS	TIMENTIN (ADD-VANTAGE) 100 MG-3 GM	1 EA	VL	IV	EA		1 EA		1	01/01/2002	08/27/2012						
00029-6579-21		J3490		01/01/2002	12/02/2014	UNCLASSIFIED DRUGS	TIMENTIN (BULK VIAL) 1 GM-30 GM	1 EA	VL	IV	EA		1 EA		1	01/01/2002	12/02/2014						
00039-0017-10		J0698		01/01/2002	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	CLAFORAN (VIAL) 500 MG	1 EA	VL	IJ	EA		1 GM		0.5	01/01/2002	99/99/9999						
00039-0018-10		J0698		01/01/2002	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	CLAFORAN (VIAL) 1 GM	1 EA	VL	IJ	EA		1 GM		1	01/01/2002	99/99/9999						
00039-0018-49		J0698		04/01/2006	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	NOVAPLUS CLAFORAN 1 GM	1 EA	VL	IJ	EA		1 GM		1	04/01/2006	99/99/9999						
00039-0019-10		J0698		01/01/2002	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	CLAFORAN (VIAL) 2 GM	1 EA	VL	IJ	EA		1 GM		2	01/01/2002	99/99/9999						
00039-0019-49		J0698		06/01/2005	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	NOVAPLUS CLAFORAN 2 GM	1 EA	VL	IJ	EA		1 GM		2	06/01/2005	99/99/9999						
00039-0020-01		J0698		01/01/2002	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	CLAFORAN (BULK VIAL) 10 GM	1 EA	GC	IJ	EA		1 GM		10	01/01/2002	99/99/9999						
00039-0020-49		J0698		06/01/2005	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	NOVAPLUS CLAFORAN (PHARMACY BULK PACKAGE) 10 GM	1 EA	GC	IJ	EA		1 GM		10	06/01/2005	99/99/9999						
00039-0023-25		J0698		01/01/2002	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	CLAFORAN (ADD-VANTAGE) 1 GM	1 EA	VL	IJ	EA		1 GM		1	01/01/2002	99/99/9999						
00039-0023-49		J0698		06/01/2005	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	NOVAPLUS CLAFORAN (ADD-VANTAGE SYSTEM) 1 GM	1 EA	VL	IJ	EA		1 GM		1	06/01/2005	99/99/9999						
00039-0023-61		J0698		04/03/2006	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	AMERINET CLAFORAN 1 GM	1 EA	VL	IJ	EA		1 GM		1	04/03/2006	99/99/9999						
00039-0024-25		J0698		01/01/2002	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	CLAFORAN (ADD-VANTAGE) 2 GM	1 EA	VL	IJ	EA		1 GM		2	01/01/2002	99/99/9999						
00039-0024-49		J0698		06/01/2005	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	NOVAPLUS CLAFORAN (ADD-VANTAGE SYSTEM) 2 GM	1 EA	VL	IJ	EA		1 GM		2	06/01/2005	99/99/9999						
00039-0024-50		J0698		01/01/2002	09/11/2013	INJECTION, CEFOTAXIME SODIUM, PER GM	CLAFORAN (ADD-VANTAGE) 2 GM	1 EA	VL	IJ	EA		1 GM		2	01/01/2002	09/11/2013						
00046-0749-05		J1410		01/01/2002	99/99/9999	INJECTION, ESTROGEN CONJUGATED, PER 25 MG	PREMARIN INTRAVENOUS (W/SECULE VIAL) 25 MG	1 EA	VL	IV	EA		25 MG		1	01/01/2002	99/99/9999						
00049-0013-83		J0295		01/01/2002	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	UNASYN (VIAL) 1 GM-0.5 GM	1 EA	VL	IJ	EA		1.5 GM		1	01/01/2002	99/99/9999						
00049-0014-83		J0295		01/01/2002	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	UNASYN (VIAL) 2 GM-1 GM	1 EA	VL	IJ	EA		1.5 GM		2	01/01/2002	99/99/9999						
00049-0022-83		J0295		01/01/2002	04/05/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	UNASYN (P.B.,ADD-VANTAGE) 1 GM-0.5 GM	1 EA	VL	IV	EA		1.5 GM		1	01/01/2002	04/05/2013						
00049-0024-28		J0295		01/01/2002	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	UNASYN (BULK PACKAGE) 10 GM-5 GM	1 EA	VL	IV	EA		1.5 GM		10	01/01/2002	99/99/9999						
00074-0124-03		J0135		08/06/2018	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA PEN STARTER PACK (PF,LATEX-FREE) 80 MG/0.8 ML	3 EA	BX	SC	EA		20 MG		4	08/06/2018	99/99/9999						
64380-0725-07		J7517		05/01/2014	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (USP,FILM-COATED) 500 MG	500 EA	BO	PO	EA		250 MG		2	05/01/2014	99/99/9999						
00049-0520-83		J2540		01/01/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PFIZERPEN (VIAL, PHARMACY BOTTLE) 5 Million U	1 EA	VL	IV	EA		600000 U		8.33333	01/01/2002	99/99/9999						
00049-0530-28		J2540		01/01/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PFIZERPEN (VIAL, PHARMACY BOTTLE) 20 Million U	1 EA	VL	IV	EA		600000 U		33.33333	01/01/2002	99/99/9999						
00049-3190-28		J3465		01/01/2004	99/99/9999	INJECTION, VORICONAZOLE, 10 MG	VFEND I.V. (S.D.V.) 200 MG	1 EA	VL	IV	EA		10 MG		20	01/01/2004	99/99/9999						
00049-3382-25		J3490		10/19/2005	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (FTV,LATEX-FREE) 50 MCG/ML	5 ML	VL	IJ	ML		1 EA		1	10/19/2005	99/99/9999						
00049-3920-83		J3486		01/01/2004	99/99/9999	INJECTION, ZIPRASIDONE MESYLATE, 10 MG	GEODON 20 MG	1 EA	VL	IM	EA		10 MG		2	01/01/2004	99/99/9999						
00051-0021-21		Q0167		01/01/2002	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A																	

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00051-0022-21		Q0168		08/14/2006	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFT GELATIN) 5 MG	60	EA	BO	PO	EA	5	MG	1	08/14/2006	12/31/2013							
00051-0023-21		Q0168		01/01/2002	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 10 MG	60	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2013							
00052-0301-51		J3490		05/01/2003	99/99/9999	UNCLASSIFIED DRUGS	GANIRELIX ACETATE 250 MCG/0.5 ML	0.5	ML	SR	SC	ML	1	EA	1	05/01/2003	99/99/9999							
00052-0315-10		J0725		01/01/2002	99/99/9999	INJECTION, CHORIONIC GONADOTROPIN, PER 1,000 USP UNITS	PREGNYL (W/DILUENT) 10000 U	1	EA	VL	IM	EA	1000	Units	10	01/01/2002	99/99/9999							
00052-0602-02		J9031		01/01/2002	99/99/9999	BCG (INTRAVESICAL) PER INSTILLATION	TICE BCG (VIAL) 800 Million CFU	1	EA	VL	IL	EA	1	INSTILLATION	1	01/01/2002	99/99/9999							
00052-0603-02		J9031		01/01/2002	99/99/9999	BCG (INTRAVESICAL) PER INSTILLATION	BCG VACCINE (VIAL)	1	EA	VL	ID	EA	1	INSTILLATION	1	01/01/2002	99/99/9999							
00053-7596-10		J1562		01/01/2007	05/06/2013	INJECTION, IMMUNE GLOBULIN (VIVAGLOBIN), 100 MG	VIVAGLOBIN (PF) 160 MG/ML	10	ML	VL	SC	ML	100	MG	1.6	01/01/2007	05/06/2013							
00053-7596-20		J1562		01/01/2007	06/08/2013	INJECTION, IMMUNE GLOBULIN (VIVAGLOBIN), 100 MG	VIVAGLOBIN (PF) 160 MG/ML	20	ML	VL	SC	ML	100	MG	1.6	01/01/2007	06/08/2013							
00054-0017-20		J7506		12/01/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (10X10) 10 MG	100	EA	BX	PO	EA	5	MG	2	12/01/2004	12/31/2015							
00054-0017-25		J7506		01/01/2005	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2005	12/31/2015							
00054-0017-29		J7506		12/01/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	500	EA	BO	PO	EA	5	MG	2	12/01/2004	12/31/2015							
00054-0018-20		J7506		09/07/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (10X10) 20 MG	100	EA	BX	PO	EA	5	MG	4	09/07/2004	12/31/2015							
00054-0018-25		J7506		10/14/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	100	EA	BO	PO	EA	5	MG	4	10/14/2004	12/31/2015							
00054-0018-29		J7506		10/08/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	500	EA	BO	PO	EA	5	MG	4	10/08/2004	12/31/2015							
00054-0019-20		J7506		09/24/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (10X10) 50 MG	100	EA	BX	PO	EA	5	MG	10	09/24/2004	12/31/2015							
00054-0019-25		J7506		08/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 50 MG	100	EA	BO	PO	EA	5	MG	10	08/10/2004	12/31/2015							
00054-3025-02		J7608		01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 10%	30	ML	VL	IH	ML	1	GM	0.1	01/01/2002	04/03/2014							
00054-3025-02	KO	J7608	KO	01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 10%	30	ML	VL	IH	ML	1	GM	0.1	01/01/2002	04/03/2014							
00054-3026-02		J7608		01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 20%	30	ML	VL	IH	ML	1	GM	0.2	01/01/2002	04/03/2014							
00054-3026-02	KO	J7608	KO	01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 20%	30	ML	VL	IH	ML	1	GM	0.2	01/01/2002	04/03/2014							
00054-3027-02		J7608		01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 10%	10	ML	VL	IH	ML	1	GM	0.1	01/01/2002	04/03/2014							
00054-3027-02	KO	J7608	KO	01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 10%	10	ML	VL	IH	ML	1	GM	0.1	01/01/2002	04/03/2014							
00054-3028-02		J7608		01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 20%	10	ML	VL	IH	ML	1	GM	0.2	01/01/2002	04/03/2014							
00054-3028-02	KO	J7608	KO	01/01/2002	04/03/2014	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 20%	10	ML	VL	IH	ML	1	GM	0.2	01/01/2002	04/03/2014							
00054-3176-44		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE INTENSOL 1 MG/ML	30	ML	BO	PO	ML	0.25	MG	4	01/01/2006	99/99/9999							
00054-3542-58		J8999		04/11/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (LEMON LIME) 40 MG/ML	240	ML	BO	PO	ML	1	EA	1	04/11/2002	99/99/9999							
00054-3721-44		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE INTENSOL 5 MG/ML	30	ML	BO	PO	ML	5	MG	1	01/01/2002	12/31/2015							
00054-3722-50		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (PEPPERMINT-VANILLA) 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	01/01/2002	12/31/2015							
00054-3722-63		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (PEPPERMINT-VANILLA) 5 MG/5 ML	500	ML	BO	PO	ML	5	MG	0.2	01/01/2002	12/31/2015							
00054-4084-25		J7500		01/01/2002	04/01/2017	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	04/01/2017							
00054-4129-25		None		03/28/2000	07/11/2016	CYCLOPHOSPHAMIDE, 25 MG, ORAL	CYCLOPHOSPHAMIDE 25 MG	100	EA	BO	PO	EA	25	MG	1	03/28/2000	07/11/2016							
00054-4130-25		None		03/28/2000	07/11/2016	CYCLOPHOSPHAMIDE, 50 MG, ORAL	CYCLOPHOSPHAMIDE 50 MG	100	EA	BO	PO	EA	50	MG	1	03/28/2000	07/11/2016							
00054-4179-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	100	EA	BO	PO	EA	0.25	MG	2	01/01/2006	99/99/9999							
00054-4180-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	100	EA	BO	PO	EA	0.25	MG	3	01/01/2006	99/99/9999							
00054-4181-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 1 MG	100	EA	BO	PO	EA	0.25	MG	4	01/01/2006	99/99/9999							
00054-4182-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 1.5 MG	100	EA	BO	PO	EA	0.25	MG	6	01/01/2006	99/99/9999							
00054-4183-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 2 MG	100	EA	BO	PO	EA	0.25	MG	8	01/01/2006	99/99/9999							
00054-4184-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	100	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999							
00054-4550-15		None		09/27/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	36	EA	BO	PO	EA	2.5	MG	1	09/27/1994	99/99/9999							
00054-4550-25		None		09/27/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	100	EA	BO	PO	EA	2.5	MG	1	09/27/1994	99/99/9999							
00054-4581-11		J8999		02/19/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MERCAPTOPYRINE (USP) 50 MG	25	EA	BO	PO	EA	1	EA	1	02/19/2004	99/99/9999							
00054-4581-27		J8999		02/19/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MERCAPTOPYRINE (USP) 50 MG	250	EA	BO	PO	EA	1	EA	1	02/19/2004	99/99/9999							
00054-4603-25		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 20 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
00054-4604-25		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
00054-4728-25		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
00054-4728-31		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	1000	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
00054-4741-25		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 1 MG	100	EA	BO	PO	EA	5	MG	0.2	01/01/2002	12/31/2015							
00054-4741-31		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 1 MG	1000	EA	BO	PO	EA	5	MG	0.2	01/01/2002	12/31/2015							
00054-4742-25		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 2.5 MG	100	EA	BO	PO	EA	5	MG	0.5	01/01/2002	12/31/2015							
00054-8084-25		J7500		01/01/2002																				

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00054-8175-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (10X10) 4 MG	100	EA	BX	PO	EA	0.25	MG	16	01/01/2006	99/99/9999							
00054-8176-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (10X10) 2 MG	100	EA	BX	PO	EA	0.25	MG	8	01/01/2006	99/99/9999							
00054-8179-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (10X10) 0.5 MG	100	EA	BX	PO	EA	0.25	MG	2	01/01/2006	99/99/9999							
00054-8180-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (10X10) 0.75 MG	100	EA	BX	PO	EA	0.25	MG	3	01/01/2006	99/99/9999							
00054-8181-25		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (10X10) 1.5 MG	100	EA	BX	PO	EA	0.25	MG	6	01/01/2006	99/99/9999							
00054-8550-25	None			09/27/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM (10X10) 2.5 MG	100	EA	BX	PO	EA	2.5	MG	1	09/27/1994	99/99/9999							
00054-8603-25	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (10X10) 20 MG	100	EA	BX	PO	EA	1	EA	1	01/01/2002	99/99/9999							
00054-8604-25	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (10X10) 40 MG	100	EA	BX	PO	EA	1	EA	1	01/01/2002	99/99/9999							
00054-8722-16	J7506			01/01/2002	09/09/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE (PEPPERMINT-VANILLA) 5 MG/5 ML	5	ML	CP	PO	ML	5	MG	0.2	01/01/2002	09/09/2014							
00054-8724-25	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (10X10) 5 MG	100	EA	BX	PO	EA	5	MG	1	01/01/2002	12/31/2015							
00054-8739-25	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (10X10) 1 MG	100	EA	BX	PO	EA	5	MG	0.2	01/01/2002	12/31/2015							
00054-8740-25	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (10X10) 2.5 MG	100	EA	BX	PO	EA	5	MG	0.5	01/01/2002	12/31/2015							
00065-0543-01	J3301			11/29/2007	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIESENCE 40 MG/ML	1	ML	VL	IJ	ML	10	MG	4	11/29/2007	99/99/9999							
00068-0597-01	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	RIFADIN IV (VIAL) 600 MG	1	EA	VL	IV	EA	1	EA	1	01/01/2002	99/99/9999							
00069-3051-07	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX (SINGLE DOSE PACKETS) 1 GM/Package	10	EA	BX	PO	EA	1	GM	1	01/01/2002	99/99/9999							
00069-3051-75	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX (SINGLE DOSE PACKETS) 1 GM/Package	3	PK	BX	PO	EA	1	GM	1	01/01/2002	99/99/9999							
00069-3060-30	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	30	EA	BO	PO	EA	1	GM	0.25	01/01/2002	99/99/9999							
00069-3060-75	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX Z-PAK (3X6) 250 MG	18	EA	DP	PO	EA	1	GM	0.25	01/01/2002	99/99/9999							
00069-3060-86	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	50	EA	BX	PO	EA	1	GM	0.25	01/01/2002	99/99/9999							
00069-3070-30	Q0144			08/06/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 500 MG	30	EA	BO	PO	EA	1	GM	0.5	08/06/2002	99/99/9999							
00069-3070-75	Q0144			08/06/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX TRI-PAK (3X3) 500 MG	9	EA	DP	PO	EA	1	GM	0.5	08/06/2002	99/99/9999							
00069-3070-86	Q0144			10/21/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX (5 X 10) 500 MG	50	EA	BX	PO	EA	1	GM	0.5	10/21/2002	99/99/9999							
00069-3080-30	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 600 MG	30	EA	BO	PO	EA	1	GM	0.6	01/01/2002	99/99/9999							
00069-3110-19	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	01/01/2002	99/99/9999							
00069-3120-19	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.04	01/01/2002	99/99/9999							
00069-3130-19	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	22.5	ML	BO	PO	ML	1	GM	0.04	01/01/2002	99/99/9999							
00069-3140-19	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	30	ML	BO	PO	ML	1	GM	0.04	01/01/2002	99/99/9999							
00069-3150-14	J0456			02/25/2002	01/10/2013	INJECTION, AZITHROMYCIN, 500 MG	ZITHROMAX (W/VIAL M/TE) 500 MG	1	EA	VL	IV	EA	500	MG	1	02/25/2002	01/10/2013							
00069-3150-83	J0456			01/01/2002	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	ZITHROMAX (VIAL) 500 MG	1	EA	VL	IV	EA	500	MG	1	01/01/2002	99/99/9999							
00069-5410-66	Q0177			01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VISTARIL 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	99/99/9999							
00069-5420-66	Q0178			01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VISTARIL 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013							
00074-1658-01	J2501			01/01/2003	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	ZEMPLAR (S.D.V., FLIPTOP) 0.005 MG/ML	1	ML	VL	IV	ML	1	MCG	5	01/01/2003	99/99/9999							
68982-0820-84	J1599			11/12/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), NOT OTHERWISE SPECIFIED, 500 MG	PANZYGA (INNER PACK,PF) 100 MG/1 ML SODIUM CHLORIDE (INTERLINK,50X2ML,PF) 0.9%	10	ML	BO	IV	ML	500	MG	0.2	11/12/2018	99/99/9999							
00074-1812-22	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	2	ML	SR	IV	ML	10	ML	0.1	01/01/2007	02/03/2016							
00074-2287-54	J1885			01/01/2002	10/17/2016	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE NOVATION (LL LATEX-FREE, CARPUJECT) 30 MG/ML	1	ML	SR	IJ	ML	15	MG	2	01/01/2002	10/17/2016							
00074-3454-25	J1642			02/20/2002	10/17/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (ANSYR, LATEX-FREE) 100 U/ML	5	ML	SR	IV	ML	10	U	10	02/20/2002	10/17/2016							
00074-3799-02	J0135			01/01/2005	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PF, PREFILLED SYRINGE) 40 MG/0.8 ML	2	EA	BX	MR	EA	20	MG	2	01/01/2005	99/99/9999							
00069-0313-10	J2185			05/29/2018	99/99/9999	INJECTION, MEROPENEM, 100 MG	MERREM IV 500 MG	10	EA	VL	IV	EA	100	MG	5	05/29/2018	99/99/9999							
00069-0314-10	J2185			05/29/2018	99/99/9999	INJECTION, MEROPENEM, 100 MG	MERREM IV 1 GM	10	EA	VL	IV	EA	100	MG	10	05/29/2018	99/99/9999							
00074-3834-02	J3480			01/01/2002	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (AMP, LATEX-FREE) 2 MEQ/ML	20	ML	AM	IV	ML	2	MEQ	1	01/01/2002	10/17/2016							
00074-4141-03	J1265			01/01/2006	10/17/2016	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL 5%-80 MG/100 ML	500	ML	GC	IV	ML	40	MG	0.02	01/01/2006	10/17/2016							
00074-4332-01	J3370			01/01/2002	02/03/2016	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (VIAL, FLIPTOP) 500 MG	1	EA	VL	IV	EA	500	MG	1	03/01/2009	02/03/2016	01/01/2002	04/24/2005	1				
00074-4339-02	J0135			07/17/2006	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (SINGLE-USE PEN; 2X1ML) 40 MG/0.8 ML	2	EA	BX	MR	EA	20	MG	2	07/17/2006	99/99/9999							
00074-4339-06	J0135			02/27/2007	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (SINGLE-USE PEN; 6X1ML) 40 MG/0.8 ML	6	EA	BX	MR	EA	20	MG	2	02/27/2007	99/99/9999							
00074-4637-01	J2501			01/01/2003	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	ZEMPLAR (VIAL, FLIPTOP) 0.002 MG/ML	1	ML	VL	IV	ML	1	MCG	2	01/01/2003	99/99/9999							
00074-4729-01	J1250			01/01/2002	10/17/2016	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	AMINOPHYLLINE (10X10ML, ABBOJECT) 25 MG/ML	100	ML	VL	IV	ML	250	MG	0.05	01/01/2002	10/17/2016							
00074-4909-18	J0280			01/01/2002	03/24/2016	INJECTION, AMINOPHYLLIN, UP TO 250 MG	SODIUM CHLORIDE (ANSYR, FOR IV, 50X5ML, PF) 0.9%	10	ML	SR	IV	ML	250	MG	0.1	01/01/2002	03/24/2016							
00074-5365-05	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, D																		

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00074-5641-25		J799		01/01/2002	10/17/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRSE (1000 ML CONTAINER) 10% BUPIVACAINE HCL (W/MALE ADAPTER)	500 ML	GC	IV	ML		1 EA		1	01/01/2002	10/17/2016						
00074-5749-22	J3490			01/01/2002	03/25/2016	UNCLASSIFIED DRUGS		50 ML	SR	IJ	ML		1 EA		1	01/01/2002	03/25/2016						
00074-6463-32	J7515			01/01/2002	12/07/2015	CYCLOSPORINE, ORAL, 25 MG	GENGRAF (BLISTER PACK) 25 MG	30 EA	BX	PO	EA		25 MG		1	01/01/2002	12/07/2015						
00074-6476-44	J1364			01/01/2002	10/17/2016	INJECTION, ERYTHROMYCIN LACTOBIONATE, PER 500 MG	ERYTHROCIN LACTOBIONATE (ADD-VANTAGE,LATEX-FREE) 500 MG	1 EA	VL	IV	EA		500 MG		1	03/01/2009	10/17/2016	01/01/2002	03/09/2006	1			
00074-6479-32	J7502			01/01/2002	11/09/2015	CYCLOSPORINE, ORAL, 100 MG	GENGRAF (BLISTER PACK) 100 MG	30 EA	BX	PO	EA		100 MG		1	01/01/2002	11/09/2015						
00074-7269-50	J7502			01/18/2002	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	GENGRAF 100 MG/ML	50 ML	BO	PO	ML		100 MG		1	01/18/2002	99/99/9999						
00074-8065-15	J0330			01/01/2002	10/17/2016	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	QUELICIN 20 MG/ML	5 ML	SR	IV	ML		20 MG		1	01/01/2002	10/17/2016						
00074-9374-02	J0135			02/22/2008	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (SINGLE-DOSE,PF) 20 MG/0.4 ML FUROSEMIDE (ANSYR,LATEX-FREE) 10 MG/ML	2 EA	BX	SC	EA		20 MG		1	02/22/2008	99/99/9999						
00074-9631-04	J1940			01/01/2002	02/03/2016	INJECTION, FUROSEMIDE, UP TO 20 MG		4 ML	SR	IJ	ML		20 MG		0.5	03/01/2009	02/03/2016	01/01/2002	04/20/2006	0.5			
00075-0620-40	J1650			01/01/2002	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX 40 MG/0.4 ML	0.4 ML	SR	IJ	ML		10 MG		10	01/01/2002	99/99/9999						
00075-0620-41	J1650			03/17/2008	04/01/2015	INJECTION, ENOXAPARIN SODIUM, 10 MG	NOVAPLUS LOVENOX (10X0.4ML,SINGLE-DOSE,PF) 40 MG/0.4 ML	0.4 ML	SR	SC	ML		10 MG		10	03/17/2008	04/01/2015						
00075-0621-60	J1650			01/01/2002	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (SRN,PREFILLED) 60 MG/0.6 ML	0.6 ML	SR	IJ	ML		10 MG		10	01/01/2002	99/99/9999						
00075-0621-61	J1650			03/11/2008	04/01/2015	INJECTION, ENOXAPARIN SODIUM, 10 MG	NOVAPLUS LOVENOX (10X0.6ML,SINGLE-DOSE,PF) 60 MG/0.6 ML	0.6 ML	SR	SC	ML		10 MG		10	03/11/2008	04/01/2015						
00075-0622-80	J1650			01/01/2002	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (SRN,PREFILLED) 80 MG/0.8 ML	0.8 ML	SR	IJ	ML		10 MG		10	01/01/2002	99/99/9999						
00075-0622-81	J1650			03/11/2008	04/01/2015	INJECTION, ENOXAPARIN SODIUM, 10 MG	NOVAPLUS LOVENOX (10X0.8ML,SINGLE-DOSE,PF) 80 MG/0.8 ML	0.8 ML	SR	SC	ML		10 MG		10	03/11/2008	04/01/2015						
00075-0623-00	J1650			01/01/2002	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (SRN,PREFILLED) 100 MG/ML	1 ML	SR	IJ	ML		10 MG		10	01/01/2002	99/99/9999						
00075-0623-01	J1650			03/11/2008	04/01/2015	INJECTION, ENOXAPARIN SODIUM, 10 MG	NOVAPLUS LOVENOX (10X1ML,SINGLE-DOSE,PF) 100 MG/ML	1 ML	SR	SC	ML		10 MG		10	03/11/2008	04/01/2015						
00075-0624-30	J1650			01/01/2002	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (SRN) 30 MG/0.3 ML	0.3 ML	SR	IJ	ML		10 MG		10	01/01/2002	99/99/9999						
00075-0624-31	J1650			03/17/2008	04/01/2015	INJECTION, ENOXAPARIN SODIUM, 10 MG	NOVAPLUS LOVENOX (10X0.3ML,SINGLE-DOSE,PF) 30 MG/0.3 ML	0.3 ML	SR	SC	ML		10 MG		10	03/17/2008	04/01/2015						
00075-0626-03	J1650			03/07/2003	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (VIAL,MULTIPLE DOSE VIAL) 100 MG/ML	3 ML	VL	SC	ML		10 MG		10	03/07/2003	99/99/9999						
00075-0626-04	J1650			03/11/2008	04/01/2015	INJECTION, ENOXAPARIN SODIUM, 10 MG	NOVAPLUS LOVENOX (1X3ML,MULTIPLE-DOSE) 100 MG/ML	3 ML	VL	IJ	ML		10 MG		10	03/11/2008	04/01/2015						
00075-2451-01	J2597			01/01/2002	04/14/2015	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DDAVP (AMP) 4 MCG/ML	1 ML	AM	IJ	ML		1 MCG		4	01/01/2002	04/14/2015						
00075-2451-53	J2597			01/01/2002	05/09/2015	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DDAVP (VIAL) 4 MCG/ML	10 ML	VL	IJ	ML		1 MCG		4	01/01/2002	05/09/2015						
00075-2912-01	J1650			01/01/2002	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX 120 MG/0.8 ML	0.8 ML	SR	IJ	ML		10 MG		15	01/01/2002	99/99/9999						
00075-2915-01	J1650			01/01/2002	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (W/AUTO SAFETY DEVICE) 150 MG/ML	1 ML	SR	IJ	ML		10 MG		15	01/01/2002	99/99/9999						
00078-0053-03	J2210			01/01/2002	04/16/2012	INJECTION, METHYLERGONOVINE MALEATE, UP TO 0.2 MG	METHERGINE (AMP) 0.2 MG/ML	1 ML	AM	IJ	ML		0.2 MG		1	01/01/2002	04/16/2012						
00078-0109-01	J7516			01/01/2002	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	SANDIMMUNE (AMP) 50 MG/ML	5 ML	AM	IV	ML		250 MG		0.2	01/01/2002	99/99/9999						
00078-0110-22	J7502			01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	SANDIMMUNE 100 MG/ML	50 ML	BO	PO	ML		100 MG		1	01/01/2002	99/99/9999						
00078-0149-23	J0630			01/01/2002	08/30/2015	INJECTION, CALCITONIN SALMON, UP TO 400 UNITS	MIACALCIN (VIAL) 200 IU/ML	2 ML	VL	IJ	ML		400 U		0.5	01/01/2002	08/30/2015						
00078-0180-01	J2354			01/01/2004	99/99/9999	INJECTION, 25 MCG	SANDOSTATIN (AMP) 50 MCG/ML	1 ML	AM	IJ	ML		25 MCG		2	01/01/2004	99/99/9999						
00078-0181-01	J2354			01/01/2004	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	SANDOSTATIN (AMP) 100 MCG/ML	1 ML	AM	IJ	ML		25 MCG		4	01/01/2004	99/99/9999						
00078-0182-01	J2354			01/01/2004	99/99/9999	INJECTION, 25 MCG	SANDOSTATIN (AMP) 500 MCG/ML	1 ML	AM	IJ	ML		25 MCG		20	01/01/2004	99/99/9999						
00078-0183-25	J2354			01/01/2004	03/15/2018	INJECTION, 25 MCG	SANDOSTATIN (M.D.V.) 200 MCG/ML	5 ML	VL	IJ	ML		25 MCG		8	01/01/2004	03/15/2018						
00078-0184-25	J2354			01/01/2004	06/05/2016	INJECTION, 25 MCG	SANDOSTATIN (M.D.V.) 1000 MCG/ML	5 ML	VL	IJ	ML		25 MCG		40	01/01/2004	06/05/2016						
00078-0240-15	J7515			01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	SANDIMMUNE (SANDOPAK,SOFTGEL) 25 MG	30 EA	BX	PO	EA		25 MG		1	01/01/2002	99/99/9999						
00078-0241-15	J7502			01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	SANDIMMUNE (SOFTGEL) 100 MG	30 EA	BX	PO	EA		100 MG		1	01/01/2002	99/99/9999						
00078-0246-15	J7515			01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	NEORAL (SOFTGEL) 25 MG	30 EA	BX	PO	EA		25 MG		1	01/01/2002	99/99/9999						
00078-0248-15	J7502			01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	NEORAL (SOFTGEL) 100 MG	30 EA	BX	PO	EA		100 MG		1	01/01/2002	99/99/9999						
00078-0274-22	J7502			01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	NEORAL 100 MG/ML	50 ML	BO	PO	ML		100 MG		1	01/01/2002	99/99/9999						
00078-0331-84	J0480			01/01/2006	99/99/9999	INJECTION, BASILIXIMAB, 20 MG	SIMULECT (S.D.V.,PF) 20 MG	1 EA	VL	IV	EA		20 MG		1	01/01/2006	99/99/9999						
00078-0340-61	J2353			07/26/2004	09/23/2015	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1&1/2"X19G.PFS) 10 MG	1 EA	BX	IM	EA		1 MG		10	07/26/2004	09/23/2015						
00078-0341-61	J2353			08/18/2004	09/23/2015	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1&1/2"X19G.PFS) 20 MG	1 EA	BX	IM	EA		1 MG		20	08/18/2004	09/23/2015						
00078-0342-61	J2353			07/14/2004	09/23/2015	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1&1/2"X19G.PFS) 30 MG	1 EA	BX	IM	EA		1 MG		30	07/14/2004	09/23/2015						
00078-0347-51	J0895			01/01/2002	08/14/2015	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DESFERAL (VIAL) 2 GM	1 EA	VL	IJ	EA		500 MG		4	01/01/2002	08/14/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00078-0385-66		J7518		01/01/2005	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYFORTIC (K-30,FILM-COATED) 180 MG	120 EA	BO	PO	EA	EA	180 MG		1	01/01/2005	99/99/9999						
00078-0386-66		J7518		01/01/2005	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYFORTIC (K-30,FILM-COATED) 360 MG	120 EA	BO	PO	EA	EA	180 MG		2	01/01/2005	99/99/9999						
00078-0393-61		J0480		01/01/2006	99/99/9999	INJECTION, BASILIXIMAB, 20 MG	SIMULECT (S.D.V.,PF) 10 MG	1 EA	VL	IV	EA	EA	20 MG		0.5	01/01/2006	99/99/9999						
00078-0435-61		J3488		01/01/2008	12/31/2013	INJECTION, ZOLEDRONIC ACID (RECLAST), 1 MG	RECLAST	100 ML	PC	IV	ML	ML	1 MG		0.05	01/01/2008	12/31/2013						
00078-0467-91		J0895		05/01/2007	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DESFERAL (USP) 500 MG	1 EA	VL	IJ	EA	EA	500 MG		1	05/01/2007	99/99/9999						
00078-0494-71		J7682		04/01/2008	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBI (56X5ML,SDA,PF)	5 ML	PC	IH	ML	ML	300 MG		0.2	04/01/2008	99/99/9999						
00078-0494-71	KO	J7682	KO	04/01/2008	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBI (56X5ML,SDA,PF)	5 ML	PC	IH	ML	ML	300 MG		0.2	04/01/2008	99/99/9999						
47335-0177-95	J3490			09/17/2018	12/31/2018	UNCLASSIFIED DRUGS	ILUMYA (PF) 100 MG/1 ML	1 ML	SR	SC	ML	ML	1 MG		1	09/17/2018	12/31/2018						
69794-0001-01	J3490			11/15/2017	12/31/2018	UNCLASSIFIED DRUGS	MEPSEVII (PF) 2 MG/1 ML	5 ML	VL	IV	ML	ML	1 MG		1	11/15/2017	12/31/2018						
50419-0385-01	J3490			09/18/2017	12/31/2018	UNCLASSIFIED DRUGS	ALIQOPA (LYOPHILIZED) 60 MG	1 EA	VL	IV	EA	EA	1 MG		1	09/18/2017	12/31/2018						
00085-0539-01	J9214			01/01/2002	05/28/2016	INJECTION, INTERFERON, ALFA-2B, RECOMBINANT, 1 MILLION UNITS	INTRON A (W/DILUENT IN VIAL) 50 Million IU	1 EA	VL	IJ	EA	EA	1 MU		50	01/01/2002	05/28/2016						
00085-0571-02	J9214			01/01/2002	07/31/2016	INJECTION, INTERFERON, ALFA-2B, RECOMBINANT, 1 MILLION UNITS	INTRON A (W/DILUENT IN VIAL) 10 Million IU	1 EA	VL	IJ	EA	EA	1 MU		10	01/01/2002	07/31/2016						
00085-1110-01	J9214			01/01/2002	05/28/2016	INJECTION, INTERFERON, ALFA-2B, RECOMBINANT, 1 MILLION UNITS	INTRON A (W/DILUENT IN VIAL) 18 Million IU	1 EA	VL	IJ	EA	EA	1 MU		18	01/01/2002	05/28/2016						
00085-1133-01	J9214			01/01/2002	99/99/9999	INJECTION, INTERFERON, ALFA-2B, RECOMBINANT, 1 MILLION UNITS	INTRON A (M.D.V.,AF) 10 Million IU/ML	2.5 ML	VL	IJ	ML	ML	1 MU		10	01/01/2002	99/99/9999						
00085-1136-01	J1327			01/01/2002	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	INTEGRILIN (VIAL) 0.75 MG/ML	100 ML	VL	IV	ML	ML	5 MG		0.15	01/01/2002	99/99/9999						
00085-1168-01	J9214			01/01/2002	99/99/9999	INTERFERON, ALFA-2B, RECOMBINANT, 1 MILLION UNITS	INTRON A (M.D.V.,AF) 6 Million IU/ML	3 ML	VL	IJ	ML	ML	1 MU		6	01/01/2002	99/99/9999						
00085-1177-01	J1327			01/01/2002	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	INTEGRILIN (VIAL) 2 MG/ML	10 ML	VL	IV	ML	ML	5 MG		0.4	01/01/2002	99/99/9999						
00085-1177-02	J1327			01/01/2002	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	INTEGRILIN (VIAL) 2 MG/ML	100 ML	VL	IV	ML	ML	5 MG		0.4	01/01/2002	99/99/9999						
00085-1248-03	None			04/09/2007	05/16/2014	TEMODAR, 5 MG, ORAL	TEMODAR 5 MG	14 EA	BO	PO	EA	EA	5 MG		1	04/09/2007	05/16/2014						
00085-1279-01	J3490			01/01/2002	10/28/2015	UNCLASSIFIED DRUGS	PEG-INTRON (VIAL/SRN/DILUENT,PF) 150 MCG	1 EA	BX	MR	EA	EA	1 EA		1	01/01/2002	10/28/2015						
00085-1291-01	J3490			01/01/2002	10/15/2015	UNCLASSIFIED DRUGS	PEG-INTRON (VIAL/SRN/DILUENT,PF) 80 MCG	1 EA	BX	MR	EA	EA	1 EA		1	01/01/2002	10/15/2015						
00085-1297-01	J3490			02/02/2004	03/31/2015	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 120 MCG	1 EA	BX	MR	EA	EA	1 EA		1	02/02/2004	03/31/2015						
00085-1297-02	J3490			03/07/2005	08/31/2016	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 120 MCG	1 EA	BX	MR	EA	EA	1 EA		1	03/07/2005	08/31/2016						
00085-1304-01	J3490			01/01/2002	11/22/2015	UNCLASSIFIED DRUGS	PEG-INTRON (VIAL/SRN/DILUENT,PF) 120 MCG	1 EA	BX	MR	EA	EA	1 EA		1	01/01/2002	11/22/2015						
00085-1316-01	J3490			02/02/2004	03/31/2015	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 80 MCG	1 EA	BX	MR	EA	EA	1 EA		1	02/02/2004	03/31/2015						
00085-1316-02	J3490			03/07/2005	06/30/2015	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 80 MCG	1 EA	BX	MR	EA	EA	1 EA		1	03/07/2005	06/30/2015						
00085-1323-01	J3490			02/02/2004	03/31/2015	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 50 MCG	1 EA	BX	MR	EA	EA	1 EA		1	02/02/2004	03/31/2015						
00085-1323-02	J3490			03/07/2005	04/30/2015	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 50 MCG	1 EA	BX	MR	EA	EA	1 EA		1	03/07/2005	04/30/2015						
00085-1366-01	None			04/09/2007	08/31/2014	TEMODAR, 100 MG, ORAL	TEMODAR 100 MG	14 EA	BO	PO	EA	EA	100 MG		1	04/09/2007	08/31/2014						
00085-1366-02	None			04/09/2007	12/31/2014	TEMODAR, 100 MG, ORAL	TEMODAR 100 MG	5 EA	BO	PO	EA	EA	100 MG		1	04/09/2007	12/31/2014						
00085-1368-01	J3490			01/01/2002	03/06/2016	UNCLASSIFIED DRUGS	PEG-INTRON (VIAL/SRN/DILUENT,PF) 50 MCG	1 EA	BX	MR	EA	EA	1 EA		1	01/01/2002	03/06/2016						
00085-1370-01	J3490			02/02/2004	03/31/2015	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 150 MCG	1 EA	BX	MR	EA	EA	1 EA		1	02/02/2004	03/31/2015						
00085-1370-02	J3490			03/07/2005	07/31/2015	UNCLASSIFIED DRUGS	PEG-INTRON (PF,REDIPEN) 150 MCG	1 EA	BX	MR	EA	EA	1 EA		1	03/07/2005	07/31/2015						
00085-1417-01	None			04/09/2007	12/31/2014	TEMODAR, 250 MG, ORAL	TEMODAR 250 MG	5 EA	BO	PO	EA	EA	250 MG		1	04/09/2007	12/31/2014						
00085-1425-01	None			04/09/2007	08/31/2015	TEMODAR, 20 MG, ORAL	TEMODAR 140 MG	5 EA	BO	PO	EA	EA	20 MG		7	04/09/2007	08/31/2015						
00085-1425-02	None			04/09/2007	08/31/2015	TEMODAR, 20 MG, ORAL	TEMODAR 140 MG	14 EA	BO	PO	EA	EA	20 MG		7	04/09/2007	08/31/2015						
00085-1430-01	None			04/09/2007	08/31/2015	TEMODAR, 20 MG, ORAL	TEMODAR 180 MG	5 EA	BO	PO	EA	EA	20 MG		9	04/09/2007	08/31/2015						
00085-1430-02	None			04/09/2007	11/30/2014	TEMODAR, 20 MG, ORAL	TEMODAR 180 MG	14 EA	BO	PO	EA	EA	20 MG		9	04/09/2007	11/30/2014						
00085-1519-01	None			04/09/2007	07/31/2015	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	14 EA	BO	PO	EA	EA	20 MG		1	04/09/2007	07/31/2015						
00085-1519-02	None			04/09/2007	08/31/2014	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	5 EA	BO	PO	EA	EA	20 MG		1	04/09/2007	08/31/2014						
00085-1737-01	J2280			08/17/2005	03/31/2017	INJECTION, MOXIFLOXACIN, 100 MG	AVELOX I.V. (FLEXIBAG,PF) 400 MG/250 ML	250 ML	FC	IV	ML	ML	100 MG		0.016	08/17/2005	03/31/2017						
00085-3004-01	None			01/30/2008	07/31/2014	TEMODAR, 5 MG, ORAL	TEMODAR 5 MG	14 EA	BO	PO	EA	EA	5 MG		1	01/30/2008	07/31/2014						
00085-3004-02	None			01/30/2008	05/21/2014	TEMODAR, 5 MG, ORAL	TEMODAR 5 MG	5 EA	BO	PO	EA	EA	5 MG		1	01/30/2008	05/21/2014						
00088-1202-05	Q0180			01/01/2002	99/99/9999	DOLASETRON MESYLATE, 100 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	ANZEMET 50 MG	5 EA	BO	PO	EA	EA	100 MG		0.5	01/01/2002	99/99/9999						
00088-1203-05	Q0180			01/01/2002	99/99/9999	DOLASETRON MESYLATE, 100 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	ANZEMET 100 MG	5 EA	BO	PO	EA	EA	100 MG		1	01/01/2002	99/99/9999						
00088-1206-32	J1260			01/01/2002	99/99/9999	INJECTION, DOLASETRON MESYLATE, 10 MG	ANZEMET (S.D.V.) 20 MG/ML	5 ML	VL	IV	ML	ML	10 MG		2	01/01/2002	99/99/9999						
00143-9275-01	J9000			08/10/2018	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 10 MG	1 EA	VL	IV	EA	EA	10 MG		1	08/10/2018	99/99/9999						
00143-9277-01	J9000			08/10/2018	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 10 MG	1 EA	VL	IV	EA	EA	10 MG		1	08/10/2018	99/99/9999						
00378-9690-52	J7614			07/23/2018	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3 ML	VL	IH	ML	ML	0.5 MG		0.20666	07/23/2018	99/99/9999						
00088-1208-06	J1260			12/15/2003	99/99/9999	INJECTION, DOLASETRON MESYLATE, 10 MG	ANZEMET (S.D.V.) 20 MG/ML	0.625 ML	VL	IV	ML	ML	10 MG		2	12/15/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00088-1209-26	J1260			07/21/2003	99/99/9999	INJECTION, DOLASETRON MESYLATE, 10 MG	ANZEMET (M.D.V.) 20 MG/ML	25 ML	VL	IV	ML		10 MG		2	07/21/2003	99/99/9999						
00088-2220-33	J1815			01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	LANTUS 100 U/ML	10 ML	VL	SC	ML		5 U		20	01/01/2003	99/99/9999						
00088-2500-33	J1817			01/24/2006	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	APIDRA 100 U/ML	10 ML	VL	SC	ML		50 U		2	01/24/2006	99/99/9999						
00091-1110-20	J0270			09/02/2004	05/22/2012	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	EDEX (29GX1/2",KIT) 10 MCG	1 EA	BX	MR	EA		1.25 MCG		8	09/02/2004	05/22/2012						
00091-1140-16	J0270			09/02/2004	05/30/2012	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	EDEX (29GX1/2",KIT) 40 MCG	1 EA	BX	MR	EA		1.25 MCG		32	09/02/2004	05/30/2012						
00091-1140-20	J0270			09/02/2004	05/07/2012	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	EDEX (29GX1/2",KIT) 40 MCG	1 EA	BX	MR	EA		1.25 MCG		32	09/02/2004	05/07/2012						
00093-0782-01	J8999			02/20/2003	10/20/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	100 EA	BO	PO	EA		1 EA		1	02/20/2003	10/20/2016						
00093-0782-05	J8999			01/09/2008	10/20/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	500 EA	BO	PO	EA		1 EA		1	01/09/2008	10/20/2016						
00093-0782-10	J8999			01/09/2008	10/20/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	1000 EA	BO	PO	EA		1 EA		1	01/09/2008	10/20/2016						
00093-0782-56	J8999			02/20/2003	07/17/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	30 EA	BO	PO	EA		1 EA		1	02/20/2003	07/17/2016						
00093-0784-05	J8999			01/09/2008	10/20/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 10 MG	500 EA	BO	PO	EA		1 EA		1	01/09/2008	10/20/2016						
00093-0784-06	J8999			02/20/2003	07/17/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 10 MG	60 EA	BO	PO	EA		1 EA		1	02/20/2003	07/17/2016						
00093-0784-10	J8999			01/09/2008	10/20/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 10 MG	1000 EA	BO	PO	EA		1 EA		1	01/09/2008	10/20/2016						
00093-0784-86	J8999			02/20/2003	08/02/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 10 MG	180 EA	BO	PO	EA		1 EA		1	02/20/2003	08/02/2016						
00093-5420-88	J8515			03/07/2007	99/99/9999	CABERGOLINE, ORAL, 0.25 MG	CABERGOLINE 0.5 MG	8 EA	BO	PO	EA		0.25 MG		2	03/07/2007	99/99/9999						
00093-5510-06	J8999			04/27/2005	03/26/2015	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MERCAPTOPYRINE (USP) 50 MG	60 EA	BO	PO	EA		1 EA		1	04/27/2005	03/26/2015						
00093-6118-16	J7510			01/01/2002	08/13/2018	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	480 ML	BO	PO	ML		5 MG		0.6	01/01/2002	08/13/2018						
00093-6118-87	J7510			01/01/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240 ML	BO	PO	ML		5 MG		0.6	01/01/2002	99/99/9999						
00093-6723-73	J7620			01/03/2008	06/04/2018	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	30 ML	VL	IH	ML		3 MG		0.33333	01/03/2008	06/04/2018						
00093-6723-74	J7620			01/03/2008	06/04/2018	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	60 ML	VL	IH	ML		3 MG		0.33333	01/03/2008	06/04/2018						
00093-7146-09	Q0144			12/06/2005	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (3X6,FILM-COATED) 250 MG	18 EA	DP	PO	EA		1 GM		0.25	12/06/2005	01/31/2014						
00093-7146-18	Q0144			11/14/2005	07/01/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	6 EA	DP	PO	EA		1 GM		0.25	11/14/2005	07/01/2016						
00093-7146-56	Q0144			11/14/2005	09/12/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	30 EA	BO	PO	EA		1 GM		0.25	11/14/2005	09/12/2017						
00093-7147-56	Q0144			11/14/2005	06/28/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 600 MG	30 EA	BO	PO	EA		1 GM		0.6	11/14/2005	06/28/2017						
00093-7169-33	Q0144			11/14/2005	01/10/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	3 EA	DP	PO	EA		1 GM		0.5	11/14/2005	01/10/2014						
00093-7169-56	Q0144			11/14/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	30 EA	BO	PO	EA		1 GM		0.5	11/14/2005	99/99/9999						
16714-0777-01	J9025			07/03/2018	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (SDV,PF,LATEX-FREE) 100 MG	1 EA	VL	IJ	EA		1 MG		100	07/03/2018	99/99/9999						
16729-0332-03	J9263			05/01/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		10	05/01/2018	99/99/9999						
00069-1305-10	J0885			05/22/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACIR (PF) 2000 U/1 ML	1 ML	VL	IJ	ML		1000 U		2	05/22/2018	12/31/2018						
00093-7485-12	Q0166			01/02/2008	11/12/2018	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISTERON HYDROCHLORIDE (2X1,FILM COATED) 1 MG	2 EA	BX	PO	EA		1 MG		1	01/02/2008	11/12/2018						
00093-7485-20	Q0166			01/02/2008	11/12/2018	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISTERON HYDROCHLORIDE (5X4,FILM COATED) 1 MG	20 EA	BX	PO	EA		1 MG		1	01/02/2008	11/12/2018						
00093-8940-01	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
00078-0741-81	J2502			08/23/2018	99/99/9999	INJECTION, PASIREOTIDE LONG ACTING, 1 MG	SIGNIFOR LAR (SINGLE USE) 30 MG	1 EA	BX	IM	EA		1 MG		30	08/23/2018	99/99/9999						
00078-0748-81	J2502			08/23/2018	99/99/9999	INJECTION, PASIREOTIDE LONG ACTING, 1 MG	SIGNIFOR LAR (SINGLE USE) 10 MG	1 EA	BX	IM	EA		1 MG		10	08/23/2018	99/99/9999						
00093-8940-05	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	500 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
00093-8940-93	J8499			11/30/2007	11/27/2012	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (USP,HARD GELATIN) 200 MG	100 EA	BX	PO	EA		1 EA		1	11/30/2007	11/27/2012						
00093-8943-01	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
00093-8943-05	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	500 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
00093-8947-01	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
00093-8947-05	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	500 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
00093-9643-01	Q0164			01/01/2002	08/06/2018	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	100 EA	BO	PO	EA		5 MG		1	01/01/2002	08/06/2018						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00093-9652-01		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013						
00113-0379-26		Q0163		01/14/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GOOD SENSE ANTIHISTAMINE ALLERGY RELIEF (ALCOHOL FREE,CHERRY) 12.5 MG/5 ML	118	ML	BO	PO	ML	50	MG	0.05	01/14/2004	99/99/9999						
00113-0431-62		Q0163		01/14/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GOOD SENSE NIGHTTIME SLEEP AID (MINI-CAPLETS) 25 MG	24	EA	NA	PO	EA	50	MG	0.5	01/14/2004	99/99/9999						
00113-0462-62		Q0163		01/14/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GOOD SENSE ANTIHISTAMINE ALLERGY RELIEF (EASY TO SWALLOW) 25 MG	24	EA	BX	PO	EA	50	MG	0.5	01/14/2004	99/99/9999						
00113-0479-62		Q0163		01/14/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GOOD SENSE ANTIHISTAMINE ALLERGY RELIEF (EASY TO SWALLOW) 25 MG	24	EA	BX	PO	EA	50	MG	0.5	01/14/2004	99/99/9999						
00113-0479-78		Q0163		01/14/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GOOD SENSE ANTIHISTAMINE ALLERGY RELIEF (EASY TO SWALLOW) 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/14/2004	99/99/9999						
00115-1040-01		Q0169		02/12/2008	11/01/2012	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 12.5 MG	100	EA	BO	PO	EA	12.5	MG	1	02/12/2008	11/01/2012						
00115-1041-01		Q0170		02/12/2008	09/19/2012	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	100	EA	BO	PO	EA	25	MG	1	02/12/2008	09/19/2012						
00115-1041-03		Q0170		04/01/2008	09/19/2012	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	1000	EA	BO	PO	EA	25	MG	1	04/01/2008	09/19/2012						
00115-1042-01		Q0170		05/20/2008	12/20/2012	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 50 MG	100	EA	BO	PO	EA	25	MG	2	05/20/2008	12/20/2012						
00121-0489-05		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	5	ML	CP	PO	ML	50	MG	0.05	01/01/2002	99/99/9999						
00121-0489-10		Q0163		01/01/2002	06/06/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	10	ML	CP	PO	ML	50	MG	0.05	01/01/2002	06/06/2017						
00121-0759-08		J7510		05/02/2005	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (AF,DYE-FREE,GRAPE) 15 MG/5 ML	237	ML	BO	PO	ML	5	MG	0.6	05/02/2005	99/99/9999						
00121-4776-10		J8999		07/07/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (40X10ML CUPS,APRICOT) 40 MG/ML	10	ML	CP	PO	ML	1	EA	1	07/07/2006	99/99/9999						
00143-1425-01		J7506		12/09/2004	11/27/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 2.5 MG	100	EA	BO	PO	EA	5	MG	0.5	12/09/2004	11/27/2013						
16729-0332-05		J9263		05/01/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	20	ML	VL	IV	ML	0.5	MG	10	05/01/2018	99/99/9999						
00143-1473-01		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
00143-1473-10		J7506		01/01/2002	11/27/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	1000	EA	BO	PO	EA	5	MG	2	01/01/2002	11/27/2013						
00143-1475-01		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
00143-1475-10		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	1000	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
00143-1477-01		J7506		01/01/2002	11/27/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	100	EA	BO	PO	EA	5	MG	4	01/01/2002	11/27/2013						
00143-1477-05		J7506		01/01/2002	11/27/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	500	EA	BO	PO	EA	5	MG	4	01/01/2002	11/27/2013						
00143-1477-10		J7506		01/01/2002	11/27/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	1000	EA	BO	PO	EA	5	MG	4	01/01/2002	11/27/2013						
00143-9891-05		J2405		04/03/2007	04/10/2012	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (USP,SINGLE DOSE) 2 MG/ML	2	ML	VL	IJ	ML	1	MG	2	04/03/2007	04/10/2012						
00169-1833-11		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN R (VIAL) 100 U/ML	10	ML	VL	IJ	ML	5	U	20	01/01/2003	99/99/9999						
00169-1834-11		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN N (VIAL) 100 U/ML	10	ML	VL	SC	ML	5	U	20	01/01/2003	99/99/9999						
16729-0419-03		J9201		01/15/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 100 MG/1 ML	10	ML	VL	IV	ML	200	MG	0.5	01/15/2018	99/99/9999						
25021-0179-15		J0878		06/15/2018	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (SDV,PF,LATEX-FREE) 350 MG	1	EA	VL	IV	EA	1	MG	350	06/15/2018	99/99/9999						
00169-1837-11		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN 70/30 (VIAL) 70 U/ML-30 U/ML	10	ML	VL	SC	ML	5	U	20	01/01/2003	99/99/9999						
00169-3303-12		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLOG (PENFILL CARTRIDGE) 100 U/ML	3	ML	CT	SC	ML	5	U	20	01/01/2003	99/99/9999						
00169-3685-12		J1815		02/10/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLOG MIX 70/30 (VIAL) 70 U/ML-30 U/ML	10	ML	VL	SC	ML	5	U	20	02/10/2003	99/99/9999						
00169-3696-19		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLOG MIX 70/30 (FLEXPEN SRN PREFILLED) 70 U/ML-30 U/ML	3	ML	SR	SC	ML	5	U	20	01/01/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00169-6339-10		J1815		02/10/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLOG FLEXPEN (PREFILLED SYRINGE) 100 U/ML	3 ML	SR	SC	ML		5 U		20	02/10/2003	99/99/9999						
00169-7065-15		J1610		06/01/2005	99/99/9999	INJECTION, GLUCAGON HYDROCHLORIDE, PER 1 MG	GLUCAGON HYPKOKIT 1 MG	1 EA	BX	IJ	EA		1 MG		1	06/01/2005	99/99/9999						
00169-7501-11		J1817		01/01/2003	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	NOVOLOG (VIAL) 100 U/ML	10 ML	VL	SC	ML		50 U		2	01/01/2003	99/99/9999						
00172-3753-96		J9265		01/24/2002	12/31/2014	INJECTION, PACLITAXEL, 30 MG	NOV-ONXOL (M.D.V.) 6 MG/ML	50 ML	VL	IV	ML		30 MG		0.2	01/24/2002	12/31/2014						
00172-3754-94		J9265		01/24/2002	12/31/2014	INJECTION, PACLITAXEL, 30 MG	NOV-ONXOL (M.D.V.) 6 MG/ML	5 ML	VL	IV	ML		30 MG		0.2	01/24/2002	12/31/2014						
25021-0179-16		J0878		06/15/2018	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (SDV,PF,LATEX-FREE) 350 MG	10 EA	VL	IV	EA		1 MG		350	06/15/2018	99/99/9999						
25021-0675-10		J2800		06/04/2018	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (LATEX-FREE) 100 MG/1 ML	10 ML	VL	IJ	ML		10 ML		0.1	06/04/2018	99/99/9999						
52609-4504-06		J0895		05/23/2018	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE 2 GM	4 EA	VL	IJ	EA		500 MG		4	05/23/2018	99/99/9999						
00172-3756-95		J9265		01/24/2002	12/31/2014	INJECTION, PACLITAXEL, 30 MG	NOV-ONXOL (M.D.V.) 6 MG/ML	25 ML	VL	IV	ML		30 MG		0.2	01/24/2002	12/31/2014						
00172-4960-58		J8999		01/01/2002	12/31/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	FLUTAMIDE 125 MG	180 EA	BO	PO	EA		1 EA		1	01/01/2002	12/31/2016						
00172-4960-70		J8999		01/01/2002	12/31/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	FLUTAMIDE 125 MG	500 EA	BO	PO	EA		1 EA		1	01/01/2002	12/31/2016						
00172-6406-49		J7631		01/01/2002	99/99/9999	CROMOLYN SODIUM, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (VIAL) 10 MG/ML	2 ML	PC	IH	ML		10 MG		1	01/01/2002	99/99/9999						
00172-6406-49	KO	J7631	KO	01/01/2002	99/99/9999	CROMOLYN SODIUM, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (VIAL) 10 MG/ML	2 ML	PC	IH	ML		10 MG		1	01/01/2002	99/99/9999						
00172-6406-59		J7631		01/01/2002	10/05/2016	CROMOLYN SODIUM, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (VIAL) 10 MG/ML	2 ML	PC	IH	ML		10 MG		1	01/01/2002	10/05/2016						
00172-6406-59	KO	J7631	KO	01/01/2002	10/05/2016	CROMOLYN SODIUM, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (VIAL) 10 MG/ML	2 ML	PC	IH	ML		10 MG		1	01/01/2002	10/05/2016						
00172-7310-46		J7515		04/14/2005	05/02/2017	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE (USP,MODIFIED,SOFTGEL) 25 MG	30 EA	BX	PO	EA		25 MG		1	04/14/2005	05/02/2017						
00172-7311-46		J7515		04/14/2005	11/03/2015	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE (USP,MODIFIED,SOFTGEL) 50 MG	30 EA	BX	PO	EA		25 MG		2	04/14/2005	11/03/2015						
00172-7312-46		J7502		04/14/2005	05/02/2017	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (USP,MODIFIED,SOFTGEL) 100 MG	30 EA	BX	PO	EA		100 MG		1	04/14/2005	05/02/2017						
00172-7313-20		J7502		04/14/2005	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (USP,MODIFIED) 100 MG/ML	50 ML	BO	PO	ML		100 MG		1	04/14/2005	99/99/9999						
00173-0260-10		J1160		01/01/2002	09/29/2013	INJECTION, DIGOXIN, UP TO 0.5 MG	LANOXIN (AMP) 0.25 MG/ML	2 ML	AM	IV	ML		0.5 MG		0.5	01/01/2002	09/29/2013						
00173-0262-10		J1160		01/01/2002	04/22/2013	INJECTION, DIGOXIN, UP TO 0.5 MG	LANOXIN PEDIATRIC (AMP) 0.1 MG/ML	1 ML	AM	IV	ML		0.5 MG		0.2	01/01/2002	04/22/2013						
00173-0352-10		J0697		02/01/2005	08/26/2013	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	ZINACEF 750 MG	1 EA	VL	IJ	EA		750 MG		1	02/01/2005	08/26/2013						
00173-0354-10		J0697		02/01/2005	08/26/2013	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	ZINACEF 1.5 GM	1 EA	VL	IJ	EA		750 MG		2	02/01/2005	08/26/2013						
54879-0021-01		None		05/08/2018	99/99/9999	CYCLOPHOSPHAMIDE, 25 MG, ORAL	ZINACEF (PREMIX) 750 MG/50 ML	100 EA	BO	PO	EA		25 MG		1	05/08/2018	99/99/9999						
00173-0362-38		J2780		01/01/2002	11/30/2014	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC (VIAL) 25 MG/ML	2 ML	VL	IJ	ML		25 MG		1	01/01/2002	11/30/2014						
00173-0363-00		J2780		01/01/2002	12/09/2013	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC (VIAL) 25 MG	40 ML	VL	IJ	ML		25 MG		1	01/01/2002	12/09/2013						
00173-0363-01		J2780		01/01/2002	12/11/2013	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC (M.D.V.) 25 MG/ML	6 ML	VL	IJ	ML		25 MG		1	01/01/2002	12/11/2013						
00173-0377-10		J0713		02/01/2005	12/16/2012	INJECTION, CEFTAZIDIME, PER 500 MG	FORTAZ 500 MG	1 EA	VL	IJ	EA		500 MG		1	02/01/2005	12/16/2012						
00173-0378-10		J0713		02/01/2005	09/18/2013	INJECTION, CEFTAZIDIME, PER 500 MG	FORTAZ 1 GM	1 EA	VL	IJ	EA		500 MG		2	02/01/2005	09/18/2013						
00173-0379-34		J0713		01/01/2002	08/05/2013	INJECTION, CEFTAZIDIME, PER 500 MG	FORTAZ (VIAL) 2 GM	1 EA	VL	IJ	EA		500 MG		4	01/01/2002	08/05/2013						
00173-0382-37		J0713		01/01/2002	06/18/2013	INJECTION, CEFTAZIDIME, PER 500 MG	FORTAZ (BULK VIAL) 6 GM	1 EA	VL	IJ	EA		500 MG		12	01/01/2002	06/18/2013						
00173-0400-00		J0697		01/01/2002	04/04/2013	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	ZINACEF 7.5 GM	1 EA	VL	IJ	EA		750 MG		10	01/01/2002	04/04/2013						
00173-0424-00		J0697		01/01/2002	06/28/2013	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	ZINACEF (PREMIX) 750 MG/50 ML	50 ML	PC	IV	ML		750 MG		0.02	01/01/2002	06/28/2013						
00173-0425-00		J0697		01/01/2002	12/12/2013	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	ZINACEF (PREMIX) 1.5 GM/50 ML	50 ML	PC	IV	ML		750 MG		0.04	01/01/2002	12/12/2013						
00173-0434-00		J0713		01/01/2002	12/01/2013	INJECTION, CEFTAZIDIME, PER 500 MG	FORTAZ (ADD-VANTAGE) 1 GM	1 EA	VL	IJ	EA		500 MG		2	01/01/2002	12/01/2013						
00173-0435-00		J0713		01/01/2002	12/01/2013	INJECTION, CEFTAZIDIME, PER 500 MG	FORTAZ (ADD-VANTAGE) 2 GM	1 EA	VL	IJ	EA		500 MG		4	01/01/2002	12/01/2013						
00173-0436-00		J0697		01/01/2002	12/29/2013	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	ZINACEF (ADD-VANTAGE) 750 MG	1 EA	VL	IJ	EA		750 MG		1	01/01/2002	12/29/2013						
00173-0437-00		J0697		01/01/2002	05/02/2013	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	ZINACEF (ADD-VANTAGE) 1.5 GM	1 EA	VL	IJ	EA		750 MG		2	01/01/2002	05/02/2013						
00173-0441-00		J2780		01/01/2002	06/14/2013	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC (PREMIX) 1 MG/ML	50 ML	FC	IV	ML		25 MG		0.04	01/01/2002	06/14/2013						
00173-0442-00		J2405		01/01/2002	05/07/2018	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ZOFRAN (M.D.V.) 2 MG/ML	20 ML	VL	IJ	ML		1 MG		2	01/01/2002	05/07/2018						
00378-9690-52	KO	J7614	KO	07/23/2018	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.20666	07/23/2018	99/99/9999						
00078-0755-61		J2502		08/23/2018	99/99/9999	INJECTION, PASIREOTIDE LONG ACTING, 1 MG	SIGNIFOR LAR (6ML VIAL) 10 MG	1 EA	VL	IM	EA		1 MG		10	08/23/2018	99/99/9999						
00078-0769-61		J2502		08/23/2018	99/99/9999	INJECTION, PASIREOTIDE LONG ACTING, 1 MG	SIGNIFOR LAR (6ML VIAL) 30 MG	1 EA	VL	IM	EA		1 MG		30	08/23/2018	99/99/9999						
00173-0449-02		J3030		01/01/2002	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX (S.D.V.) 6 MG/0.5 ML	0.5 ML	VL	SC	ML		6 MG		2	01/01/2002	99/99/9999						
00173-0635-35		J8999		01/01/2002	06/21/2012	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN 2 MG	50 EA	BO	PO	EA		1 EA		1	01/01/2002	06/21/2012						
00173-0713-25		None		01/01/2000	06/21/2012	BUSULFAN, 2 MG, ORAL	MYLERAN 2 MG	25 EA	BO	PO	EA		2 MG		1	01/01/2000	06/21/2012						
00173-0739-00		J3030		03/17/2006	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX STATDOSE 4 MG/0.5 ML	1 EA	BX	SC	EA		6 MG		0.66666	03/17/2006	99/99/9999						
00173-0739-02		J3030		03/17/2006	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX STATDOSE (REFILL W/2 SYRINGES) 4 MG/0.5 ML	1 EA	BX	SC	EA		6 MG		0.66666	03/17/2006	99/99/9999						
00173-0945-55		J8499		01/01/2002	01/08/2017	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	01/08/2017						
00173-0949-55		J8499		01/01/2002	06/08/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 400 MG	100 EA	BO														

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00182-1131-93		Q0163		05/03/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	NIGHT-TIME SLEEP AID (MAX. STR.,SOFTGEL) 50 MG	32	EA	BO	PO	EA	50	MG	1	05/03/2002	02/03/2016						
00069-1306-10		J0885		05/22/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 3000 U/1 ML	1	ML	VL	IJ	ML	1000	U	3	05/22/2018	12/31/2018						
00069-1307-10		J0885		05/22/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 4000 U/1 ML	1	ML	VL	IJ	ML	1000	U	4	05/22/2018	12/31/2018						
63323-0721-10		J9044		01/01/2019	99/99/9999	INJECTION, BORTEZOMIB, NOT OTHERWISE SPECIFIED, 0.1 MG	BORTEZOMIB, (SDV,LATEX-FREE) 3.5 MG	1	EA	VL	IV	EA	0.1	MG	35	01/01/2019	99/99/9999						
00003-2187-13		J0129		11/05/2018	99/99/9999	INJECTION, ABATACEPT, 10 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ORENCIA (W/SYRINGE,PF) 250 MG	1	EA	VL	IV	EA	10	MG	25	11/05/2018	99/99/9999						
00185-0613-01		Q0177		01/01/2002	07/29/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	07/29/2014						
00185-0613-05		Q0177		01/01/2002	07/29/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	500	EA	BO	PO	EA	25	MG	1	01/01/2002	07/29/2014						
00185-0615-01		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013						
00185-0615-05		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013						
00185-0648-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
00185-0648-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	1000	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
00185-0649-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
00185-0649-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	1000	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
00185-0932-30		J7515		01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE (SOFTGEL) 25 MG	30	EA	BO	PO	EA	25	MG	1	01/01/2002	99/99/9999						
00185-0933-30		J7502		01/01/2002	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (SOFTGEL) 100 MG	30	EA	BO	PO	EA	100	MG	1	01/01/2002	99/99/9999						
00185-1125-05		J8999		01/01/2002	04/18/2012	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	FLUTAMIDE 125 MG	500	EA	BO	PO	EA	1	EA	1	01/01/2002	04/18/2012						
00185-1125-18		J8999		01/01/2002	04/18/2012	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	FLUTAMIDE 125 MG	180	EA	BO	PO	EA	1	EA	1	01/01/2002	04/18/2012						
00185-1125-88		J8999		01/01/2002	04/18/2012	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	FLUTAMIDE (BLISTER PACK,10X10) 125 MG	100	EA	BX	PO	EA	1	EA	1	01/01/2002	04/18/2012						
00185-7203-70		Q0144		09/21/2006	11/13/2012	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (USP,CHERRY) 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	09/21/2006	11/13/2012						
00185-7206-70		Q0144		09/21/2006	11/13/2012	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (USP,CHERRY) 200 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.04	09/21/2006	11/13/2012						
00185-7322-30		J7620		07/01/2007	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	30	ML	PC	IH	ML	3	MG	0.33333	07/01/2007	99/99/9999						
00185-7322-60		J7620		07/01/2007	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	60	ML	PC	IH	ML	3	MG	0.33333	07/01/2007	99/99/9999						
00186-0418-01		J0670		01/01/2002	08/31/2012	INJECTION, MEPIVACAINE HYDROCHLORIDE, PER 10 ML	POLOCAINE-MPF (S.D.V.) 1.5%	30	ML	VL	IJ	ML	10	ML	0.1	01/01/2002	08/31/2012						
00186-0859-81		J2795		01/01/2002	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	NAROPIN (S.D. INFUSION BOTTLE) 2 MG/ML	100	ML	VL	IJ	ML	1	MG	2	01/01/2002	99/99/9999						
00186-1033-91		J3490		01/01/2002	03/31/2013	UNCLASSIFIED DRUGS	SENSORCAINE-MPF (S.D.V.,STERILE-PAK) 0.5%	30	ML	VL	IJ	ML	1	EA	1	01/01/2002	03/31/2013						
00186-1988-04		J7626		01/01/2002	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES (5X6) 0.25 MG/2 ML	2	ML	PC	IH	ML	0.25	MG	0.5	01/01/2002	99/99/9999						
00186-1988-04	KO	J7626	KO	01/01/2002	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES (5X6) 0.25 MG/2 ML	2	ML	PC	IH	ML	0.25	MG	0.5	01/01/2002	99/99/9999						
00186-1989-04		J7626		01/01/2002	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES (5X6) 0.5 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	0.5	01/01/2002	99/99/9999						
00186-1989-04	KO	J7626	KO	01/01/2002	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES (5X6) 0.5 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	0.5	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00186-1990-04		J7626		08/27/2007	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES (30X2ML) 1 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	1	08/27/2007	99/99/9999							
00186-1990-04	KO	J7626	KO	08/27/2007	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES (30X2ML) 1 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	1	08/27/2007	99/99/9999							
00206-8852-16		J2543		04/05/2006	99/99/9999	INJECTION, PIPERACILLIN SODIUM/AZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	ZOSYN 2 GM-0.25 GM ZOSYN (SDV, 10X50ML) 3 GM/50 ML-0.375 GM/50 ML	1	EA	VL	IV	EA	1	GM	2	04/05/2006	99/99/9999							
00206-8854-16		J2543		03/06/2006	99/99/9999	INJECTION, PIPERACILLIN SODIUM/AZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	ZOSYN (SDV, 10X100ML) 4 GM/100 ML-0.5 GM/100 ML	1	EA	VL	IV	EA	1	GM	3	03/06/2006	99/99/9999							
00206-8855-16		J2543		03/13/2006	99/99/9999	INJECTION, PIPERACILLIN SODIUM/AZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	ZOSYN (PHARMACY BULK VIAL) 36 GM-4.5 GM	1	EA	VL	IV	EA	1	GM	4	03/13/2006	99/99/9999							
00206-8859-10		J2543		04/28/2006	99/99/9999	INJECTION, PIPERACILLIN SODIUM/AZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	ZOSYN (24 PRE-MIX BAGS OF 50ML) 2 GM/50 ML-0.25 GM/50 ML	1	EA	VL	IV	EA	1	GM	36	04/28/2006	99/99/9999							
00206-8860-02		J2543		01/09/2006	99/99/9999	INJECTION, PIPERACILLIN SODIUM/AZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	ZOSYN (24 PRE-MIX BAGS OF 50ML) 3 GM/50 ML-0.375 GM/50 ML	50	ML	PC	IV	ML	1	GM	0.04	01/09/2006	99/99/9999							
00206-8861-02		J2543		01/09/2006	99/99/9999	INJECTION, PIPERACILLIN SODIUM/AZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	ZOSYN (24 PRE-MIX BAGS OF 50ML) 3 GM/50 ML-0.375 GM/50 ML	50	ML	PC	IV	ML	1	GM	0.06	01/09/2006	99/99/9999							
00206-8862-02		J2543		01/09/2006	99/99/9999	INJECTION, PIPERACILLIN SODIUM/AZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	ZOSYN 4 GM/100 ML-0.5 GM/100 ML	100	ML	PC	IV	ML	1	GM	0.04	01/09/2006	99/99/9999							
00223-8496-02	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (AMP) 0.9%	5	ML	AM	IV	ML	10	ML	0.1	01/01/2007	02/03/2016							
00223-8496-05	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (AMP) 0.9%	5	ML	AM	IV	ML	10	ML	0.1	01/01/2007	02/03/2016							
00223-8497-10	A4216			01/01/2004	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (AMP) 0.9%	10	ML	AM	IV	ML	10	ML	0.1	01/01/2004	02/03/2016							
00223-8500-30	A4216			01/01/2004	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (VIAL) 0.9%	30	ML	VL	IV	ML	10	ML	0.1	01/01/2004	02/03/2016							
00264-1101-55	J7060			01/01/2002	12/31/2014	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (GLASS) 5%	500	ML	FC	IV	ML	500	ML	0.002	01/01/2002	12/31/2014							
00264-1102-55	J7060			01/01/2002	12/31/2014	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (GLASS W/SS, 250 ML) 5%	150	ML	GC	IV	ML	500	ML	0.002	01/01/2002	12/31/2014							
00264-1240-55	J7799			01/01/2002	11/30/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE HYPERTONIC (GLASS W/SS, 1000 ML) 30%	500	ML	GC	IV	ML	1	EA	1	01/01/2002	11/30/2014							
00264-1280-50	J7799			01/01/2002	12/31/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE HYPERTONIC (GLASS W/SOLID STOPPER) 50%	500	ML	GC	IV	ML	1	EA	1	01/01/2002	12/31/2014							
00264-1280-55	J7799			01/01/2002	09/30/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE HYPERTONIC (GLASS W/SOLID STOPPER) 50%	1000	ML	GC	IV	ML	1	EA	1	01/01/2002	09/30/2014							
00264-1290-50	J7799			01/01/2002	05/31/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE HYPERTONIC (GLASS W/SS, 1000 ML) 70%	500	ML	GC	IV	ML	1	EA	1	01/01/2002	05/31/2014							
00264-1482-55	J1265			01/01/2006	08/31/2012	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL (GLASS W/SOLID STOPPER) 5%-160 MG/100 ML	250	ML	GC	IV	ML	40	MG	0.04	01/01/2006	08/31/2012							
00264-1510-31	J7060			01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (100 ML PAB) 5%	50	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00264-1510-32	J7060			01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (150 ML PAB) 5%	100	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00264-1800-31	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (100 ML PAB) 0.9%	50	ML	FC	IV	ML	10	ML	0.1	01/01/2004	99/99/9999							
00264-1800-32	J7050			01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (150 ML PAB) 0.9%	100	ML	FC	IV	ML	250	ML	0.004	01/01/2002	99/99/9999							
00264-1800-36	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (100 ML PAB) 0.9%	25	ML	FC	IV	ML	10	ML	0.1	01/01/2004	99/99/9999							
00264-1940-20	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (CONCENTRATE) 2 MEQ/ML	250	ML	GC	IV	ML	2	MEQ	1	01/01/2002	99/99/9999							
00264-2101-00	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (PIC CONTAINER)	1000	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-2101-10	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (PIC CONTAINER)	500	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-2101-50	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (PIC CONTAINER)	2000	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00069-0291-01	Q5110			09/05/2018	99/99/9999	INJECTION, FILGRASTIM-AAFI, BIOSIMILAR, (NIVESTYM), 1 MICROGRAM	NIVESTYM (PF,LATEX-FREE) 300 MCG/0.5 ML	0.5	ML	SR	IJ	ML	1	MCG	600	09/05/2018	99/99/9999							
00264-2101-70	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (PIC CONTAINER)	4000	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-2201-00	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (PIC CONTAINER) 0.9%	1000	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-2201-10	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (PIC CONTAINER) 0.9%	500	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-2201-50	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (PIC CONTAINER) 0.9%	2000	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-2201-70	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (PIC CONTAINER) 0.9%	4000	ML	PC	IR	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-2303-50	J7799			01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	RESECTISOL 5%	2000	ML	PC	IL	ML	1	EA	1	01/01/2002	99/99/9999							
00264-3103-11	J0690			03/05/2003	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (DUPLEX) 1 GM/50 ML-4%	50	ML	FC	IV	ML	500	MG	0.04	03/05/2003	99/99/9999							
00264-3112-11	J0697			09/15/2003	03/31/2014	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	CEFUROXIME SODIUM 750 MG/50 ML	50	ML	FC	IV	ML	750	MG	0.02	09/15/2003	03/31/2014							
00264-3114-11	J0697			03/01/2004	09/30/2014	INJECTION, STERILE CEFUROXIME SODIUM, PER 750 MG	CEFUROXIME SODIUM (DUPLEX) 1.5 GM/50 ML	50	ML	FC	IV	ML	750	MG	0.04	03/01/2004	09/30/2014							
00264-3123-11	J0694			07/01/2006	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN 1 GM	1	EA	FC	IV	EA	1	GM	1	07/01/2006	99/99/9999							
00264-3125-11	J0694			07/01/2006	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN 2 GM	1	EA	FC	IV	EA	1	GM	2	07/01/2006	99/99/9999							
00264-3153-11	J0696			07/20/2005	99/99/9999	INJECTION, CEFTRIAOXONE SODIUM, PER 250 MG	CEFTRIAOXONE/DEXTROSE 1 GM/50 ML	50	ML	FC	IV	ML	250	MG	0.08	07/20/2005	99/99/9999							
00264-3155-11	J0696			07/20/2005	99/99/9999	INJECTION, CEFTRIAOXONE SODIUM, PER 250 MG	CEFTRIAOXONE/DEXTROSE 2 GM/50 ML	50	ML	FC	IV	ML	250	MG	0.16	07/20/2005	99/99/9999							
00264-4000-55	J7030			01/01/2002	06/30/2015	INFUSION, NORMAL SALINE SOLUTION , 1000 CC	SODIUM CHLORIDE (GLASS CONTAINER) 0.9%	1000	ML	GC	IV	ML	1000	ML	0.001	01/01/2002	06/30/2015							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00264-4001-55		J7040		01/01/2002	09/30/2015	INFUSION, NORMAL SALINE SOLUTION, STERILE (500 ML=1 UNIT)	SODIUM CHLORIDE (GLASS CONTAINER) 0.9%	500	ML	GC	IV	ML	500	ML	0.002	01/01/2002	09/30/2015							
00264-4002-55		J7050		01/01/2002	11/30/2013	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (250 ML GLASS CONTAINER) 0.9%	250	ML	GC	IV	ML	250	ML	0.004	01/01/2002	11/30/2013							
00264-4021-55		J7799		01/01/2002	09/30/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (GLASS CONTAINER) 0.45%	500	ML	GC	IV	ML	1	EA	1	01/01/2002	09/30/2015							
00264-5535-32		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (150 ML PAB CONTAINER) 500 MG/100 ML	100	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-5808-32		J1580		01/01/2002	03/31/2013	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (150 ML PAB CONTAINER) 80 MG/100 ML-0.9%	100	ML	FC	IV	ML	80	MG	0.01	01/01/2002	03/31/2013							
00264-5810-32		J1580		01/01/2002	04/30/2013	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (150 ML PAB CONTAINER) 100 MG/100 ML-0.9%	100	ML	FC	IV	ML	80	MG	0.0125	01/01/2002	04/30/2013							
00264-5812-38		J1580		01/01/2002	08/31/2012	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (100 ML PAB CONTAINER) 1.2 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.015	01/01/2002	08/31/2012							
00264-5816-38		J1580		01/01/2002	04/30/2013	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (100 ML PAB CONTAINER) 1.6 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.02	01/01/2002	04/30/2013							
00264-7510-00		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (EXCEL) 5%	1000	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00264-7510-10		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (EXCEL) 5%	500	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00264-7510-20		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (EXCEL) 5%	250	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00264-7520-00		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (EXCEL) 10%	1000	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7520-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (EXCEL) 10%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7578-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	MANNITOL (EXCEL) 20%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7605-00		J7799		01/01/2002	04/30/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 2.5%-0.45%	1000	ML	FC	IV	ML	1	EA	1	01/01/2002	04/30/2017							
00264-7605-10		J7799		01/01/2002	02/28/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 2.5%-0.45%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	02/28/2014							
00264-7610-00		J7042		01/01/2002	99/99/9999	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.9%	1000	ML	FC	IV	ML	5	%	0.002	01/01/2002	99/99/9999							
00264-7610-10		J7042		01/01/2002	99/99/9999	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.9%	500	ML	FC	IV	ML	5	%	0.002	01/01/2002	99/99/9999							
00264-7610-20		J7042		01/01/2002	07/31/2014	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.9%	250	ML	FC	IV	ML	5	%	0.002	01/01/2002	07/31/2014							
00264-7612-00		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.45%	1000	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7612-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.45%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7614-00		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.33%	1000	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7614-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.33%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7616-00		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.2%	1000	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7616-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.2%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7616-20		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.2%	250	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7622-00		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 10%-0.45%	1000	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7623-20		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 10%-0.2%	250	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7750-00		J7120		01/01/2002	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (EXCEL)	1000	ML	FC	IV	ML	1000	ML	0.001	01/01/2002	99/99/9999							
00264-7750-10		J7120		01/01/2002	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (EXCEL)	500	ML	FC	IV	ML	1000	ML	0.001	01/01/2002	99/99/9999							
00264-7750-20		J7120		01/01/2002	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (EXCEL)	250	ML	FC	IV	ML	1000	ML	0.001	01/01/2002	99/99/9999							
00264-7751-00		J7120		01/01/2002	12/31/2015	RINGERS LACTATE INFUSION, UP TO 1000 CC	DEXTROSE 5%/LACTATED RINGERS (EXCEL)	1000	ML	FC	IV	ML	1000	ML	0.0005	01/01/2002	12/31/2015							
00264-7751-10		J7120		01/01/2002	12/31/2015	RINGERS LACTATE INFUSION, UP TO 1000 CC	DEXTROSE 5%/LACTATED RINGERS (EXCEL)	500	ML	FC	IV	ML	1000	ML	0.0005	01/01/2002	12/31/2015							
00264-7800-00		J7030		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 1000 CC	SODIUM CHLORIDE (EXCEL) 0.9%	1000	ML	FC	IV	ML	1000	ML	0.001	01/01/2002	99/99/9999							
00264-7800-10		J7040		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION, STERILE (500 ML=1 UNIT)	SODIUM CHLORIDE (EXCEL) 0.9%	500	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00264-7800-20		J7050		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (EXCEL) 0.9%	250	ML	FC	IV	ML	250	ML	0.004	01/01/2002	99/99/9999							
00264-7802-00		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (EXCEL) 0.45%	1000	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7802-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (EXCEL) 0.45%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7805-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (HYPERTONIC, EXCEL) 3%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7806-10		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (HYPERTONIC, EXCEL) 5%	500	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00264-7850-00		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION (EXCEL)	1000	ML	FC	IV	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-7850-10		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION (EXCEL)	500	ML	FC	IV	ML	500	ML	0.002	01/01/2004	99/99/9999							
00264-7850-20		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION (EXCEL)	250	ML	FC	IV	ML	500	ML	0.002	01/01/2004	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00264-7865-00		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE/SODIUM CHLORIDE (EXCEL) 2 MEQ/100 ML-0.9% WATER FOR INJECTION (GLASS W/SOLID STOPPER)	1000 ML	FC	IV	ML		2 MEQ		0.01	01/01/2002	99/99/9999							
00264-9200-55		A4217		01/01/2004	08/31/2013	STERILE WATER/SALINE, 500 ML	STERILE WATER/SALINE, 500 ML	1000 ML	GC	IV	ML		500 ML		0.002	01/01/2004	08/31/2013							
00264-9201-55		A4217		01/01/2004	09/30/2013	STERILE WATER/SALINE, 500 ML	STERILE WATER/SALINE, 500 ML	500 ML	GC	IV	ML		500 ML		0.002	01/01/2004	09/30/2013							
00264-9554-10		J2810		01/01/2002	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	DEXTROSE/THEOPHYLLINE (EXCEL) 5%-80 MG/100 ML	500 ML	FC	IV	ML		40 MG		0.02	01/01/2002	99/99/9999							
00264-9558-10		J2810		01/01/2002	09/30/2013	INJECTION, THEOPHYLLINE, PER 40 MG	DEXTROSE/THEOPHYLLINE (EXCEL) 5%-160 MG/100 ML	500 ML	FC	IV	ML		40 MG		0.04	01/01/2002	09/30/2013							
00264-9567-10		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	DEXTROSE/HEPARIN SODIUM (EXCEL) 5%-4000 U/100 ML	500 ML	FC	IV	ML		1000 U		0.04	01/01/2002	99/99/9999							
00264-9577-10		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	DEXTROSE/HEPARIN SODIUM (EXCEL) 5%-5000 U/100 ML	500 ML	FC	IV	ML		1000 U		0.05	01/01/2002	99/99/9999							
00264-9587-20		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	DEXTROSE/HEPARIN SODIUM (EXCEL) 5%-10000 U/100 ML	250 ML	FC	IV	ML		1000 U		0.1	01/01/2002	99/99/9999							
00264-9594-10		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTROSE/LIDOCAINE HCL (EXCEL) 5%-0.4%	500 ML	FC	IV	ML		10 MG		0.4	01/01/2004	99/99/9999							
00264-9594-20		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTROSE/LIDOCAINE HCL (EXCEL) 5%-0.4%	250 ML	FC	IV	ML		10 MG		0.4	01/01/2004	99/99/9999							
00264-9598-20		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTROSE/LIDOCAINE HCL (EXCEL) 5%-0.8%	250 ML	FC	IV	ML		10 MG		0.8	01/01/2004	99/99/9999							
00264-9872-10		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE 200 U/100 ML-0.9%	500 ML	FC	IV	ML		1000 U		0.002	01/01/2002	99/99/9999							
00270-0556-15		J2805		01/01/2006	99/99/9999	INJECTION, SINGALIDE, 5 MICROGRAMS	KINEVAC (VIAL) 5 MCG	1 EA	VL	IV	EA		5 MCG		1	01/01/2006	99/99/9999							
00310-0201-30		J8999		01/01/2002	07/01/2018	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	ARIMIDEX 1 MG	30 EA	BO	PO	EA		1 EA		1	08/07/2008	07/01/2018	01/01/2002	06/02/2008	1				
00310-0321-30		J2185		01/01/2004	99/99/9999	INJECTION, MEROPENEM, 100 MG	MERREM IV (VIAL) 1 GM	1 EA	VL	IV	EA		100 MG		10	01/01/2004	99/99/9999							
00310-0321-65		J2185		07/17/2006	10/22/2012	INJECTION, MEROPENEM, 100 MG	NOVAPLUS MERREM 1 GM	1 EA	VL	IV	EA		100 MG		10	07/17/2006	10/22/2012							
00310-0325-20		J2185		01/01/2004	99/99/9999	INJECTION, MEROPENEM, 100 MG	MERREM IV (VIAL) 500 MG	1 EA	VL	IV	EA		100 MG		5	01/01/2004	99/99/9999							
00310-0325-64		J2185		07/17/2006	10/22/2012	INJECTION, MEROPENEM, 100 MG	NOVAPLUS MERREM 500 MG	1 EA	VL	IV	EA		100 MG		5	07/17/2006	10/22/2012							
00310-0950-36		J9202		05/05/2003	04/05/2018	GOSERELIN ACETATE IMPLANT, PER 3.6 MG	ZOLADEX (SAFESYSTEM SRN) 3.6 MG	1 EA	SR	SC	EA		3.6 MG		1	05/05/2003	04/05/2018							
00310-0951-30		J9202		05/05/2003	02/01/2018	GOSERELIN ACETATE IMPLANT, PER 3.6 MG	ZOLADEX (SAFESYSTEM SRN) 10.8 MG	1 EA	SR	SC	EA		3.6 MG		3	05/05/2003	02/01/2018							
00338-0003-44		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	1000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0003-46		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	2000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0003-47		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	3000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0003-49		A4217		01/01/2004	05/31/2012	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	5000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	05/31/2012							
00338-0004-02		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	250 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0004-03		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	500 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0004-04		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	1000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0004-05		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	1500 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0013-04		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION	1000 ML	FC	IV	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0013-06		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION	2000 ML	FC	IV	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0013-08		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION	3000 ML	FC	IV	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0013-29		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION	5000 ML	FC	IV	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0017-01		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE 5%	150 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-02		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE 5%	250 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-03		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE 5%	500 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-04		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE 5%	1000 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-10		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (QUAD PACK, MINI-BAG) 5%	25 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-11		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (QUAD PACK, MINI-BAG) 5%	50 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-18		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (QUAD PACK, MINI-BAG) 5%	100 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-31		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (MULTI PACK, MINI-BAG) 5%	50 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-38		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (MULTI PACK, MINI-BAG) 5%	100 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-41		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (SINGLE PACK MINI-BAG) 5%	50 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0017-48		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (SINGLE PACK MINI-BAG) 5%	100 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0023-02		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE 10%	250 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0023-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE 10%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0023-04		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE 10%	1000 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0043-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE 0.45%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0043-04		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE 0.45%	1000 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0047-24		A4217		01/01/2004	12/31/2012	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (ARTHROMATIC P.C.) 0.9%	1000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	12/31							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00338-0047-29		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE 0.9% SODIUM CHLORIDE (UROMATIC P.C.) 0.9%	5000 ML	PC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0047-44		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE 0.9%	1000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0047-46		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE 0.9%	2000 ML	BO	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0047-47		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE 0.9%	3000 ML	FC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0048-02		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE 0.9%	250 ML	PC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0048-03		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE 0.9%	500 ML	PC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0048-04		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (P.C.) 0.9%	1000 ML	PC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0048-05		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE 0.9%	1500 ML	PC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0049-01		J7050		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE 0.9%	150 ML	FC	IV	ML		250 ML		0.004	01/01/2002	99/99/9999							
00338-0049-02		J7050		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE 0.9%	250 ML	FC	IV	ML		250 ML		0.004	01/01/2002	99/99/9999							
00338-0049-03		J7040		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION, STERILE (500 ML=1 UNIT)	SODIUM CHLORIDE 0.9%	500 ML	FC	IV	ML		500 ML		0.002	01/01/2002	99/99/9999							
00338-0049-04		J7030		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 1000 CC	SODIUM CHLORIDE 0.9%	1000 ML	FC	IV	ML		1000 ML		0.001	01/01/2002	99/99/9999							
00338-0049-10		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (QUAD PACK, MINI-BAG) 0.9%	25 ML	FC	IV	ML		10 ML		0.1	01/01/2004	99/99/9999							
00338-0049-11		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (QUAD PACK, MINI-BAG) 0.9%	50 ML	FC	IV	ML		10 ML		0.1	01/01/2004	99/99/9999							
00338-0049-18		J7050		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (QUAD PACK, MINI-BAG) 0.9%	100 ML	FC	IV	ML		250 ML		0.004	01/01/2002	99/99/9999							
00338-0049-31		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (MULTI PACK, MINI-BAG) 0.9%	50 ML	FC	IV	ML		10 ML		0.1	01/01/2004	99/99/9999							
00338-0049-38		J7050		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (MULTI PACK, MINI-BAG) 0.9%	100 ML	FC	IV	ML		250 ML		0.004	01/01/2002	99/99/9999							
00338-0049-41		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (SINGLE PACK, MINI-BAG) 0.9%	50 ML	FC	IV	ML		10 ML		0.1	01/01/2004	99/99/9999							
00338-0049-48		J7050		01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (SINGLE PACK, MINI-BAG) 0.9%	100 ML	FC	IV	ML		250 ML		0.004	01/01/2002	99/99/9999							
00338-0050-47		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (PROCESSING) 0.9%	3000 ML	PC	IR	ML		500 ML		0.002	01/01/2004	99/99/9999							
00338-0054-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE 3%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0056-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE 5% DEXTROSE/SODIUM CHLORIDE 2.5%-0.45%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0073-04		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.2%	1000 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0077-02		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.2%	250 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0077-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.2%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0077-04		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.2% RHOGAM ULTRA-FILTERED PLUS (INNER PACK,PF) 300 MCG	1000 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00562-7805-00		J2790		01/08/2014	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, FULL DOSE, 300 MICROGRAMS (1500 I.U.)		1 EA	SR	IM	EA		300 MCG		1	01/08/2014	99/99/9999							
00338-0081-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.33%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0085-02		J7799		01/01/2002	07/16/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.45%	250 ML	FC	IV	ML		1 EA		1	01/01/2002	07/16/2016							
00338-0085-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.45%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0085-04		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE 5%-0.45%	1000 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0089-03		J7042		01/01/2002	99/99/9999	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTROSE/SODIUM CHLORIDE 5%-0.9%	500 ML	FC	IV	ML		5 %		0.002	01/01/2002	99/99/9999							
00338-0089-04		J7042		01/01/2002	99/99/9999	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTROSE/SODIUM CHLORIDE 5%-0.9%	1000 ML	FC	IV	ML		5 %		0.002	01/01/2002	99/99/9999							
00338-0117-02		J7120		01/01/2002	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S	250 ML	FC	IV	ML		1000 ML		0.001	01/01/2002	99/99/9999							
00338-0117-03		J7120		01/01/2002	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S	500 ML	FC	IV	ML		1000 ML		0.001	01/01/2002	99/99/9999							
00338-0117-04		J7120		01/01/2002	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S	1000 ML	FC	IV	ML		1000 ML		0.001	01/01/2002	99/99/9999							
00338-0125-03		J7120		01/01/2002	12/31/2015	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S/DEXTROSE 5%	500 ML	FC	IV	ML		1000 ML		0.0005	01/01/2002	12/31/2015							
00338-0125-04		J7120		01/01/2002	12/31/2015	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S/DEXTROSE 5%	1000 ML	FC	IV	ML		1000 ML		0.0005	01/01/2002	12/31/2015							
00338-0351-04		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	OSMITROL (VIAFLEX AF) 5%	1000 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0353-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	OSMITROL (VIAFLEX) 10%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0355-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	OSMITROL (VIAFLEX AF) 15%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0357-02		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	OSMITROL (VIAFLEX) 20%	250 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0357-03		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	OSMITROL (VIAFLEX) 20%	500 ML	FC	IV	ML		1 EA		1	01/01/2002	99/99/9999							
00338-0409-02		J2001		01/01/2004	05/31/2012	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTOSE/LIDOCAINE HCL 5%-0.4%	250 ML	FC	IV	ML		10 MG		0.4	01/01/2004	05/31/2012							
00338-0409-03		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTOSE/LIDOCAINE HCL 5%-0.4%	500 ML	FC	IV	ML		10 MG		0.4	01/01/2004	99/99/9999							
00338-0411-02		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTOSE/LIDOCAINE HCL 5%-0.8%	250 ML	FC	IV	ML		10 MG		0.8	01/01/2004	99/99/9999							
00338-0431-03		J1644		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE 200 U/100 ML-0.9%	500 ML	FC	IV	ML		1000 U		0.002	01/01/2002	02/03/2016							
00338-0433-04		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE 200 U/100 ML-0.9%	1000 ML	FC	IV	ML		1000 U		0.002	01/01/2002	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00338-0505-48	J1580			01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE 100 MG/100 ML-0.9%	100	ML	FC	IV	ML	80	MG	0.0125	01/01/2002	99/99/9999							
00338-0507-41	J1580			01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (24X50ML) 1.2 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.015	01/01/2002	99/99/9999							
00338-0507-48	J1580			01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (24X100ML) 1.2 MG/ML-0.9%	100	ML	FC	IV	ML	80	MG	0.015	01/01/2002	99/99/9999							
00338-0509-41	J1580			01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE 1.6 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.02	01/01/2002	99/99/9999							
00338-0511-41	J1580			01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE 2 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.025	01/01/2002	99/99/9999							
00338-0551-11	J7060			01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (MINI-BAG PLUS) 5%	50	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00338-0551-18	J7060			01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (MINI-BAG PLUS) 5%	100	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999							
00338-0553-11	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (MINI-BAG PLUS) 0.9%	50	ML	FC	IV	ML	10	ML	0.1	01/01/2004	99/99/9999							
00338-0553-18	J7050			01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (MINI-BAG PLUS) 0.9%	100	ML	FC	IV	ML	250	ML	0.004	01/01/2002	99/99/9999							
00338-0691-04	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE/SODIUM CHLORIDE 2 MEQ/100 ML-0.9%	1000	ML	FC	IV	ML	2	MEQ	0.01	01/01/2002	99/99/9999							
00338-0695-04	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE/SODIUM CHLORIDE 4 MEQ/100 ML-0.9%	1000	ML	FC	IV	ML	2	MEQ	0.02	01/01/2002	99/99/9999							
00338-0703-41	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE 20 MEQ/50 ML	50	ML	PC	IV	ML	2	MEQ	0.2	01/01/2002	99/99/9999							
00338-0703-48	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE 40 MEQ/100 ML	100	ML	PC	IV	ML	2	MEQ	0.2	01/01/2002	99/99/9999							
00338-0704-34	J3480			05/21/2003	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE/SODIUM CHLORIDE (VIAFLEX BAG,PF) 2 MEQ/100 ML-0.45%	1000	ML	FC	IV	ML	2	MEQ	0.01	05/21/2003	99/99/9999							
00338-0705-41	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE 10 MEQ/50 ML	50	ML	PC	IV	ML	2	MEQ	0.1	01/01/2002	99/99/9999							
00338-0705-48	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE 20 MEQ/100 ML	100	ML	PC	IV	ML	2	MEQ	0.1	01/01/2002	99/99/9999							
00338-0709-48	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE 10 MEQ/100 ML	100	ML	PC	IV	ML	2	MEQ	0.05	01/01/2002	99/99/9999							
00338-0719-06	J7799			01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (BULK PACKAGE) 70%	2000	ML	PC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00338-0719-13	J7799			01/01/2002	10/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (12X500ML,USP) 70%	500	ML	PC	IV	ML	1	EA	1	01/01/2002	10/31/2015							
00338-0811-04	J7120			01/01/2002	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	POTASSIUM CHLORIDE SOLUTION (5%,DEXTROSE & LAC-RING)	1000	ML	FC	IV	ML	1000	ML	0.0005	01/01/2002	99/99/9999							
00338-1005-02	J1265			01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL (PRE-MIX IN D5W) 5%-80 MG/100 ML	250	ML	PC	IV	ML	40	MG	0.02	01/01/2006	99/99/9999							
69097-0318-87	J7626			11/14/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.25 MG/2 ML	2	ML	AM	IH	ML	0.5	MG	0.25	11/14/2017	99/99/9999							
44567-0701-25	J0696			04/25/2013	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 1 GM	25	EA	VL	IJ	EA	250	MG	4	04/25/2013	99/99/9999							
00338-1005-03	J1265			01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL 5%-80 MG/100 ML	500	ML	PC	IV	ML	40	MG	0.02	01/01/2006	99/99/9999							
00338-1007-02	J1265			01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL 5%-160 MG/100 ML	250	ML	PC	IV	ML	40	MG	0.04	01/01/2006	99/99/9999							
00338-1007-03	J1265			01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL 5%-160 MG/100 ML	500	ML	PC	IV	ML	40	MG	0.04	01/01/2006	99/99/9999							
00338-1009-02	J1265			01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL 5%-320 MG/100 ML	250	ML	PC	IV	ML	40	MG	0.08	01/01/2006	99/99/9999							
00338-1013-41	J2700			01/01/2002	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN SODIUM (PREMIXED) 1 GM/50 ML	50	ML	PC	IV	ML	250	MG	0.08	01/01/2002	99/99/9999							
00338-1015-41	J2700			01/01/2002	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN SODIUM (PREMIXED) 2 GM/50 ML	50	ML	PC	IV	ML	250	MG	0.16	01/01/2002	99/99/9999							
00338-1017-41	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM (GALAXY,PREMIX) 1 GM/50 ML	50	ML	PC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00338-1019-48	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM (GALAXY,PREMIX) 1 GM/50 ML	100	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00338-1021-41	J2540			01/01/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PENICILLIN G POTASSIUM (GALAXY,PREMIX) 1 Million U/50 ML	50	ML	PC	IV	ML	600000	U	0.03333	01/01/2002	99/99/9999							
00338-1023-41	J2540			01/01/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PENICILLIN G POTASSIUM (GALAXY,PREMIX) 2 Million U/50 ML	50	ML	PC	IV	ML	600000	U	0.06666	01/01/2002	99/99/9999							
00338-1025-41	J2540			01/01/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PENICILLIN G POTASSIUM (GALAXY,PREMIX) 3 Million U/50 ML	50	ML	PC	IV	ML	600000	U	0.1	01/01/2002	99/99/9999							
00338-1055-48	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE 500 MG/100 ML	100	ML	FC	IV	ML	1	EA	1	01/01/2002	99/99/9999							
00338-1073-02	J1250			01/01/2002	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DEXTROSE/DOBUTAMINE 5%-100 MG/100 ML	250	ML	FC	IV	ML	250	MG	0.004	01/01/2002	99/99/9999							
00338-1075-02	J1250			01/01/2002	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DEXTROSE/DOBUTAMINE 5%-200 MG/100 ML	250	ML	FC	IV	ML	250	MG	0.008	01/01/2002	99/99/9999							
00338-1077-02	J1250			01/01/2002	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DEXTROSE/DOBUTAMINE 5%-400 MG/100 ML	250	ML	FC	IV	ML	250	MG	0.016	01/01/2002	99/99/9999							
00338-1762-41	J2405			12/27/2006	07/25/2012	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (50MLX10,SD,USP,PREMIX) 32 MG/50 ML	50	ML	PC	IV	ML	1	MG	0.64	12/27/2006	07/25/2012							
00338-3503-41	J0690			01/01/2002	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (GALAXY P.C.) 1 GM/50 ML	50	ML	FC	IV	ML	500	MG	0.04	01/01/2002	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00338-3551-48		J3370		01/01/2002	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANOCIN HCL (S.D. GALAXY PLASTIC) 5%-500 MG/100 ML	100	ML	PC	IV	ML	500	MG	0.01	01/01/2002	99/99/9999						
00338-3552-48		J3370		01/01/2002	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANOCIN HCL (S.D. GALAXY PLASTIC) 5%-500 MG/100 ML	200	ML	PC	IV	ML	500	MG	0.01	01/01/2002	99/99/9999						
00338-5002-41		J0696		09/06/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 1 GM/50 ML	50	ML	PC	IV	ML	250	MG	0.08	09/06/2005	99/99/9999						
00338-5003-41		J0696		09/06/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 2 GM/50 ML	50	ML	PC	IV	ML	250	MG	0.16	09/06/2005	99/99/9999						
00338-5197-41		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	FAMOTIDINE (GALAXY PC-PF) 0.4 MG/ML	50	ML	PC	IV	ML	1	EA	1	01/01/2002	99/99/9999						
00338-6010-48		J2260		06/05/2002	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	DEXTROSE/MILRINONE LACTATE (BAG,INTRAVIA) 5%-20 MG/100 ML	100	ML	FC	IV	ML	5	MG	0.04	06/05/2002	99/99/9999						
00338-6011-37		J2260		06/05/2002	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	DEXTROSE/MILRINONE LACTATE (BAG,INTRAVIA) 5%-20 MG/100 ML	200	ML	FC	IV	ML	5	MG	0.04	06/05/2002	99/99/9999						
00338-6045-37		J1450		07/29/2004	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (INTRAVIA CONTAINER) 400 MG/200 ML	200	ML	PC	IV	ML	200	MG	0.01	07/29/2004	99/99/9999						
00338-6046-48		J1450		07/29/2004	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (INTRAVIA CONTAINERS) 200 MG/100 ML	100	ML	PC	IV	ML	200	MG	0.01	07/29/2004	99/99/9999						
00338-6307-02		J7120		10/17/2007	06/30/2016	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (USP,LATEX-FREE)	250	ML	FC	IV	ML	1000	ML	0.001	10/17/2007	06/30/2016						
00338-6346-02		J7060		03/01/2007	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (USP,40X250ML,AVIVA) 5%	250	ML	FC	IV	ML	500	ML	0.002	03/01/2007	99/99/9999						
00378-0014-01	None			01/01/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	100	EA	BO	PO	EA	2.5	MG	1	01/01/1994	99/99/9999						
00378-0014-50	None			02/23/1998	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	5000	EA	BO	PO	EA	2.5	MG	1	02/23/1998	99/99/9999						
00378-0144-05	J8999			02/20/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 10 MG	500	EA	BO	PO	EA	1	EA	1	02/20/2003	99/99/9999						
00378-0144-91	J8999			02/20/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 10 MG	60	EA	BO	PO	EA	1	EA	1	02/20/2003	99/99/9999						
00378-0253-01	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
00378-0274-01	J8999			02/20/2003	07/12/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 20 MG	100	EA	BO	PO	EA	1	EA	1	02/20/2003	07/12/2016						
00378-0274-93	J8999			02/20/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 20 MG	30	EA	BO	PO	EA	1	EA	1	02/20/2003	99/99/9999						
00378-0302-01	J8499			01/01/2002	01/14/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	01/14/2016						
00378-1003-94	Q0166			01/30/2007	11/30/2012	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (FILM-COATED) 1 MG	20	EA	BO	PO	EA	1	MG	1	01/30/2007	11/30/2012						
00378-3266-94	None			10/19/2001	99/99/9999	ETOPOSIDO, 50 MG, ORAL	ETOPOSIDO (BLISTER PACK,SOFTGEL) 50 MG	20	EA	BX	PO	EA	50	MG	1	10/19/2001	99/99/9999						
00378-3547-25	J8999			07/01/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MERCAPTOPYRINE (U.S.P.) 50 MG	250	EA	BO	PO	EA	1	EA	1	07/01/2005	99/99/9999						
00378-3547-52	J8999			07/01/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MERCAPTOPYRINE (U.S.P.) 50 MG	25	EA	BO	PO	EA	1	EA	1	07/01/2005	99/99/9999						
00378-5105-01	Q0164			01/01/2002	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2002	99/99/9999						
00378-5110-01	Q0165			01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013						
00378-6988-58	J7620			12/28/2007	09/25/2013	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	ALBUTEROL SULFATE (30X3ML,5 VIALS/POUCH) 3 MG/3 ML-0.5 MG/3 ML	30	ML	PC	IH	ML	3	MG	0.33333	12/28/2007	09/25/2013						
00378-6988-91	J7620			12/28/2007	12/31/2014	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	ALBUTEROL SULFATE (60X3ML,5 VIALS/POUCH) 3 MG/3 ML-0.5 MG/3 ML	60	ML	PC	IH	ML	3	MG	0.33333	12/28/2007	12/31/2014						
00378-6988-93	J7620			12/28/2007	06/12/2013	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML,1 VIAL/POUCH) 3 MG/3 ML-0.5 MG/3 ML	30	ML	PC	IH	ML	3	MG	0.33333	12/28/2007	06/12/2013						
00406-0646-02	J0706			01/01/2002	99/99/9999	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATED (PURIFIED) 50 MG	1	EA	BO	NA	GM	5	MG	200	01/01/2002	99/99/9999						
00406-0672-52	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P.)	1	EA	BO	NA	GM	1	EA	1	01/01/2002	99/99/9999						
00406-1130-52	J3010			01/01/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE	1	EA	BO	NA	GM	0.1	MG	10000	01/01/2002	99/99/9999						
00406-1395-04	J3520			01/01/2002	99/99/9999	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM (U.S.P.)	1	EA	BO	NA	GM	150	MG	6.66666	01/01/2002	99/99/9999						
00406-1492-52	J2310			01/01/2002	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
00406-1510-56	J1230			01/01/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL	1	EA	BO	NA	GM	10	MG	100	01/01/2002	99/99/9999						
00406-1510-57	J1230			01/01/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL	1	EA	BO	NA	GM	10	MG	100	01/01/2002	99/99/9999						
00406-1510-59	J1230			01/01/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL	1	EA	BO	NA	GM	10	MG	100	01/01/2002	99/99/9999						
00406-1521-53	J2271			01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE	1	EA	BO	NA	GM	100	MG	10	01/01/2002	12/31/2014						
00406-1521-55	J2271			01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE	1	EA	BO	NA	GM	100	MG	10	01/01/2002	12/31/2014						
00406-1521-56	J2271			01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE	1	EA	BO	NA	GM	100	MG	10	01/01/2002	12/31/2014						
00406-1521-57	J2271			01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE	1	EA	BO	NA	GM	100	MG	10	01/01/2002	12/31/2014						
00406-1548-32	J0745			01/01/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE	1	EA	BO	NA	GM	30	MG	33.33333	01/01/2002	99/99/9999						
00406-1548-35	J0745			01/01/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE	1	EA	BO	NA	GM	30	MG	33.33333	01/01/2002	99/99/9999						
00406-1585-55	J2175			01/01/2002	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	01/01/2002	99/99/9999						
00406-3245-52	J1170			01/01/2002	09/30/2016	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL	1	EA	BO	NA	GM	4	MG	250	01/01/2002	09/30/2016						
00406-4200-12	J3475			01/01/2002	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (U.S.P.)	1	EA	BO	NA	GM	500	MG	2	01/01/2002	99/99/9999						
00406-6838-04	J3480			01/01/2002	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	2	MEQ	6.71141	01/01/2002	10/17/2016						
00406-6838-06	J3480			01/01/2002	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	2	MEQ	6.71141	01/01/2002	10/17/2016						
00406-6845-04	J3480			01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	2	MEQ	6.71141	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00406-6858-04		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (A.C.S.)	1 EA	NA	NA	GM		2 MEQ		6.71141	01/01/2002	99/99/9999						
00406-8050-03		J9218		01/01/2002	10/17/2016	LEUPROLIDE ACETATE, PER 1 MG	LEUPROLIDE ACETATE	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	10/17/2016						
00406-8642-12		J3350		01/01/2002	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P.)	1 EA	BO	NA	GM		40 GM		0.025	01/01/2002	99/99/9999						
00409-0801-01		J9268		07/20/2007	99/99/9999	INJECTION, PENTOSTATIN, 10 MG	NIPENT (SDV) 10 MG	1 EA	VL	IV	EA		10 MG		1	07/20/2007	99/99/9999						
00409-1036-30		J0670		03/21/2006	99/99/9999	INJECTION, MEPIVACAINE HYDROCHLORIDE, PER 10 ML	CARBOCAINE (MDV)	30 ML	VL	IJ	ML		10 ML		0.1	03/21/2006	99/99/9999						
00409-1038-50		J0670		10/08/2007	99/99/9999	INJECTION, MEPIVACAINE HYDROCHLORIDE, PER 10 ML	CARBOCAINE (MDV) 1%	50 ML	VL	IJ	ML		10 ML		0.1	10/08/2007	99/99/9999						
00409-1041-30		J0670		04/26/2006	99/99/9999	INJECTION, MEPIVACAINE HYDROCHLORIDE, PER 10 ML	CARBOCAINE (PF) 1.5%	30 ML	VL	IJ	ML		10 ML		0.1	04/26/2006	99/99/9999						
00409-1067-20		J0670		01/15/2007	99/99/9999	INJECTION, MEPIVACAINE HYDROCHLORIDE, PER 10 ML	CARBOCAINE (SDV,USP,PF) 2%	20 ML	VL	IJ	ML		10 ML		0.1	01/15/2007	99/99/9999						
69097-0318-87	KO	J7626	KO	11/14/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.25 MG/2 ML	2 ML	AM	IH	ML		0.5 MG		0.25	11/14/2017	99/99/9999						
00409-1081-51		A4216		12/27/2006	09/11/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (THERMOJECT, 25X10ML) 0.9%	10 ML	VL	IV	ML		10 ML		0.1	12/27/2006	09/11/2016						
00409-1082-01		J7060		04/25/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (THERMOJECT KIT) 5%	10 ML	VL	IV	EA		500 ML		0.08	04/25/2005	99/99/9999						
00409-1082-51		J7060		03/29/2006	06/01/2012	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (USP,PF) 5%	10 ML	VL	IV	ML		500 ML		0.002	03/29/2006	06/01/2012						
00409-1120-62		J2405		01/22/2007	03/01/2013	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (10X2ML,SDPFS,USP) 2 MG/ML	2 ML	SR	IJ	ML		1 MG		2	01/22/2007	03/01/2013						
00409-1130-02		J7799		05/13/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE 23.4%	250 ML	GC	IV	ML		1 EA		1	05/13/2005	99/99/9999						
00069-0291-10		Q5110		09/05/2018	99/99/9999	INJECTION, FILGRASTIM-AAFI, BIOSIMILAR, (NIVESTYM), 1 MICROGRAM	NIVESTYM (PF,LATEX-FREE) 300 MCG/0.5 ML	0.5 ML	SR	IJ	ML		1 MCG		600	09/05/2018	99/99/9999						
00409-1134-03		J2271		09/14/2005	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (VIAL, FLIPTOP) 50 MG/ML	20 ML	VL	IJ	ML		100 MG		0.5	09/14/2005	12/31/2014						
00409-1134-05		J2271		08/08/2005	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (LATEX-FREE) 50 MG/ML	50 ML	VL	IJ	ML		100 MG		0.5	08/08/2005	12/31/2014						
00409-1135-02		J2275		07/21/2005	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (HIGH CONCENTRATION,PF) 25 MG/ML	10 ML	VL	IJ	ML		10 MG		2.5	07/21/2005	12/31/2014						
00409-1141-02		J7799		04/13/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (VIAL,FLIPTOP,BULK PKG) 23.4%	100 ML	VL	IV	ML		1 EA		1	04/13/2005	99/99/9999						
00409-1158-01		J3490		07/27/2005	11/01/2016	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (AMP,5X30ML,LATEX-FREE) 0.25%	30 ML	AM	IJ	ML		1 EA		1	07/27/2005	11/01/2016						
00409-1159-01		J3490		06/29/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (USP,25X2ML,LATEX-FREE) 0.25%	10 ML	VL	IJ	ML		1 EA		1	06/29/2005	99/99/9999						
00409-1159-02		J3490		08/10/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (25X30ML,LATEX-FREE) 0.25%	30 ML	VL	IJ	ML		1 EA		1	08/10/2005	99/99/9999						
00409-1160-01		J3490		04/12/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (VIAL,FLIPTOP,LATEX-FREE) 0.25%	50 ML	VL	IJ	ML		1 EA		1	04/12/2005	99/99/9999						
00409-1161-01		J3490		10/18/2004	12/08/2017	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (AMP,LATEX-FREE) 0.5%	30 ML	AM	IJ	ML		1 EA		1	10/18/2004	12/08/2017						
00409-1162-01		J3490		03/08/2006	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (25X10ML) 0.5%	10 ML	VL	IJ	ML		1 EA		1	03/08/2006	99/99/9999						
00409-1162-02		J3490		11/22/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (VIAL,LATEX-FREE) 0.5%	30 ML	VL	IJ	ML		1 EA		1	11/22/2005	99/99/9999						
00409-1163-01		J3490		03/30/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (VIAL,FLIPTOP,LATEX-FREE) 0.5%	50 ML	VL	IJ	ML		1 EA		1	03/30/2005	99/99/9999						
00409-1165-01		J3490		12/08/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (VIAL,LATEX-FREE) 0.75%	10 ML	VL	IJ	ML		1 EA		1	12/08/2005	99/99/9999						
00409-1165-02		J3490		05/24/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (TTV,LATEX-FREE) 0.75%	30 ML	VL	IJ	ML		1 EA		1	05/24/2005	99/99/9999						
00409-1176-30		J2175		08/25/2005	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE (LLK,SLIM PK,LATEX-FREE) 25 MG/ML	1 ML	SR	IJ	ML		100 MG		0.25	08/25/2005	99/99/9999						
00409-1178-30		J2175		09/14/2005	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE (LATEX-FREE,CARPUJECT) 50 MG/ML	1 ML	SR	IJ	ML		100 MG		0.5	09/14/2005	99/99/9999						
00069-0292-01		Q5110		09/05/2018	99/99/9999	INJECTION, FILGRASTIM-AAFI, BIOSIMILAR, (NIVESTYM), 1 MICROGRAM	NIVESTYM (PF,LATEX-FREE) 480 MCG/0.8 ML	0.5 ML	SR	IJ	ML		1 MCG		600	09/05/2018	99/99/9999						
00409-1179-30		J2175		12/08/2005	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE (LATEX-FREE,CARPUJECT) 75 MG/ML	1 ML	SR	IJ	ML		100 MG		0.75	12/08/2005	99/99/9999						
00409-1180-69		J2175		09/14/2005	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE (CARPUJECT) 100 MG/ML	1 ML	SR	IJ	ML		100 MG		1	09/14/2005	99/99/9999						
00409-1181-30		J2175		01/31/2006	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL (USP,MDV,STERILE) 50 MG/ML	30 ML	VL	IJ	ML		100 MG		0.5	01/31/2006	99/99/9999						
00409-1187-01		J1790		08/23/2005	99/99/9999	INJECTION, DROPERIDOL, UP TO 5 MG	DROPERIDOL (10X2ML AMP,LATEX-FREE) 2.5 MG/ML	2 ML	AM	IJ	ML		5 MG		0.5	08/23/2005	99/99/9999						
00409-1201-20		J2175		03/09/2006	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL (MDV) 100 MG/ML	20 ML	VL	IJ	ML		100 MG		1	03/09/2006	99/99/9999						
00409-1203-01		J2175		12/16/2005	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE (UNI-AMP, 5X5,LATEX-FREE) 50 MG/ML	0.5 ML	AM	IJ	ML		100 MG		0.5	12/16/2005	99/99/9999						
00409-1207-03		J1580		08/30/2005	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (VIAL,FLIPTOP) 40 MG/ML	2 ML	VL	IJ	ML		80 MG		0.5	08/30/2005	99/99/9999						
00409-1215-01		J2310		07/08/2005	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (VIAL,FLIPTOP,10X1ML) 0.4 MG/ML	1 ML	VL	IJ	ML		1 MG		0.4	07/08/2005	99/99/9999						
00409-1219-01		J2310		04/03/2006	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HYDROCHLORIDE 0.4 MG/ML	10 ML	VL	IJ	ML		1 MG		0.4	04/03/2006	99/99/9999						
00409-1253-01		J2175		01/04/2006	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE (LATEX-FREE) 50 MG/ML	1 ML	AM	IJ	ML		100 MG		0.5	01/04/2006	99/99/9999						
00409-1254-01		J2175		03/20/2006	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL (25X1.5ML) 50 MG/ML	1.5 ML	AM	IJ	ML		100 MG		0.5	03/20/2006	99/99/9999						

NDC	NDC Mod	HPCS	HPCS Mod	Relationship Start Date	Relationship End Date	HPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Scouts of Administration	Billing Units	HPCS Amount #1	HPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-1255-02		J2175		11/23/2005	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMEROL HYDROCHLORIDE (UNI-AMP 5X5,LATEX-FREE) 50 MG/ML	2 ML	AM	IJ	ML	100 MG		0.5	11/23/2005	99/99/9999							
00409-1256-01		J2175		01/26/2006	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMEROL HYDROCHLORIDE (25X1ML,LATEX-FREE) 100 MG/ML	1 ML	AM	IJ	ML	100 MG		1	01/26/2006	99/99/9999							
00409-1258-30		J2270		05/10/2005	09/01/2013	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (LUER LOCK,U.S.P., 10X1ML) 4 MG/ML	1 ML	CR	IJ	ML	10 MG		0.4	05/10/2005	09/01/2013							
00409-1260-69		J2270		03/22/2006	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE 8 MG/ML	1 ML	SR	IJ	ML	10 MG		0.8	03/22/2006	99/99/9999							
00409-1261-30		J2270		07/21/2005	03/01/2014	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (LLK,SLIM PK, 10X1ML) 10 MG/ML	1 ML	SR	IJ	ML	10 MG		1	07/21/2005	03/01/2014							
00409-1264-31		J2271		12/16/2005	06/01/2013	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (LUER LOCK,LATEX-FREE) 15 MG/ML	1 ML	CR	IJ	ML	100 MG		0.15	12/16/2005	06/01/2013							
00409-1273-32		J3360		08/23/2005	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (10X2ML, LUER LOCK) 5 MG/ML	2 ML	CR	IJ	ML	5 MG		1	08/23/2005	99/99/9999							
00409-1276-32		J3010		07/27/2005	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (LUER LOCK,10X2ML,PF) 0.05 MG/ML	2 ML	SR	IJ	ML	0.1 MG		0.5	07/27/2005	99/99/9999							
00069-0292-10		Q5110		09/05/2018	99/99/9999	INJECTION, FILGRASTIM-AAFI, BIOSIMILAR, (NIVESTYM), 1 MICROGRAM	NIVESTYM (PF,LATEX-FREE) 480 MCG/0.8 ML	0.5 ML	SR	IJ	ML	1 MCG		600	09/05/2018	99/99/9999							
00409-1283-31		J1170		06/14/2005	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (LUER LOCK,10X1ML) 1 MG/ML	1 ML	CR	IJ	ML	4 MG		0.25	06/14/2005	99/99/9999							
00409-1304-31		J1170		07/13/2005	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (LUER LOCK,10X1ML) 4 MG/ML	1 ML	CR	IJ	ML	4 MG		1	07/13/2005	99/99/9999							
00409-1312-30		J1170		07/07/2005	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (10X1ML,LLK,SLIM PK) 2 MG/ML	1 ML	CR	IJ	ML	4 MG		0.5	07/07/2005	99/99/9999							
00409-1316-25		J1644		10/29/2007	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (10X0.5ML,W/ LUER LOCK) 5000 U/0.5 ML	0.5 ML	SR	IJ	ML	1000 U		10	10/29/2007	99/99/9999							
00409-1316-32		J1644		03/23/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM 10000 U/ML	0.5 ML	SR	IJ	ML	1000 U		10	03/23/2005	99/99/9999							
00409-1316-66		J1644		02/11/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (PF,CARPUJECT) 10000 U/ML	0.5 ML	SR	IJ	ML	1000 U		10	02/11/2005	99/99/9999							
00409-1317-02		J1165		03/30/2005	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (AMP,LATEX-FREE) 50 MG/ML	5 ML	AM	IV	ML	50 MG		1	03/30/2005	99/99/9999							
00409-1323-05		J2001		12/08/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (10X5ML, ANSYR) 2%	5 ML	SR	IJ	ML	10 MG		2	12/08/2005	99/99/9999							
00409-1402-31		J1644		03/21/2005	07/01/2012	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (LUER LOCK,CARPUJECT) 5000 U/ML	1 ML	SR	IJ	ML	1000 U		5	03/21/2005	07/01/2012							
00409-1410-01		J7660		01/01/2007	11/30/2013	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ISUPREL (AMP,25X1ML,LATEX-FREE) 0.2 MG/ML	1 ML	AM	IV	ML	1 MG		0.2	01/01/2007	11/30/2013							
00143-9270-01		J9200		09/21/2018	99/99/9999	INJECTION, FLOXURIDINE, 500 MG	FLOXURIDINE (LYOPHILIZED) 0.5 GM	1 EA	VL	IJ	EA	500 MG		1	09/21/2018	99/99/9999							
00409-1410-01	KO	J7660	KO	01/01/2007	11/30/2013	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ISUPREL (AMP,25X1ML,LATEX-FREE) 0.2 MG/ML	1 ML	AM	IV	ML	1 MG		0.2	01/01/2007	11/30/2013							
00409-1410-05		J7660		01/01/2007	11/30/2013	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ISUPREL (AMP,25X1ML,LATEX-FREE) 0.2 MG/ML	5 ML	AM	IV	ML	1 MG		0.2	01/01/2007	11/30/2013							
00409-1410-05	KO	J7660	KO	01/01/2007	11/30/2013	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ISUPREL (10X5ML,AMP,LATEX-FREE) 0.2 MG/ML	5 ML	AM	IV	ML	1 MG		0.2	01/01/2007	11/30/2013							
00409-1412-04		J3490		06/14/2006	99/99/9999	UNCLASSIFIED DRUGS	BUMETANIDE (SDFLIPTOP VIAL,USP) 0.25 MG/ML	4 ML	VL	IJ	ML	1 EA		1	06/14/2006	99/99/9999							
00409-1412-10		J3490		06/29/2006	99/99/9999	UNCLASSIFIED DRUGS	BUMETANIDE (MDV,USP,10X10ML) 0.25 MG/ML	10 ML	VL	IJ	ML	1 EA		1	06/29/2006	99/99/9999							
00409-1463-01		J2300		03/09/2005	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL (AMP,LATEX-FREE) 10 MG/ML	1 ML	AM	IJ	ML	10 MG		1	03/09/2005	99/99/9999							
00409-1464-01		J2300		07/13/2005	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL (25X10ML) 10 MG/ML	10 ML	VL	IJ	ML	10 MG		1	07/13/2005	99/99/9999							
00409-1465-01		J2300		11/18/2004	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL (AMP,LATEX-FREE) 20 MG/ML	1 ML	AM	IJ	ML	10 MG		2	11/18/2004	99/99/9999							
00409-1467-01		J2300		05/12/2005	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL (VIAL,FLIPTOP) 20 MG/ML	10 ML	VL	IJ	ML	10 MG		2	05/12/2005	99/99/9999							
00409-1508-05		J7799		08/31/2005	05/18/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (6X1000ML) 2.5%	1000 ML	GC	IV	ML	1 EA		1	08/31/2005	05/18/2016							
00264-7055-10		J2400		09/17/2018	99/99/9999	INJECTION, CHLOROPROCAINE HYDROCHLORIDE, PER 30 ML	CLOTROKAL 10 MG/1 ML	5 ML	VL	IN	ML	30 ML		0.03333	09/17/2018	99/99/9999							
00409-1513-02		J3480		06/16/2005	06/01/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (12X250ML,LATEX-FREE) 2 MEQ/ML	250 ML	VL	IV	ML	2 MEQ		1	06/16/2005	06/01/2016							
00409-1522-01		J7060		04/11/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (12X150ML) 5%	150 ML	GC	IV	ML	500 ML		0.002	04/11/2005	99/99/9999							
00409-1522-02		J7060		03/09/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (12X250ML) 5%	250 ML	GC	IV	ML	500 ML		0.002	03/09/2005	99/99/9999							
00409-1522-03		J7060		06/16/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (12X500ML) 5%	500 ML	GC	IV	ML	500 ML		0.002	06/16/2005	99/99/9999							
00409-1523-01		J7060		09/16/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (50/150ML PART FILL) 5%	50 ML	GC	IV	ML	500 ML		0.002	09/16/2005	99/99/9999							
00409-1523-11		J7060		07/27/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (12X100ML) 5%	100 ML	GC	IV	ML	500 ML		0.002	07/27/2005	99/99/9999							
00409-1534-05		J7799		02/24/2006	05/18/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE AND SODIUM CHLORIDE (6X1000ML) 10%-0.9%	1000 ML	GC	IV	ML	1 EA		1	02/24/2006	05/18/2016							
00409-1535-03		J7799		09/08/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (12X500ML) 20%	500 ML	GC	IV	ML	1 EA		1	09/08/2005	99/99/9999							
00409-1539-31		J2060		12/23/2005	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (10X1ML, LUER LOCK) 4 MG/ML	1 ML	CR	IJ	ML	2 MG		2	12/23/2005	99/99/9999							
00409-1559-10		J3490		08/22/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (10X10ML, S.D.V.) 0.25%	10 ML	VL	IJ	ML	1 EA		1	08/22/2005	99/99/9999							
00409-1559-30		J3490		09/07/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (S.D.V.,LATEX-FREE) 0.25%	30 ML	VL	IJ	ML	1 EA		1	09/07/2005	99/99/9999							
00409-1560-10		J3490		08/31/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (S.D.V.) 0.5%	10 ML	VL	IJ	ML	1 EA		1	08/31/2005	99/99/9999							
00409-1560-29		J3490		08/05/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (S.D.V.) 0.5%	30 ML	VL	IJ	ML	1 EA		1	08/05/2005	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-1582-10		J3490		07/22/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (10X10ML, S.D.V.) 0.75%	10 ML	VL	IJ	ML		1 EA		1	07/22/2005	99/99/9999						
00409-1582-29		J3490		08/04/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (10X30ML,LATEX-FREE) 0.75%	30 ML	VL	IJ	ML		1 EA		1	08/04/2005	99/99/9999						
00409-1583-01		J7050		07/20/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (12X150ML,PF) 0.9%	150 ML	FC	IV	ML		250 ML		0.004	07/20/2005	99/99/9999						
00409-1583-02		J7050		09/14/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (12X250ML,PF) 0.9%	250 ML	GC	IV	ML		250 ML		0.004	09/14/2005	99/99/9999						
00409-1584-11		J7050		09/16/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (12X100ML,150ML VIAL,PF) 0.9%	100 ML	GC	IV	ML		250 ML		0.004	09/16/2005	99/99/9999						
00409-1586-03		J7799		03/24/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (12X500ML) 5%	500 ML	GC	IV	ML		1 EA		1	03/24/2006	99/99/9999						
00409-1587-50		J3490		01/10/2006	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (M.D.V.,LATEX-FREE) 0.25%	50 ML	VL	IJ	ML		1 EA		1	01/10/2006	99/99/9999						
00409-1590-02		A4217		08/05/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION (12X250ML,PF,LATEX-FREE)	250 ML	GC	IV	ML		500 ML		0.002	08/05/2005	99/99/9999						
00409-1610-50		J3490		11/22/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL (M.D.V.) 0.5%	50 ML	VL	IJ	ML		1 EA		1	11/22/2005	99/99/9999						
00409-1623-01		J0595		09/20/2005	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (10X1ML) 1 MG/ML	1 ML	VL	IJ	ML		1 MG		1	09/20/2005	99/99/9999						
00378-9692-52	KO	J7614	KO	09/10/2018	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	LEVALBUTEROL (PF) 1.25 MG/3 ML BUTORPHANOL TARTRATE NOVATION (10X1ML) 1 MG/ML	3 ML	VL	IH	ML		0.5 MG		0.83333	09/10/2018	99/99/9999						
00409-1623-49		J0595		10/19/2005	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (10X1ML) 2 MG/ML	1 ML	VL	IJ	ML		1 MG		1	10/19/2005	99/99/9999						
00409-1626-01		J0595		03/21/2006	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (10X2ML) 2 MG/ML	1 ML	VL	IJ	ML		1 MG		2	03/21/2006	99/99/9999						
00409-1626-02		J0595		12/21/2005	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (10X2ML) 2 MG/ML	2 ML	VL	IJ	ML		1 MG		2	12/21/2005	99/99/9999						
00409-1626-49		J0595		05/24/2006	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	NOVAPLUS BUTORPHANOL TARTRATE (VHA,10X1ML) 2 MG/ML	1 ML	VL	IJ	ML		1 MG		2	05/24/2006	99/99/9999						
00409-1626-51		J0595		12/08/2005	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE NOVATION (10X2ML) 2 MG/ML	2 ML	VL	IJ	ML		1 MG		2	12/08/2005	99/99/9999						
00409-1639-10		J1940		01/23/2006	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (10X10ML, ANSYR) 10 MG/ML	10 ML	SR	IJ	ML		20 MG		0.5	01/23/2006	99/99/9999						
00409-1754-10		J3475		11/27/2006	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (10X10ML,SINGLE-DOSE USP) 500 MG/ML	10 ML	SR	IJ	ML		500 MG		1	11/27/2006	99/99/9999						
00409-1761-02		J3490		06/06/2005	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE SPINAL (AMP,W/DEXTROSE,PF) 0.75%	2 ML	AM	IJ	ML		1 EA		1	06/06/2005	99/99/9999						
00409-1762-30		J2270		05/27/2005	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (LLK,SLIM PK,CARPUJECT) 2 MG/ML	1 ML	CR	IJ	ML		10 MG		0.2	05/27/2005	99/99/9999						
00409-1775-10		J7799		02/20/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTOSE (2.5GM INFANT ANSYR SYR) 25%	10 ML	SR	IV	ML		1 EA		1	02/20/2006	99/99/9999						
00409-1782-69		J2310		09/29/2005	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (10X1ML, CARPUJECT) 0.4 MG/ML	1 ML	SR	IJ	ML		1 MG		0.4	09/29/2005	99/99/9999						
00409-1800-01		J2370		04/14/2005	99/99/9999	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	NEO-SYNEPHRINE HCL (AMP,25X1ML) 10 MG/ML	1 ML	AM	IJ	ML		1 ML		1	04/14/2005	99/99/9999						
00409-1902-01		J2690		03/10/2006	99/99/9999	INJECTION, PROCAINAMIDE HCL, UP TO 1 GM	PROCAINAMIDE HYDROCHLORIDE (25X10ML,FTV) 100 MG/ML	10 ML	VL	IJ	ML		1 GM		0.1	03/10/2006	99/99/9999						
00409-1903-01		J2690		08/24/2005	99/99/9999	INJECTION, PROCAINAMIDE HCL, UP TO 1 GM	PROCAINAMIDE HCL 500 MG/ML	2 ML	VL	IV	ML		1 GM		0.5	08/24/2005	99/99/9999						
00409-1918-32		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (LUER LOCK,50X2ML,PF) 0.9%	2 ML	CR	IV	ML		10 ML		0.1	01/01/2007	99/99/9999						
00409-1918-33		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (LUER LOCK,PF,LATEX-FREE) 0.9%	5 ML	CR	IV	ML		10 ML		0.1	01/01/2007	99/99/9999						
00409-1918-35		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (LUER LOCK,PF,LATEX-FREE) 0.9%	5 ML	CR	IV	ML		10 ML		0.1	01/01/2007	99/99/9999						
00409-1920-10		J3070		09/29/2005	99/99/9999	INJECTION, PENTAZOCINE, 30 MG	TALWIN LACTATE (VIAL,LATEX-FREE) 30 MG/ML	10 ML	VL	IJ	ML		30 MG		1	09/29/2005	99/99/9999						
00409-1941-01		J3070		11/18/2005	03/01/2018	INJECTION, PENTAZOCINE, 30 MG	TALWIN LACTATE (UNI-AMP,LATEX-FREE) 30 MG/ML	1 ML	AM	IJ	ML		30 MG		1	11/18/2005	03/01/2018						
00409-1955-01		J0278		01/01/2006	11/15/2012	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (VIAL,FLIPTOP,LATEX-FREE) 50 MG/ML	2 ML	VL	IJ	ML		100 MG		0.5	01/01/2006	11/15/2012						
00409-1956-01		J0278		01/01/2006	11/01/2012	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (10X2ML) 250 MG/ML	2 ML	VL	IJ	ML		100 MG		2.5	01/01/2006	11/01/2012						
00378-9692-52	KO	J7614	KO	09/10/2018	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	LEVALBUTEROL (PF) 1.25 MG/3 ML SODIUM CHLORIDE BACTERIOSTATIC (25X20ML,LATEX-FREE) 0.9%	3 ML	VL	IH	ML		0.5 MG		0.83333	09/10/2018	99/99/9999						
00409-1966-05		A4216		05/02/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE BACTERIOSTATIC (VIAL,FLIPTOP,PLASTIC) 0.9%	20 ML	VL	IV	ML		10 ML		0.1	05/02/2005	99/99/9999						
00409-1966-07		A4216		04/05/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE BACTERIOSTATIC (VIAL,FLIPTOP,PLASTIC) 0.9%	30 ML	VL	IV	ML		10 ML		0.1	04/05/2005	99/99/9999						
00409-1966-12		A4216		10/06/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE BACTERIOSTATIC (25X10ML, LS-PLASTIC) 0.9%	10 ML	VL	IV	ML		10 ML		0.1	10/06/2005	99/99/9999						
00409-1966-14		A4216		06/01/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE BACTERIOSTATIC (FLIPTOP,LS-PLASTIC) 0.9%	30 ML	VL	IV	ML		10 ML		0.1	06/01/2005	99/99/9999						
00641-6151-25		J1170		10/01/2018	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (PF,LATEX-FREE) 2 MG/1 ML	1 ML	VL	IJ	ML		4 MG		0.5	10/01/2018	99/99/9999						
00409-1985-05		J2060		02/08/2008	07/01/2012	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (10X1ML) 2 MG/ML	1 ML	SR	IJ	ML		2 MG		1	02/08/2008	07/01/2012						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00409-1985-30		J2060		06/01/2005	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (LUER LOCK,CARPUJECT) 2 MG/ML	1	ML	CR	IJ	ML	2	MG	1	06/01/2005	99/99/9999							
00409-2012-32		J0592		06/17/2005	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE (10X1ML,CARPUJECT) 0.3 MG/ML	1	ML	SR	IJ	ML	0.1	MG	3.24	06/17/2005	99/99/9999							
00409-2025-20		J1250		02/20/2006	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE (10X20ML) 12.5 MG/ML	20	ML	VL	IV	ML	250	MG	0.05	02/20/2006	99/99/9999							
00409-2025-54		J1250		11/10/2005	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE HCL (10X40ML) 12.5 MG/ML	40	ML	VL	IV	ML	250	MG	0.05	11/10/2005	99/99/9999							
00409-2043-02		J1245		03/31/2005	10/05/2016	INJECTION, DIPYRIDAMOLE, PER 10 MG	DIPYRIDAMOLE (AMIP,UNI-NEST,LATEX-FREE) 5 MG/ML	2	ML	AM	IV	ML	10	MG	0.5	03/31/2005	10/05/2016							
00409-2047-50		J0670		09/22/2006	99/99/9999	INJECTION, MEPIVACAINE HYDROCHLORIDE, PER 10 ML	CARBOCAINE (M.D.V. USP) 2%	50	ML	VL	IJ	ML	10	ML	0.1	09/22/2006	99/99/9999							
00409-2066-05		J2001		09/06/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (VIAL,LATEX-FREE) 2%	5	ML	VL	IJ	ML	10	ML	2	09/06/2005	99/99/9999							
00409-2102-02		A4216		01/01/2007	07/02/2013	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (25X2ML,PF) 0.9%	2	ML	VL	IV	ML	10	ML	0.1	01/01/2007	07/02/2013							
00409-2102-05		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (25X5ML,PF) 0.9%	5	ML	VL	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
00409-2168-02		J3475		01/31/2005	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (VIAL, FLIPTOP) 500 MG/ML	20	ML	VL	IJ	ML	500	MG	1	01/31/2005	99/99/9999							
00409-2265-01		J2597		02/04/2005	99/99/9999	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DESMOPRESSIN ACETATE (UNI-AMP) 4 MCG/ML	1	ML	AM	IJ	ML	1	MCG	4	02/04/2005	99/99/9999							
00409-2287-21		J1885		06/22/2007	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (10X1ML, USP) 30 MG/ML	1	ML	CT	IJ	ML	15	MG	2	06/22/2007	99/99/9999							
00409-2287-22		J1885		06/22/2007	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (10X2ML) 30 MG/ML	2	ML	CT	IJ	ML	15	MG	2	06/22/2007	99/99/9999							
00904-6745-61		Q0167		10/01/2018	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (USP,SOFT GELATIN) 2.5 MG	100	EA	ST	PO	EA	2.5	MG	1	10/01/2018	99/99/9999							
00904-6746-04		Q0167		10/01/2018	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (USP,SOFT GELATIN) 5 MG	30	EA	ST	PO	EA	2.5	MG	2	10/01/2018	99/99/9999							
00409-2287-31		J1885		04/25/2005	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (LUER LOCK,CARPUJECT) 30 MG/ML	1	ML	CR	IJ	ML	15	MG	2	04/25/2005	99/99/9999							
00409-2287-61		J1885		06/20/2005	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE ((LUER LOCK),10X2ML) 30 MG/ML	2	ML	SR	IM	ML	15	MG	2	06/20/2005	99/99/9999							
00409-2288-31		J1885		08/29/2005	03/01/2015	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (LUER LOCK,LATEX-FREE) 15 MG/ML	1	ML	SR	IJ	ML	15	MG	1	08/29/2005	03/01/2015							
00409-2290-31		J1200		04/25/2005	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (LUER LOCK,CARPUJECT) 50 MG/ML	1	ML	CR	IJ	ML	50	MG	1	04/25/2005	99/99/9999							
25021-0812-30		J0132		08/29/2018	99/99/9999	INJECTION, ACETYLCYSTEINE, 100 MG	ACETYLCYSTEINE (SDV,PF,LATEX-FREE) 200 MG/1 ML	30	ML	VL	IV	ML	100	MG	2	08/29/2018	99/99/9999							
00409-2305-02		J2250		08/15/2005	05/01/2012	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL, FLIPTOP,PF) 1 MG/ML	2	ML	VL	IJ	ML	1	MG	1	08/15/2005	05/01/2012							
00409-2305-05		J2250		12/21/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (PF) 1 MG/ML	5	ML	VL	IJ	ML	1	MG	1	12/21/2005	99/99/9999							
00409-2305-49		J2250		08/02/2005	06/20/2016	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL NOVATION (10X2ML,PF) 1 MG/ML	2	ML	VL	IJ	ML	1	MG	1	08/02/2005	06/20/2016							
00409-2305-50		J2250		09/13/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL NOVATION (FTV,10X5ML,PF) 1 MG/ML	5	ML	VL	IJ	ML	1	MG	1	09/13/2005	99/99/9999							
00409-2305-61		J2250		10/03/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL AMERINET CHOICE (VIAL,FLIPTOP,PF) 1 MG/ML	2	ML	VL	IJ	ML	1	MG	1	10/03/2005	99/99/9999							
00409-2305-62		J2250		10/03/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL AMERINET CHOICE (VIAL,FLIPTOP,PF) 1 MG/ML	5	ML	VL	IJ	ML	1	MG	1	10/03/2005	99/99/9999							
00409-2306-62		J2250		03/10/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (LUER LOCK,STERILE,PF) 1 MG/ML	2	ML	SR	IJ	ML	1	MG	1	03/10/2005	99/99/9999							
00409-2307-21		J2250		07/20/2007	06/01/2012	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HYDROCHLORIDE (10X1ML,PF) 5 MG/ML	1	ML	SR	IJ	ML	1	MG	5	07/20/2007	06/01/2012							
00409-2307-60		J2250		04/25/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (10X1ML,PF,CARPUJECT) 5 MG/ML	1	ML	CR	IJ	ML	1	MG	5	04/25/2005	99/99/9999							
00409-2308-01		J2250		06/07/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (10X1ML,PF) 5 MG/ML	1	ML	VL	IJ	ML	1	MG	5	06/07/2005	99/99/9999							
00409-2308-02		J2250		10/10/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,FLIPTOP,PF) 5 MG/ML	2	ML	VL	IJ	ML	1	MG	5	10/10/2005	99/99/9999							
00409-2308-49		J2250		12/29/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL NOVATION (FLIPTOP VIAL,PF) 5 MG/ML	1	ML	VL	IJ	ML	1	MG	5	12/29/2005	99/99/9999							
00409-2308-50		J2250		11/18/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL NOVATION (VIAL,FLIPTOP,PF) 5 MG/ML	2	ML	VL	IJ	ML	1	MG	5	11/18/2005	99/99/9999							
00409-2312-31		J2550		04/05/2005	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (LUER LOCK,CARPUJECT) 25 MG/ML	1	ML	SR	IJ	ML	50	MG	0.5	04/05/2005	99/99/9999							
00409-2336-10		J0895		04/25/2005	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE (LATEX-FREE) 500 MG	1	EA	VL	IJ	EA	500	MG	1	04/25/2005	99/99/9999							
00409-2337-25		J0895		03/21/2005	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE (LATEX-FREE) 2 GM	1	EA	VL	IJ	EA	500	MG	4	03/21/2005	99/99/9999							
00409-2344-01		J1250		07/27/2005	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE HCL (VIAL,FLIPTOP) 12.5 MG/ML	20	ML	VL	IV	ML	250	MG	0.05	07/27/2005	99/99/9999							
00409-2344-02		J1250		06/29/2005	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE (10X20ML,FTV) 12.5 MG/ML	20	ML	VL	IV	ML	250	MG	0.05	06/29/2005	99/99/9999							
00409-2344-88		J1250		03/21/2005	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE NOVAPLUS (S.D.V., U.S.P.) 12.5 MG/ML	20	ML	VL	IV	ML	250	MG	0.05	03/21/2005	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00078-0675-15		Q0162		03/20/2018	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFANR 4 MG	30	EA	BO	PO	EA	1	MG	4	03/20/2018	99/99/9999							
00409-2346-32		J1250		08/11/2005	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE IN DEXTROSE (12X250ML,LATEX-FREE) 5%-100 MG/100 ML	250	ML	FC	IV	ML	250	MG	0.004	08/11/2005	99/99/9999							
00409-2346-34		J1250		02/07/2006	10/05/2016	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE IN DEXTROSE (12X500ML,LIFECARE) 5%-100 MG/100 ML	500	ML	FC	IV	ML	250	MG	0.004	02/07/2006	10/05/2016							
00409-2347-32		J1250		01/11/2006	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DEXTROSE/DOBUTAMINE NOVAPLUS (U.S.P.) 5%-200 MG/100 ML	250	ML	FC	IV	ML	250	MG	0.008	01/11/2006	99/99/9999							
00409-2347-33		J1250		03/21/2005	02/01/2015	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE IN DEXTROSE (12X500ML,LIFECARE) 5%-100 MG/100 ML	250	ML	FC	IV	ML	250	MG	0.008	03/21/2005	02/01/2015							
00409-2349-31		J2560		09/07/2005	04/28/2016	INJECTION, PHENOBARBITAL SODIUM, UP TO 120 MG	LUMINAL SODIUM (LUSER LOCK CARPUJECT) 130 MG/ML	1	ML	SR	IJ	ML	120	MG	1.08333	09/07/2005	04/28/2016							
00409-2540-01		J1170		09/21/2005	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (USP,10X1ML) 4 MG/ML	1	ML	AM	IJ	ML	4	MG	1	09/21/2005	99/99/9999							
00409-2552-01		J1170		09/21/2005	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (USP,10X1ML) 1 MG/ML	1	ML	AM	IJ	ML	4	MG	0.25	09/21/2005	99/99/9999							
00409-2581-02		J1644		03/24/2006	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (ADD-VANTAGE VIAL) 2000 U/ML	5	ML	VL	IV	ML	1000	U	2	03/24/2006	99/99/9999							
00409-2584-02		J1644		07/01/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (25X10ML,PF,LATEX-FREE) 2500 U/ML	10	ML	VL	IJ	ML	1000	U	2.5	07/01/2005	99/99/9999							
00409-2585-01		J0690		06/27/2007	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN (SDV,ADD-VANTAGE) 1 GM	25	EA	VL	IV	EA	500	MG	2	06/27/2007	99/99/9999							
00409-2587-05		J2250		01/27/2006	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HYDROCHLORIDE (10X10ML,FLIPTOP/VIAL) 1 MG/ML	10	ML	VL	IJ	ML	1	MG	1	01/27/2006	99/99/9999							
00409-2587-53		J2250		03/07/2006	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	NOVAPLUS MIDAZOLAM HCL (10X10ML,FTV) 1 MG/ML	10	ML	VL	IJ	ML	1	MG	1	03/07/2006	99/99/9999							
00409-2596-03		J2250		10/28/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,FLIPTOP,LATEX-FREE) 5 MG/ML	5	ML	VL	IJ	ML	1	MG	5	10/28/2005	99/99/9999							
00409-2596-05		J2250		01/11/2006	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL, FLIPTOP) 5 MG/ML	10	ML	VL	IJ	ML	1	MG	5	01/11/2006	99/99/9999							
00409-2596-52		J2250		01/23/2006	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	NOVAPLUS MIDAZOLAM HYDROCHLORIDE (10X5ML) 5 MG/ML	5	ML	VL	IJ	ML	1	MG	5	01/23/2006	99/99/9999							
00409-2596-53		J2250		09/27/2005	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL NOVATION (FTV,10X10ML,LATEX-FREE) 5 MG/ML	10	ML	VL	IJ	ML	1	MG	5	09/27/2005	99/99/9999							
00409-2687-15		J0295		06/22/2007	06/01/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM 10 GM-5 GM	1	EA	VL	IV	EA	1.5	GM	10	06/22/2007	06/01/2013							
00409-2689-11		J0295		07/01/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	NOVAPLUS AMPICILLIN AND SULBACTAM (USP,ADD-VANTAGE) 1 GM-0.5 GM	1	EA	VL	IV	EA	1.5	GM	1	07/01/2007	99/99/9999							
00409-2776-02		J2260		03/08/2006	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (IN 5% DEXTROSE,10X200ML) 5%-20 MG/100 ML	200	ML	FC	IV	ML	5	MG	0.04	03/08/2006	99/99/9999							
00409-2776-23		J2260		06/15/2005	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	DEXTROSE/MILRINONE LACTATE (10X100ML,LATEX-FREE) 5%-20 MG/100 ML	100	ML	FC	IV	ML	5	MG	0.04	06/15/2005	99/99/9999							
00409-2987-13		J0295		07/01/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	NOVAPLUS AMPICILLIN AND SULBACTAM (USP,ADD-VANTAGE) 2 GM-1 GM	1	EA	VL	IV	EA	1.5	GM	2	07/01/2007	99/99/9999							
00409-2988-01		J0295		07/20/2007	10/01/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP) 1 GM-0.5 GM	10	EA	VL	IJ	EA	1.5	GM	1	07/20/2007	10/01/2013							
00409-2998-03		J0295		07/20/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP) 2 GM-1 GM	10	EA	VL	IJ	EA	1.5	GM	2	07/20/2007	99/99/9999							
25021-0231-20		J0894		09/07/2018	99/99/9999	INJECTION, DECITABINE, 1 MG	DECITABINE (PF,LATEX-FREE) 50 MG	1	EA	VL	IV	EA	1	MG	50	09/07/2018	99/99/9999							
00409-3213-12		J3360		10/01/2007	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (10X10ML,USP,MDV,FLIPTOP) 5 MG/ML	10	ML	VL	IJ	ML	5	MG	1	10/01/2007	99/99/9999							
00409-3307-03		J7608		04/11/2005	99/99/9999	ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCHOLINE 10%	30	ML	VL	IH	ML	1	GM	0.1	04/11/2005	99/99/9999							
25021-0408-51		J1327		09/17/2018	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (PF,LATEX-FREE) 0.75 MG/1 ML	100	ML	VL	IV	ML	5	MG	0.15	09/17/2018	99/99/9999							
25021-0409-10		J1327		09/17/2018	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (PF,LATEX-FREE) 2 MG/1 ML	10	ML	VL	IV	ML	5	MG	0.4	09/17/2018	99/99/9999							
25021-0783-05		J2469		09/19/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL (PF,LATEX-FREE) 0.05 MG/1 ML	5	ML	VL	IV	ML	25	MCG	2	09/19/2018	99/99/9999							
43598-0635-10		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (10X100ML) 5 MG/1 ML	100	ML	BG	IV	ML	10	MG	0.5	06/13/2018	99/99/9999							
00409-3307-03	KO	J7608	KO	04/11/2005	99/99/9999	ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCHOLINE 10%	30	ML	VL	IH	ML	1	GM	0.1	04/11/2005	99/99/9999							
00409-3308-03		J7608		05/25/2005	99/99/9999	ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCHOLINE (3X30ML) 20%	30	ML	VL	IH	ML	1	GM	0.2	05/25/2005	99/99/9999							
00409-3308-03	KO	J7608	KO	05/25/2005	99/99/9999	ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCHOLINE (3X30ML) 20%	30	ML	VL	IH	ML	1	GM	0.2	05/25/2005	99/99/9999							
00409-3356-01		J1170		09/21/2005	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (10X1ML,USP) 2 MG/ML	1	ML	AM	IJ	ML	4	MG	0.5	09/21/2005	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-3365-01		J1170		09/21/2005	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (SDV,25X1ML) 2 MG/ML	1	ML	VL	IJ	ML	4	MG	0.5	09/21/2005	99/99/9999						
00409-3380-31		J3490		09/01/2005	11/03/2013	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (LATEX-FREE) 50 MCG/ML	1	ML	AM	IJ	ML	1	EA	1	09/01/2005	11/03/2013						
00409-3380-32		J3490		11/03/2005	08/01/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (AMP,10X2ML,LATEX-FREE) 50 MCG/ML	2	ML	AM	IJ	ML	1	EA	1	11/03/2005	08/01/2015						
00409-3380-35		J3490		12/28/2005	08/01/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (AMP,LATEX-FREE) 50 MCG/ML	5	ML	AM	IJ	ML	1	EA	1	12/28/2005	08/01/2015						
00409-3380-49		J3490		11/29/2005	02/23/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE NOVAPLUS (AMP,PF,LATEX-FREE) 50 MCG/ML	1	ML	AM	IJ	ML	1	EA	1	11/29/2005	02/23/2015						
00409-3380-50		J3490		11/07/2005	02/23/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE NOVAPLUS (10X2ML,PF,LATEX-FREE) 50 MCG/ML	2	ML	AM	IJ	ML	1	EA	1	11/07/2005	02/23/2015						
00409-3380-51		J3490		10/12/2005	02/23/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE NOVAPLUS (AMP,10X5ML,PF) 50 MCG/ML	5	ML	AM	IJ	ML	1	EA	1	10/12/2005	02/23/2015						
00409-3382-21		J3490		07/15/2005	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (10X1ML,LATEX-FREE) 50 MCG/ML	1	ML	VL	IJ	ML	1	EA	1	07/15/2005	99/99/9999						
00409-3382-22		J3490		07/18/2005	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (10X2ML,LATEX-FREE) 50 MCG/ML	2	ML	VL	IJ	ML	1	EA	1	07/18/2005	99/99/9999						
00409-3382-25		J3490		10/19/2005	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (USP,10X5ML) 50 MCG/ML	5	ML	VL	IJ	ML	1	EA	1	10/19/2005	99/99/9999						
00409-3400-01		J1580		03/24/2006	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (25X5ML,ADD-VANTAGE) 10 MG/ML	6	ML	VL	IV	ML	80	MG	0.125	03/24/2006	99/99/9999						
00409-3401-01		J1580		01/09/2006	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (VIAL-ADD-VANTAGE) 10 MG/ML	8	ML	VL	IJ	ML	80	MG	0.125	01/09/2006	99/99/9999						
00409-3402-01		J1580		06/05/2006	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (SD ADD-VANTGE,USP) 10 MG/ML	10	ML	VL	IV	ML	80	MG	0.125	06/05/2006	99/99/9999						
00409-3470-23		J3260		09/26/2005	04/01/2014	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	SODIUM CHLORIDE/TOBAMYCIN SULFATE (PREMX,24X100ML) 0.9%-80 MG/100 ML	100	ML	FC	IV	ML	80	MG	0.01	09/26/2005	04/01/2014						
00409-3577-01		J3260		03/31/2005	02/01/2016	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (VIAL,FLIPTOP,LATEX-FREE) 10 MG/ML	2	ML	VL	IJ	ML	80	MG	0.125	03/31/2005	02/01/2016						
00409-3578-01		J3260		11/02/2004	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (VIAL,FLIPTOP) 40 MG/ML	2	ML	VL	IJ	ML	80	MG	0.5	11/02/2004	99/99/9999						
00409-3590-02		J3260		02/15/2006	07/01/2012	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (BULK PACKAGE) 40 MG/ML	50	ML	VL	IJ	ML	80	MG	0.5	02/15/2006	07/01/2012						
00409-3613-01		J3490		01/07/2005	99/99/9999	UNCLASSIFIED DRUGS	BUPIVACAINE SPINAL AMPUL (AMP,LATEX-FREE) 0.25%	2	ML	AM	IJ	ML	1	EA	1	01/07/2005	99/99/9999						
00409-3724-32		J1250		10/07/2005	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DEXTROSE/DOBUTAMINE (LATEX-FREE) 5%-400 MG/100 ML	250	ML	FC	IV	ML	250	MG	0.016	10/07/2005	99/99/9999						
00409-3793-01		J1885		05/31/2005	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP,FLIPTOP VIAL) 15 MG/ML	1	ML	VL	IJ	ML	15	MG	1	05/31/2005	99/99/9999						
00409-3793-49		J1885		04/19/2005	04/01/2016	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE NOVAPLUS (U.S.P.,25X1ML) 15 MG/ML	1	ML	VL	IJ	ML	15	MG	1	04/19/2005	04/01/2016						
00409-3795-01		J1885		01/06/2006	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (LATEX-FREE) 30 MG/ML	1	ML	VL	IJ	ML	15	MG	2	01/06/2006	99/99/9999						
00409-3795-49		J1885		09/21/2005	04/01/2016	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE NOVATION (FTV,25X1ML,2ML VIAL) 30 MG/ML	1	ML	VL	IJ	ML	15	MG	2	09/21/2005	04/01/2016						
00409-3796-01		J1885		12/21/2005	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (VIAL, FLIPTOP) 30 MG/ML	2	ML	VL	IM	ML	15	MG	2	12/21/2005	99/99/9999						
00409-3796-49		J1885		11/07/2005	02/01/2016	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE NOVATION (FTV,25X2ML,LATEX-FREE) 30 MG/ML	2	ML	VL	IM	ML	15	MG	2	11/07/2005	02/01/2016						
00409-3814-12		J2275		07/19/2005	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (5X10ML,PF,LATEX-FREE) 0.5 MG/ML	10	ML	VL	IJ	ML	10	MG	0.05	07/19/2005	12/31/2014						
67877-0266-01		J7517		08/01/2013	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	100	EA	BO	PO	EA	250	MG	1	08/01/2013	99/99/9999						
00409-3977-03		A4216		04/07/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION BACTERIOSTATIC (VIAL,FLIPTOP,LATEX-FREE)	30	ML	VL	IV	ML	10	ML	0.1	04/07/2005	99/99/9999						
00409-4029-03		A4216		03/01/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (AMP,PF,LATEX-FREE)	20	ML	AM	IV	ML	10	ML	0.1	03/01/2005	99/99/9999						
00409-4031-01		J2150		10/19/2004	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (VIAL, FLIPTOP) 25%	50	ML	VL	IV	ML	50	ML	0.02	10/19/2004	99/99/9999						
00409-4044-02		A4216		02/09/2006	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (25X10ML,PF,LATEX-FREE)	10	ML	AM	IV	ML	10	ML	0.1	02/09/2006	99/99/9999						
00409-4050-01		J3490		05/13/2005	09/02/2015	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (VIAL,FLIPTOP,LATEX-FREE) 150 MG/ML	2	ML	VL	IJ	ML	1	EA	1	05/13/2005	09/02/2015						
00409-4051-01		J3490		05/31/2005	09/02/2015	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (VIAL,FLIPTOP,LATEX-FREE) 150 MG/ML	4	ML	VL	IJ	ML	1	EA	1	05/31/2005	09/02/2015						
00409-4052-01		J3490		07/05/2005	09/02/2015	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (25X5ML,LATEX-FREE) 150 MG/ML	6	ML	VL	IJ	ML	1	EA	1	07/05/2005	09/02/2015						
00409-4053-03		J3490		05/11/2005	09/02/2015	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (ADD-VANTAGE,25X2ML) 150 MG/ML	2	ML	VL	IJ	ML	1	EA	1	05/11/2005	09/02/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00409-4054-03		J3490		02/18/2005	09/02/2015	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (VIAL,ADVANTAGE) 150 MG/ML	4 ML	VL	IJ	ML		1 EA		1	02/18/2005	09/02/2015							
00409-4055-03		J3490		02/24/2005	09/02/2015	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (VIAL,ADVANTAGE) 150 MG/ML	6 ML	VL	IJ	ML		1 EA		1	02/24/2005	09/02/2015							
00409-4056-01		J2001		10/31/2005	11/01/2015	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (AMP,PF) 1.5%	20 ML	AM	IJ	ML		10 MG		1.5	10/31/2005	11/01/2015							
00409-4057-12		J2275		12/13/2005	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (PF,LATEX-FREE) 0.5 MG/ML	5 ML	AM	IJ	ML		10 MG		0.05	12/13/2005	12/31/2014							
00409-4089-02		J7799		05/18/2005	06/08/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (AMP,LATEX-FREE) 10%	5 ML	AM	IV	ML		1 EA		1	05/18/2005	06/08/2016							
00409-4169-01		J2400		06/20/2005	01/01/2013	INJECTION, CHLOROPROCAINE HYDROCHLORIDE, PER 30 ML	CHLOROPROCAINE HCL (25X30ML) 2% CHLOROPROCAINE HCL (VIAL,25X30ML) 3%	30 ML	VL	IJ	ML		30 ML		0.03333	06/20/2005	01/01/2013							
00409-4170-01		J2400		04/20/2005	07/01/2013	INJECTION, CHLOROPROCAINE HYDROCHLORIDE, PER 30 ML	CHLOROPROCAINE HCL (VIAL,25X30ML) 3%	30 ML	VL	IJ	ML		30 ML		0.03333	04/20/2005	07/01/2013							
00409-4197-01		J3490		03/31/2005	09/02/2015	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (VIAL,BULK,LATEX-FREE) 150 MG/ML	60 ML	VL	IJ	ML		1 EA		1	03/31/2005	09/02/2015							
00409-4219-02		J7799		03/30/2005	09/03/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE 2.5%	250 ML	GC	IV	ML		1 EA		1	03/30/2005	09/03/2016							
00409-4265-01		J1265		01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HCL (25X10ML) 80 MG/ML LEVETIRACETAM (1X100ML, INNER PACK) 5 MG/1 ML	10 ML	VL	IV	ML		40 MG		2	01/01/2006	99/99/9999							
43598-0635-52		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (1X100ML, INNER PACK) 5 MG/1 ML	100 ML	BG	IV	ML		10 MG		0.5	06/13/2018	99/99/9999							
00409-4270-01		J2001		02/27/2006	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (STERILE PACK,SDV) 1%	30 ML	VL	EP	ML		10 MG		1	02/27/2006	99/99/9999							
00409-4272-01		J3490		04/06/2006	02/01/2015	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (AMP,STERILE,USP,5X20ML) 0.25% BUPIVACAINE HYDROCHLORIDE (SINGLE-DOSE,5X20ML,PF) 0.5%	20 ML	AM	IJ	ML		1 EA		1	04/06/2006	02/01/2015							
00409-4273-01		J3490		06/28/2006	10/01/2015	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (AMP,STERILE,USP,5X20ML) 0.75% LIDOCAINE HCL (VIAL,FLIPTOP) 0.5%	20 ML	AM	IJ	ML		1 EA		1	06/28/2006	10/01/2015							
00409-4274-01		J3490		03/31/2006	08/05/2016	UNCLASSIFIED DRUGS	BUPIVACAINE HCL (AMP,STERILE,USP,5X20ML) 0.5%	20 ML	AM	IJ	ML		1 EA		1	03/31/2006	08/05/2016							
00409-4275-01		J2001		12/30/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (VIAL,FLIPTOP) 0.5%	50 ML	VL	IJ	ML		10 MG		0.5	12/30/2005	99/99/9999							
00409-4276-01		J2001		08/12/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (FTV,25X20ML) 1%	20 ML	VL	EP	ML		10 MG		1	08/12/2005	99/99/9999							
00409-4276-02		J2001		07/07/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (25X50ML) 1% LIDOCAINE HCL (25X20ML,LATEX-FREE) 2%	50 ML	VL	EP	ML		10 MG		1	07/07/2005	99/99/9999							
00409-4277-01		J2001		06/13/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (FTV,25X50ML,LATEX-FREE) 2%	20 ML	VL	IJ	ML		10 MG		2	06/13/2005	99/99/9999							
00409-4277-02		J2001		08/12/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (FTV,25X50ML,LATEX-FREE) 2%	50 ML	VL	IJ	ML		10 MG		2	08/12/2005	99/99/9999							
00409-4278-01		J2001		06/29/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (25X50ML) 0.5%	50 ML	VL	IJ	ML		10 MG		0.5	06/29/2005	99/99/9999							
00409-4279-02		J2001		08/31/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (TEARDROP BOTTLE) 1% LIDOCAINE HCL (AMP,25X2ML,LATEX-FREE) 2%	30 ML	VL	EP	ML		10 MG		1	08/31/2005	99/99/9999							
00409-4282-01		J2001		09/09/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HYDROCHLORIDE (USP,25X10ML,SDA,PF) 2% LIDOCAINE HCL (AMP,LATEX-FREE) 4%	2 ML	AM	IJ	ML		10 MG		2	09/09/2005	99/99/9999							
00409-4282-02		J2001		02/08/2006	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (AMP,LATEX-FREE) 4%	10 ML	AM	IJ	ML		10 MG		2	02/08/2006	99/99/9999							
00409-4283-01		J2001		05/16/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	VANCOMYCIN HCL (VIAL,FLIPTOP) 500 MG VANCOMYCIN HCL NOVATION (FTV,LATEX-FREE) 500 MG	5 ML	AM	IJ	ML		10 MG		4	05/16/2005	99/99/9999							
00409-4332-01		J3370		04/25/2005	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (VIAL,FLIPTOP) 500 MG	1 EA	VL	IV	EA		500 MG		1	04/25/2005	99/99/9999							
00409-4332-49		J3370		08/04/2005	01/01/2016	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL NOVATION (FTV,LATEX-FREE) 500 MG AMINOCAPROIC ACID (VIAL,FLIPTOP) 250 MG/ML	1 EA	VL	IV	EA		500 MG		1	08/04/2005	01/01/2016							
00409-4346-73		J3490		04/13/2005	99/99/9999	UNCLASSIFIED DRUGS	AMIODARONE HYDROCHLORIDE (3MLX10,SINGLE-DOSE) 50 MG/ML FLUCONAZOLE (6X200ML,LATEX-FREE) 400 MG/200 ML	20 ML	VL	IV	ML		1 EA		1	04/13/2005	99/99/9999							
00409-4348-35		J0282		09/27/2006	08/01/2015	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HYDROCHLORIDE (3MLX10,SINGLE-DOSE) 50 MG/ML FLUCONAZOLE (6X200ML,LATEX-FREE) 400 MG/200 ML	3 ML	AM	IV	ML		30 MG		1.66666	09/27/2006	08/01/2015							
00409-4684-02		J1450		03/06/2007	09/01/2015	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (6X100ML) 200 MG/100 ML	200 ML	FC	IV	ML		200 MG		0.01	03/06/2007	09/01/2015							
00409-4684-23		J1450		04/14/2006	11/17/2016	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (6X200ML) 400 MG/200 ML FLUCONAZOLE (6X100ML,LATEX FREE) 200 MG/100 ML	100 ML	FC	IV	ML		200 MG		0.01	04/14/2006	11/17/2016							
00409-4688-02		J1450		07/27/2006	11/01/2016	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (6X200ML) 400 MG/200 ML FLUCONAZOLE (6X100ML,LATEX FREE) 200 MG/100 ML	200 ML	FC	IV	ML		200 MG		0.01	07/27/2006	11/01/2016							
00409-4688-23		J1450		06/16/2006	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	AMERINET CHOICE FLUCONAZOLE (100MLX6,LATEX-FREE) 200 MG/100 ML NOVAPLUS FLUCONAZOLE (6X100ML,LATEX-FREE) 200 MG/100 ML	100 ML	FC	IV	ML		200 MG		0.01	06/16/2006	99/99/9999							
00409-4688-27		J1450		05/27/2006	06/10/2013	INJECTION FLUCONAZOLE, 200 MG	AMERINET CHOICE FLUCONAZOLE (6X200ML,LATEX-FREE) 200 MG/100 ML NOVAPLUS FLUCONAZOLE (6X100ML,LATEX-FREE) 200 MG/100 ML	100 ML	FC	IV	ML		200 MG		0.01	05/27/2006	06/10/2013							
00409-4688-28		J1450		06/01/2005	12/01/2015	INJECTION FLUCONAZOLE, 200 MG	AMERINET CHOICE FLUCONAZOLE (6X200ML,LATEX-FREE) 200 MG/100 ML NOVAPLUS FLUCONAZOLE (6X100ML,LATEX-FREE) 200 MG/100 ML	100 ML	FC	IV	ML		200 MG		0.01	06/01/2005	12/01/2015							
00409-4688-33		J1450		10/25/2006	06/10/2013	INJECTION FLUCONAZOLE, 200 MG	AMERINET CHOICE FLUCONAZOLE (6X200ML,LATEX-FREE) 200 MG/100 ML NOVAPLUS FLUCONAZOLE (6X100ML,LATEX-FREE) 200 MG/100 ML	200 ML	FC	IV	ML		200 MG		0.01	10/25/2006	06/10/2013							
00409-4688-34		J1450		03/02/2006	02/01/2016	INJECTION FLUCONAZOLE, 200 MG	AMERINET CHOICE FLUCONAZOLE (6X200ML,LATEX-FREE) 200 MG/100 ML NOVAPLUS FLUCONAZOLE (6X100ML,LATEX-FREE) 200 MG/100 ML	200 ML	FC	IV	ML		200 MG		0.01	03/02/2006	02/01/2016							
00409-4699-24		J3490		03/22/2006	99/99/9999	UNCLASSIFIED DRUGS	PROPOFOL (FLIPTOP VIAL) 10 MG/ML	100 ML	VL	IV	ML		1 EA		1	03/22/2006	99/99/9999							
00409-4699-30		J3490		03/22/2006	99/99/9999	UNCLASSIFIED DRUGS	PROPOFOL (FLIPTOP VIAL) 10 MG/ML	20 ML	VL	IV	ML		1 EA		1	03/22/2006	99/99/9999							
00409-4699-33		J3490		03/22/2006	99/99/9999	UNCLASSIFIED DRUGS	PROPOFOL (FLIPTOP VIAL) 10 MG/ML	50 ML	VL	IV	ML		1 EA		1	03/22/2006	99/99/9999							
00409-4699-61		J3490		12/01/2007	08/26/2014	UNCLASSIFIED DRUGS	AMERINET CHOICE PROPOFOL (5X20ML,SDV,PF) 10 MG/ML LIDOCAINE HCL (25X5ML,LATEX-FREE) 1%	20 ML	VL	IV	ML		1 EA		1	12/01/2007	08/26/2014							
00409-4713-02		J2001		11/21/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (AMP,LATEX-FREE) 1%	5 ML	AM	EP	ML		10 MG		1	11/21/2005	99/99/9999							
00409-4713-32		J2001		09/06/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (LATEX-FREE) 1% ONDANSETRON (SINGLEDOSE,USP,10X2ML) 2 MG/ML	2 ML	AM	EP	ML		10 MG		1	09/06/2005	99/99/9999							
00409-4755-02		J2405		08/24/2007	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (SINGLEDOSE,USP,10X2ML) 2 MG/ML	2 ML	VL	IJ	ML		1 MG		2	08/24/2007	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00409-4755-61	J2405			12/26/2006	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	AMERINET CHOICE ONDANSETRON (5X2ML,SDV,USP) 2 MG/ML	2	ML	VL	IJ	ML	1	MG	2	12/26/2006	99/99/9999							
00409-4759-01	J2405			12/26/2006	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV,USP) 2 MG/ML	20	ML	VL	IJ	ML	1	MG	2	12/26/2006	99/99/9999							
00409-4765-86	J0744			08/29/2006	08/01/2015	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	CIPROFLOXACIN (SINGLE-DOSE,USP) 10 MG/ML	20	ML	VL	IV	ML	200	MG	0.05	08/29/2006	08/01/2015							
00409-4776-01	J2001			02/06/2006	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HYDROCHLORIDE (25X20ML,PF) 1.5% CIPROFLOXACIN (24X200ML,SINGLEDSE,USP) 400 MG/200 ML	20	ML	AM	IJ	ML	10	MG	1.5	02/06/2006	99/99/9999							
00409-4777-02	J0744			03/19/2008	99/99/9999	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	AMERINET CHOICE CIPROFLOXACIN (24X100ML,SINGLEDSE,USP) 200 MG/100 ML	200	ML	FC	IV	ML	200	MG	0.01	03/19/2008	99/99/9999							
00069-1305-10	Q5106			01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 2000 U/1 ML CIPROFLOXACIN (24X100ML,SINGLEDSE,USP) 200 MG/100 ML	1	ML	VL	IJ	ML	1000	U	2	01/01/2019	99/99/9999							
00409-4777-23	J0744			03/19/2008	99/99/9999	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	AMERINET CHOICE CIPROFLOXACIN (24X100ML,SINGLEDSE,USP) 200 MG/100 ML	100	ML	FC	IV	ML	200	MG	0.01	03/19/2008	99/99/9999							
00409-4777-61	J0744			05/19/2008	99/99/9999	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	AMERINET CHOICE CIPROFLOXACIN (24X100ML,SINGLEDSE,USP) 200 MG/100 ML	100	ML	FC	IV	ML	200	MG	0.01	05/19/2008	99/99/9999							
00409-4856-05	J1720			06/27/2006	99/99/9999	INJECTION, HYDROCORTISONE SODIUM SUCCINATE, UP TO 100 MG	A-HYDROCORT (SINGLE-DOSE) 100 MG	10	EA	VL	IJ	EA	100	MG	1	06/27/2006	99/99/9999							
00378-9691-52	J7614			07/23/2018	99/99/9999	COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.63 MG/3 ML DEXTRROSE/SODIUM CHLORIDE 10%-0.225%	3	ML	VL	IH	ML	0.5	MG	0.42	07/23/2018	99/99/9999							
00409-4862-02	J7799			03/09/2005	05/18/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE/SODIUM CHLORIDE 10%-0.225%	250	ML	GC	IV	ML	1	EA	1	03/09/2005	05/18/2016							
00409-4862-03	J7799			04/04/2005	05/18/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE/SODIUM CHLORIDE 10%-0.225%	500	ML	GC	IV	ML	1	EA	1	04/04/2005	05/18/2016							
00409-4887-10	A4216			08/18/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (FTV,25X100ML,PF)	10	ML	VL	IV	ML	10	ML	0.1	08/18/2005	99/99/9999							
00409-4887-20	A4216			06/16/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (25X20ML,STERILE,PF)	20	ML	VL	IV	ML	10	ML	0.1	06/16/2005	99/99/9999							
00409-4887-50	A4216			08/05/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (FTV,25X50ML,PF)	50	ML	VL	IV	ML	10	ML	0.1	08/05/2005	99/99/9999							
00409-4887-99	A4216			08/03/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (FTV,25X100ML,PF)	100	ML	VL	IV	ML	10	ML	0.1	08/03/2005	99/99/9999							
00409-4888-10	A4216			04/22/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (VIAL,FLIPTOP,ADDITIVE) 0.9% SODIUM CHLORIDE (25X10ML,PF,LATEX-FREE) 0.9%	10	ML	VL	IV	ML	10	ML	0.1	04/22/2005	99/99/9999							
00409-4888-12	A4216			07/15/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (VIAL, FLIPTOP, ADDITIVE) 0.9%	10	ML	VL	IV	ML	10	ML	0.1	07/15/2005	99/99/9999							
00409-4888-20	A4216			02/23/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (VIAL,FLIPTOP,ADDITIVE) 0.9%	20	ML	VL	IV	ML	10	ML	0.1	02/23/2005	99/99/9999							
00409-4888-50	A4216			02/14/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (VIAL,FLIPTOP,ADDITIVE) 0.9%	50	ML	VL	IV	ML	10	ML	0.1	02/14/2005	99/99/9999							
00409-4902-34	J7799			12/08/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (LIFESHEILD, 18G1-1/2) 50% LIDOCAINE HCL (21GX1-1/2',LATEX-FREE) 2%	1	ML	SR	IV	ML	1	EA	1	12/08/2005	99/99/9999							
00409-4903-34	J2001			12/01/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (10X5ML,LATEX-FREE) 1%	5	ML	SR	IJ	ML	10	MG	2	12/01/2005	99/99/9999							
00409-4904-34	J2001			08/23/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (10X5ML,LATEX-FREE) 1%	5	ML	SR	EP	ML	10	MG	1	08/23/2005	99/99/9999							
00409-5082-16	J0713			10/24/2005	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	TAZICEF (LATEX-FREE) 1 GM	1	EA	VL	IJ	EA	500	MG	2	10/24/2005	99/99/9999							
00409-5082-52	J0713			10/04/2005	03/01/2016	INJECTION, CEFTAZIDIME, PER 500 MG	NOVAPLUS TAZICEF 1 GM	1	EA	VL	IJ	EA	500	MG	2	10/04/2005	03/01/2016							
00409-5084-11	J0713			12/05/2005	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	TAZICEF 2 GM	1	EA	VL	IJ	EA	500	MG	4	12/05/2005	99/99/9999							
00409-5084-51	J0713			10/04/2005	11/01/2015	INJECTION, CEFTAZIDIME, PER 500 MG	NOVAPLUS TAZICEF 2 GM	1	EA	VL	IJ	EA	500	MG	4	10/04/2005	11/01/2015							
00409-5086-11	J0713			04/19/2006	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	TAZICEF (BULK PHARMACY) 6 GM NOVAPLUS TAZICEF (BULK PACKAGE) 6 GM	1	EA	VL	IV	EA	500	MG	12	04/19/2006	99/99/9999							
00409-5086-51	J0713			10/04/2005	03/24/2016	INJECTION, CEFTAZIDIME, PER 500 MG	TAZICEF (SINGLE-DOSE ADD-VANTAGE) 1 GM	1	EA	VL	IJ	EA	500	MG	12	10/04/2005	03/24/2016							
00409-5092-16	J0713			05/02/2006	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	NOVAPLUS TAZICEF 1 GM	1	EA	VL	IJ	EA	500	MG	2	05/02/2006	99/99/9999							
00409-5092-52	J0713			06/27/2006	04/22/2016	INJECTION, CEFTAZIDIME, PER 500 MG	NOVAPLUS TAZICEF 1 GM	1	EA	VL	IJ	EA	500	MG	2	06/27/2006	04/22/2016							
00409-5093-11	J0713			04/03/2006	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	TAZICEF (ADD-VANTAGE,USP) 2 GM NOVAPLUS TAZICEF (ADD-VANTAGE) 2 GM	1	EA	VL	IJ	EA	500	MG	4	04/03/2006	99/99/9999							
00409-5093-51	J0713			10/01/2006	10/30/2014	INJECTION, CEFTAZIDIME, PER 500 MG	NOVAPLUS TAZICEF (ADD-VANTAGE) 2 GM	1	EA	VL	IJ	EA	500	MG	4	10/01/2006	10/30/2014							
00409-5684-01	J2920			11/01/2005	09/22/2016	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 40 MG	A-METHAPRED (UNIVIAL,LATEX-FREE) 40 MG	1	EA	VL	IJ	EA	40	MG	1	11/01/2005	09/22/2016							
00409-5685-02	J2930			11/01/2005	10/17/2016	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	A-METHAPRED (UNIVIAL,LATEX-FREE) 125 MG	1	EA	VL	IJ	EA	125	MG	1	11/01/2005	10/17/2016							
00409-5820-01	J1265			01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HCL (FLIPTOP) 40 MG/ML AMINOPHYLLINE (VIAL,FLIPTOP,25X10ML) 25 MG/ML	5	ML	VL	IV	ML	40	MG	1	01/01/2006	99/99/9999							
00409-5921-01	J0280			04/25/2005	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE (VIAL,FLIPTOP,ABJECT) 25 MG/ML MORPHINE SULFATE (SDV,30MLX10) 5 MG/ML	10	ML	VL	IV	ML	250	MG	0.1	04/25/2005	99/99/9999							
00409-5922-01	J0280			12/24/2004	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	MORPHINE SULFATE (SDV,30MLX10) 5 MG/ML	20	ML	VL	IV	ML	250	MG	0.1	12/24/2004	99/99/9999							
00409-6028-04	J2271			03/23/2007	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (SDV,30MLX10) 5 MG/ML	30	ML	VL	IV	ML	100	MG	0.05	03/23/2007	12/31/2014							
00409-6030-04	J2175			01/02/2007	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HYDROCHLORIDE (SDV,USP,10X30ML) 10 MG/ML	30	ML	VL	IV	ML	100	MG	0.1	01/02/2007	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00409-6062-02		J2270		01/10/2006	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE IN 5% DEXTROSE (PREMIX) 5%-100 MG/100 ML	250	ML	GC	IV	ML	10	MG	0.1	01/10/2006	99/99/9999							
00409-6102-02		J1940		02/18/2005	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (VIAL,FLIPTOP,ABBOJECT) 10 MG/ML	2	ML	VL	IJ	ML	20	MG	0.5	02/18/2005	99/99/9999							
43598-0636-10		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (10X100ML) 10 MG/1 ML	100	ML	BG	IV	ML	10	MG	1	06/13/2018	99/99/9999							
43598-0636-52		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (1X100ML, INNER PACK) 10 MG/1 ML	100	ML	BG	IV	ML	10	MG	1	06/13/2018	99/99/9999							
00409-6102-04		J1940		02/21/2005	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (VIAL,FLIPTOP,ABBOJECT) 10 MG/ML	4	ML	VL	IJ	ML	20	MG	0.5	02/21/2005	99/99/9999							
00409-6102-10		J1940		03/24/2005	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (VIAL,FLIPTOP,ABBOJECT) 10 MG/ML	10	ML	VL	IJ	ML	20	MG	0.5	03/24/2005	99/99/9999							
00409-6138-03		A4217		06/01/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (USP,AQUALITE,PF) 0.9%	500	ML	PC	IR	ML	500	ML	0.002	06/01/2005	99/99/9999							
00409-6138-22		A4217		09/01/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (AQUALITE, 24X250ML,PF) 0.9%	250	ML	PC	IR	ML	500	ML	0.002	09/01/2005	99/99/9999							
00409-6139-03		A4217		05/09/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (AQUALITE, U.S.P.)	500	ML	PC	IR	ML	500	ML	0.002	05/09/2005	99/99/9999							
00409-6139-22		A4217		05/04/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (AQUALITE, U.S.P.)	250	ML	PC	IR	ML	500	ML	0.002	05/04/2005	99/99/9999							
00409-6177-14		J2270		07/14/2005	99/99/9999	MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (ADD-VANTAGE, 10X4ML) 25 MG/ML	4	ML	VL	IJ	ML	10	MG	2.5	07/14/2005	99/99/9999							
00409-6179-14		J2270		09/01/2005	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (ADD-VANTAGE,LATEX-FREE) 25 MG/ML	10	ML	VL	IJ	ML	10	MG	2.5	09/01/2005	99/99/9999							
00409-6476-44		J1364		03/10/2006	99/99/9999	INJECTION, ERYTHROMYCIN LACTOBIONATE, PER 500 MG	ERYTHROCIIN LACTOBIONATE (ADD-VANTAGE VIAL,PF) 500 MG	1	EA	VL	IV	EA	500	MG	1	03/10/2006	99/99/9999							
00409-6478-44		J1364		01/10/2007	99/99/9999	INJECTION, ERYTHROMYCIN LACTOBIONATE, PER 500 MG	ERYTHROCIIN LACTOBIONATE (ADD-VANTAGE VIAL) 1 GM	1	EA	VL	IV	EA	500	MG	2	01/10/2007	99/99/9999							
00409-6482-01		J1364		05/23/2005	99/99/9999	INJECTION, ERYTHROMYCIN LACTOBIONATE, PER 500 MG	ERYTHROCIIN LACTOBIONATE (LATEX-FREE) 500 MG	1	EA	VL	IV	EA	500	MG	1	05/23/2005	99/99/9999							
00409-6509-01		J3370		06/06/2005	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (BULK,LATEX-FREE) 5 GM	1	EA	VL	IV	EA	500	MG	10	06/06/2005	99/99/9999							
00409-6509-49		J3370		06/03/2005	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL NOVAPLUS (BULK) 5 GM	1	EA	VL	IV	EA	500	MG	10	06/03/2005	99/99/9999							
00409-6533-01		J3370		03/15/2005	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (VIAL,FLIPTOP,LATEX-FREE) 1 GM	1	EA	VL	IV	EA	500	MG	2	03/15/2005	99/99/9999							
00409-6533-49		J3370		04/06/2005	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL NOVATION (VIAL,FLIPTOP,LATEX-FREE) 1 GM	1	EA	VL	IV	EA	500	MG	2	04/06/2005	99/99/9999							
00409-6534-01		J3370		06/08/2005	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (ADD-VANTAGE,LATEX-FREE) 500 MG	1	EA	VL	IV	EA	500	MG	1	06/08/2005	99/99/9999							
00143-9513-01		J2469		03/26/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL (PF) 0.25 MG/1 ML	2	ML	VL	IV	ML	25	MCG	5	03/26/2018	99/99/9999							
00409-6534-49		J3370		06/10/2005	05/01/2015	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL NOVATION (ADD-VANTAGE,10X10) 500 MG	1	EA	VL	IV	EA	500	MG	1	06/10/2005	05/01/2015							
00409-6535-01		J3370		03/29/2005	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HYDROCHLORIDE (ADD-VANTAGE,LATEX-FREE) 1 GM	1	EA	VL	IV	EA	500	MG	2	03/29/2005	99/99/9999							
00409-6535-49		J3370		04/06/2005	12/01/2015	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HYDROCHLORIDE NOVATION (ADD-VANTAGE,LATEX-FREE) 1 GM	1	EA	VL	IV	EA	500	MG	2	04/06/2005	12/01/2015							
00409-6629-02		J0330		04/25/2005	99/99/9999	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	QUELICIN (VIAL,FLIPTOP) 20 MG/ML	10	ML	VL	IV	ML	20	MG	1	04/25/2005	99/99/9999							
00409-6629-61		J0330		04/20/2006	06/05/2014	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	AMERINET CHOICE SUCCINYLCHOLINE CHLORIDE (USP,25X10ML,MD FLIPTOP) 20 MG/ML	10	ML	VL	IJ	ML	20	MG	1	04/20/2006	06/05/2014							
00409-6635-01		J3480		09/21/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (FTV,25X5ML,10ML VIAL) 2 MEQ/ML	5	ML	VL	IV	ML	2	MEQ	1	09/21/2005	99/99/9999							
00409-6636-01		J3480		08/09/2005	04/01/2013	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (FTV,30ML,LATEX-FREE) 2 MEQ/ML	15	ML	VL	IV	ML	2	MEQ	1	08/09/2005	04/01/2013							
00409-6648-02		J7799		03/29/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (VIAL,FLIPTOP,ADDITIVE) 50%	50	ML	VL	IV	ML	1	EA	1	03/29/2005	99/99/9999							
00409-6651-06		J3480		11/10/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (VIAL,FLIPTOP,20ML) 2 MEQ/ML	10	ML	VL	IV	ML	2	MEQ	1	11/10/2005	99/99/9999							
00409-6653-05		J3480		08/09/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (FTV,30ML,LATEX-FREE) 2 MEQ/ML	20	ML	VL	IV	ML	2	MEQ	1	08/09/2005	99/99/9999							
00409-6657-73		J7799		10/14/2005	01/01/2018	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (FTV,50MEQ,25X20ML) 14.6%	20	ML	VL	IV	ML	1	EA	1	10/14/2005	01/01/2018							
00409-6660-75		J7799		07/26/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (25X40ML,LATEX-FREE) 14.6%	40	ML	VL	IV	ML	1	EA	1	07/26/2005	99/99/9999							
00409-6727-23		J3475		09/20/2005	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	DEXTROSE/MAGNESIUM SULFATE (PLASTIC CONTAINER) 5%-1 GM/100 ML	100	ML	FC	IV	ML	500	MG	0.02	09/20/2005	99/99/9999							
00409-6729-03		J3475		08/16/2005	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (24X500ML,LATEX-FREE) 40 MG/ML	500	ML	PC	IV	ML	500	MG	0.08	08/16/2005	99/99/9999							
00409-6729-09		J3475		09/22/2005	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (PLASTIC CONTAINER) 40 MG/ML	1000	ML	PC	IV	ML	500	MG	0.08	09/22/2005	99/99/9999							
00409-6729-23		J3475		10/06/2005	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (24X100ML,LATEX-FREE) 40 MG/ML	100	ML	PC	IV	ML	500	MG	0.08	10/06/2005	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-6729-24		J3475		12/01/2006	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (SINGLE DOSE,LATEX-FREE) 40 MG/ML	50 ML	FC	IV	ML		500 MG		0.08	12/01/2006	99/99/9999						
00409-6730-13		J3475		04/03/2006	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (LATEX-FREE) 80 MG/ML	50 ML	FC	IV	ML		500 MG		0.16	04/03/2006	99/99/9999						
00409-6778-02		J2060		01/27/2006	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (10X1ML) 2 MG/ML	1 ML	VL	IJ	ML		2 MG		1	01/27/2006	99/99/9999						
00409-6778-62		J2060		06/28/2005	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (10X1ML) 2 MG/ML	1 ML	VL	IJ	ML		2 MG		1	06/28/2005	99/99/9999						
00409-6779-02		J2060		01/05/2006	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (VIAL, FLIPTOP) 4 MG/ML	10 ML	VL	IJ	ML		2 MG		2	01/05/2006	99/99/9999						
00409-6780-02		J2060		12/29/2005	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (VIAL,FLIPTOP) 2 MG/ML	10 ML	VL	IJ	ML		2 MG		1	12/29/2005	99/99/9999						
00409-6781-02		J2060		01/23/2006	12/08/2017	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P., 10X10ML) 4 MG/ML QUELICIN (FTV,25X10ML,20ML VIAL) 100 MG/ML	10 ML	VL	IJ	ML		2 MG		2	01/23/2006	12/08/2017						
00409-6970-10		J0330		09/30/2005	08/01/2013	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	POTASSIUM CHLORIDE (P.C.,LATEX-FREE) 10 MEQ/100 ML	10 ML	VL	IV	ML		20 MG		5	09/30/2005	08/01/2013						
00409-7074-26		J3480		04/25/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (24X50ML,LATEX-FREE) 10 MEQ/50 ML	50 ML	PC	IV	ML		2 MEQ		0.05	04/25/2005	99/99/9999						
00409-7075-14		J3480		06/08/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (PC,24X100ML,LATEX-FREE) 20 MEQ/100 ML	100 ML	FC	IV	ML		2 MEQ		0.1	06/08/2005	99/99/9999						
00409-7075-26		J3480		04/11/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (USP,100MLX24) 30 MEQ/100 ML	100 ML	FC	IV	ML		2 MEQ		0.15	02/08/2006	99/99/9999						
00409-7076-26		J3480		02/08/2006	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (24X50ML,LATEX-FREE) 20 MEQ/50 ML	50 ML	FC	IV	ML		2 MEQ		0.2	06/28/2005	99/99/9999						
00409-7077-14		J3480		06/28/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (HIGHLY CONC.,24X100ML) 40 MEQ/100 ML	100 ML	FC	IV	ML		2 MEQ		0.2	05/04/2005	99/99/9999						
00409-7077-26		J3480		05/04/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	DEXTRROSE (ADD-VANTAGE,24X250ML)	250 ML	FC	IV	ML		500 ML		0.002	07/22/2005	99/99/9999						
00409-7100-02		J7060		07/22/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTRROSE (ADD-VANTAGE,LATEX-FREE) 5%	50 ML	FC	IV	ML		500 ML		0.002	08/17/2005	99/99/9999						
00409-7100-66		J7060		08/17/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTRROSE (ADD-VANTAGE,50X100ML) 5%	100 ML	FC	IV	ML		500 ML		0.002	09/14/2005	99/99/9999						
00409-7100-67		J7060		09/14/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	SODIUM CHLORIDE (ADD-VANTAGE,24X250ML,PF) 0.9%	250 ML	FC	IV	ML		250 ML		0.004	07/08/2005	99/99/9999						
00409-7101-02		J7050		07/08/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (ADD-VANT LIFECARE) 0.9%	50 ML	FC	IV	ML		10 ML		0.1	07/28/2005	99/99/9999						
00409-7101-66		A4216		07/28/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (50X100ML, ADD-VANTAGE) 0.9%	100 ML	PC	IV	ML		250 ML		0.004	08/24/2005	99/99/9999						
00409-7101-67		J7050		08/24/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	DEXLACT. RINGERS/POTASSIUM CHL (12X1000ML,LATEX-FREE)	1000 ML	FC	IV	ML		1000 ML		0.0005	08/05/2005	99/99/9999						
00409-7111-09		J7120		08/05/2005	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	DEXTRROSE/LACTATED RINGERS/POTASSIUM CHLORIDE (5% DEXTROSE,LATEX-FREE)	1000 ML	FC	IV	ML		1000 ML		0.0005	02/21/2005	99/99/9999						
00409-7113-09		J7120		02/21/2005	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	POTASSIUM CHLORIDE/SODIUM CHLORIDE (12X1000ML,LATEX-FREE) 2 MEQ/100 ML-0.9%	1000 ML	FC	IV	ML		2 MEQ		0.01	04/06/2005	99/99/9999						
00409-7115-09		J3480		04/06/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE/SODIUM CHLORIDE (12X100ML,LATEX-FREE) 4 MEQ/100 ML-0.9%	1000 ML	FC	IV	ML		2 MEQ		0.02	06/22/2005	99/99/9999						
00409-7116-09		J3480		06/22/2005	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	WATER FOR IRRIGATION (BULK PACKAGE,PF)	2000 ML	FC	IR	ML		500 ML		0.002	08/16/2005	99/99/9999						
00409-7118-07		A4217		08/16/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	DEXTRROSE (2000MLX6) 50%	2000 ML	FC	IV	ML		1 EA		1	05/27/2006	06/10/2016						
00409-7119-07		J7799		05/27/2006	06/10/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (6X2000ML,LATEX-FREE) 70% SODIUM CHLORIDE (USP,ADD-VANTAGE) 0.45%	2000 ML	FC	IV	ML		1 EA		1	07/06/2005	99/99/9999						
00409-7120-07		J7799		07/06/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (ADD-VANTAGE,LATEX-FREE) 0.45%	50 ML	FC	IV	ML		1 EA		1	09/12/2005	99/99/9999						
00409-7132-02		J7799		05/26/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (ADD-VANTAGE,LATEX-FREE) 0.45%	100 ML	PC	IV	ML		1 EA		1	11/14/2005	99/99/9999						
00409-7132-66		J7799		09/12/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (AQUALITE,12X1000ML,PF) 0.9%	1000 ML	FC	IR	ML		500 ML		0.002	05/11/2005	99/99/9999						
00409-7132-67		J7799		11/14/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (AQUALITE,9X1500ML,PF) 0.9%	1500 ML	PC	IR	ML		500 ML		0.002	06/09/2005	99/99/9999						
00409-7138-09		A4217		05/11/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (AQUALITE W/HANGER,PF)	1000 ML	PC	IR	ML		500 ML		0.002	03/02/2005	99/99/9999						
00409-7138-36		A4217		06/09/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (AQUALITE)	1500 ML	PC	IR	ML		500 ML		0.002	05/04/2005	99/99/9999						
00409-7139-09		A4217		03/02/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	CEFTRIAXONE (USP,FLIPTOP VIAL) 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/20/2005	99/99/9999						
00409-7139-36		A4217		05/04/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	CEFTRIAXONE (USP,ADD-VANTAGE VIAL) 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/20/2005	99/99/9999						
00409-7332-01		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS (USP,ADD-VANTAGE VIAL) 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/20/2005	99/99/9999						
00409-7333-04		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP,BULK PACK) 10 GM	1 EA	VL	IJ	EA		250 MG		40	07/20/2005	99/99/9999						
00409-7333-49		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG																	
00409-7334-10		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG																	

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-7336-04		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP,ADD-VANTAGE VIAL) 2 GM	1 EA	VL	IJ	EA		250 MG		8	07/20/2005	99/99/9999						
00409-7336-49		J0696		07/20/2005	11/01/2016	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS (USP,ADD-VANTAGE VIAL) 2 GM	1 EA	VL	IJ	EA		250 MG		8	07/20/2005	11/01/2016						
00409-7337-01		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 250 MG	1 EA	VL	IJ	EA		250 MG		1	07/20/2005	99/99/9999						
00409-7338-01		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 500 MG	1 EA	VL	IJ	EA		250 MG		2	07/20/2005	99/99/9999						
00409-7385-01		J0280		12/29/2005	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE (AMP,LATEX-FREE) 25 MG/ML	10 ML	AM	IV	ML		250 MG		0.1	12/29/2005	99/99/9999						
00409-7386-01		J0280		11/29/2005	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE (AMP,LATEX-FREE) 25 MG/ML	20 ML	AM	IV	ML		250 MG		0.1	11/29/2005	99/99/9999						
00409-7418-03		J7100		02/14/2006	99/99/9999	INFUSION, DEXTRAN 40, 500 ML	LMD IN DEXTROSE (12X500ML,LATEX-FREE) 10%-5%	500 ML	FC	IV	ML		500 ML		0.002	02/14/2006	99/99/9999						
00409-7419-03		J7100		08/09/2005	99/99/9999	INFUSION, DEXTRAN 40, 500 ML	LMD W/0.9% SODIUM CHLORIDE (LATEX-FREE) 10%-0.9%	500 ML	FC	IV	ML		500 ML		0.002	08/09/2005	99/99/9999						
00409-7517-16		J7799		12/07/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (ANSYR II,LATEX-FREE) 50% MEPIVACAINE HYDROCHLORIDE	50 ML	SR	IV	ML		1 EA		1	12/07/2005	99/99/9999						
00409-7551-01		J0670		05/05/2008	06/01/2012	INJECTION, MEPIVACAINE HYDROCHLORIDE, PER 10 ML	(50X1.8ML,DENTALCARPULE) 3% HEPARIN SODIUM/SODIUM CHLORIDE (18X500ML,LATEX-FREE) 200 U/100 ML-0.9%	1.8 ML	CT	IJ	ML		10 ML		0.1	05/05/2008	06/01/2012						
00409-7620-03		J1644		04/05/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE (LATEX-FREE) 200 U/100 ML-0.9%	500 ML	FC	IV	ML		1000 U		0.002	04/05/2005	99/99/9999						
00409-7620-59		J1644		04/13/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE (LATEX-FREE) 200 U/100 ML-0.9%	1000 ML	FC	IV	ML		1000 U		0.002	04/13/2005	99/99/9999						
00409-7650-62		J1644		07/06/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE (24X250ML,LATEX-FREE) 10000 U/100 ML-0.45%	250 ML	FC	IV	ML		1000 U		0.1	07/06/2005	99/99/9999						
00409-7651-03		J1644		06/28/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE (24X500ML,LATEX-FREE) 5000 U/100 ML-0.45%	500 ML	FC	IV	ML		1000 U		0.05	06/28/2005	99/99/9999						
00409-7651-62		J1644		07/28/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM/SODIUM CHLORIDE (24X250ML,LATEX-FREE) 5000 U/100 ML-0.45%	250 ML	FC	IV	ML		1000 U		0.05	07/28/2005	99/99/9999						
00409-7666-62		J2810		01/27/2006	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE IN DEXTROSE (24X250ML,LATEX-FREE) 5%-160 MG/100 ML	250 ML	FC	IV	ML		40 MG		0.04	01/27/2006	99/99/9999						
00409-7668-23		J2810		02/06/2007	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE IN DEXTROSE (24X100ML,SINGLE-DOSE) 5%-200 MG/100 ML	100 ML	FC	IV	ML		40 MG		0.05	02/06/2007	99/99/9999						
00409-7677-13		J2810		08/10/2006	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	DEXTROSE/THEOPHYLLINE (50MLX24,DEHP,LATEX-FREE) 5%-200 MG/50 ML	50 ML	FC	IV	ML		40 MG		0.1	08/10/2006	99/99/9999						
43598-0637-10		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (10X100ML) 15 MG/1 ML	100 ML	BG	IV	ML		10 MG		1.5	06/13/2018	99/99/9999						
43598-0637-52		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (1X100ML, INNER PACK) 15 MG/1 ML	100 ML	BG	IV	ML		10 MG		1.5	06/13/2018	99/99/9999						
55150-0204-20		J3370		08/30/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF,LATEX-FREE) 1 GM	10 EA	VL	IV	EA		500 MG		2	08/30/2018	99/99/9999						
00409-7705-62		J2810		05/27/2006	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE IN DEXTROSE (USP,250MLX24) 5%-320 MG/100 ML	250 ML	FC	IV	ML		40 MG		0.08	05/27/2006	99/99/9999						
00409-7712-09		J7799		08/19/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	MANNITOL (LATEX-FREE) 5%	1000 ML	FC	IV	ML		1 EA		1	08/19/2005	99/99/9999						
00409-7713-09		J7799		04/07/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	MANNITOL (USP,LATEX-FREE) 10%	1000 ML	FC	IV	ML		1 EA		1	04/07/2006	99/99/9999						
00409-7714-03		J7799		08/30/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	MANNITOL (LATEX-FREE) 15%	500 ML	FC	IV	ML		1 EA		1	08/30/2005	99/99/9999						
00409-7715-02		J7799		11/14/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	MANNITOL (FLEX CONTAINER,24X250ML) 20%	250 ML	FC	IV	ML		1 EA		1	11/14/2005	99/99/9999						
00409-7715-03		J7799		09/16/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	MANNITOL (FLEX CONTAINER,12X500ML) 20%	500 ML	FC	IV	ML		1 EA		1	09/16/2005	99/99/9999						
00409-7730-20		J7799		07/27/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (QUAD-PK,48X25ML) 0.45%	25 ML	FC	IV	ML		1 EA		1	07/27/2005	99/99/9999						
00409-7730-36		J7799		07/11/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (80X50ML,LATEX-FREE) 0.45%	50 ML	FC	IV	ML		1 EA		1	07/11/2005	99/99/9999						
00409-7730-37		J7799		09/16/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (80X100ML,LATEX-FREE) 0.45%	100 ML	FC	IV	ML		1 EA		1	09/16/2005	99/99/9999						
00409-7760-03		J1644		08/30/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	DEXTROSE/HEPARIN SODIUM (LATEX-FREE) 5%-4000 U/100 ML	500 ML	FC	IV	ML		1000 U		0.04	08/30/2005	99/99/9999						
00409-7761-03		J1644		07/22/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	DEXTROSE/HEPARIN SODIUM (24X500ML,LATEX-FREE) 5%-5000 U/100 ML	500 ML	FC	IV	ML		1000 U		0.05	07/22/2005	99/99/9999						
00409-7793-62		J1644		10/14/2005	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	DEXTROSE/HEPARIN SODIUM (24X250ML,LATEX-FREE) 5%-10000 U/100 ML	250 ML	FC	IV	ML		1000 U		0.1	10/14/2005	99/99/9999						
00409-7794-62		J1644		06/12/2006	09/01/2017	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM IN DEXTROSE (24X250ML,USP,LATEX-FREE) 5%-5000 U/100 ML	250 ML	FC	IV	ML		1000 U		0.05	06/12/2006	09/01/2017						
00409-7808-22		J1265		01/01/2006	09/01/2017	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL (LIFECARE,12X250ML) 5%-80 MG/100 ML	250 ML	FC	IV	ML		40 MG		0.02	01/01/2006	09/01/2017						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55150-0210-10		J0583		09/27/2018	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (SINGLE-USE VIAL) 250 MG	10	EA	VL	IV	EA	1	MG	250	09/27/2018	99/99/9999						
00409-7808-24		J1265		01/01/2006	09/01/2017	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL (LIFECARE,LATEX-FREE) 5%-80 MG/100 ML	500	ML	FC	IV	ML	40	MG	0.02	01/01/2006	09/01/2017						
00409-7809-22		J1265		01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL (LIFECARE,LATEX-FREE) 5%-160 MG/100 ML	250	ML	PC	IV	ML	40	MG	0.04	01/01/2006	99/99/9999						
00409-7809-24		J1265		01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL (LIFECARE, 12X500ML) 5%-100 MG/100 ML	500	ML	FC	IV	ML	40	MG	0.025	01/01/2006	99/99/9999						
00409-7810-22		J1265		01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DEXTROSE/DOPAMINE HCL (LIFECARE, 12X250ML) 5%-320 MG/100 ML	250	ML	FC	IV	ML	40	MG	0.08	01/01/2006	99/99/9999						
00409-7811-24		J3490		08/31/2005	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (S.D.V.,LATEX-FREE) 500 MG/100 ML	100	ML	FC	IV	ML	1	EA	1	08/31/2005	99/99/9999						
00409-7811-37		J3490		09/22/2005	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (LIFECARE,QUAD PACK) 500 MG/100 ML	100	ML	FC	IV	ML	1	EA	1	09/22/2005	99/99/9999						
00409-7879-13		J1580		03/31/2006	08/01/2015	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE IN SODIUM CHLORIDE (LATEX-FREE) 1.2 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.015	03/31/2006	08/01/2015						
00409-7881-13		J1580		01/23/2006	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (LIFECARE, 24X50ML) 1.4 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.0175	01/23/2006	99/99/9999						
00409-7883-13		J1580		01/09/2006	06/01/2015	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (LIFECARE,LATEX-FREE) 1.6 MG/ML-0.9%	50	ML	FC	IV	ML	80	MG	0.02	01/09/2006	06/01/2015						
00409-7884-23		J1580		07/06/2005	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (LIFECARE,24X100ML) 80 MG/100 ML-0.9%	100	ML	FC	IV	ML	80	MG	0.01	07/06/2005	99/99/9999						
00409-7886-23		J1580		01/27/2006	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE IN SODIUM CHLORIDE (LIFECARE,24X100ML) 90 MG/100 ML-0.9%	100	ML	FC	IV	ML	80	MG	0.01125	01/27/2006	99/99/9999						
00409-7889-23		J1580		09/20/2005	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE/SODIUM CHLORIDE (LIFECARE,24X100ML) 100 MG/100 ML-0.9%	100	ML	FC	IV	ML	80	MG	0.0125	09/20/2005	99/99/9999						
00409-7918-19		J7799		07/08/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (12X500ML,LATEX-FREE) 70%	500	ML	PC	IV	ML	1	EA	1	07/08/2005	99/99/9999						
00409-7922-02		J7060		04/05/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE/PLASTIC) 5%	250	ML	FC	IV	ML	500	ML	0.002	04/05/2005	99/99/9999						
00409-7922-03		J7060		02/25/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE/PLASTIC) 5%	500	ML	FC	IV	ML	500	ML	0.002	02/25/2005	99/99/9999						
00409-7922-09		J7060		02/21/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE/PLASTIC) 5%	1000	ML	FC	IV	ML	500	ML	0.002	02/21/2005	99/99/9999						
00409-7922-30		J7060		04/14/2006	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (VISIV CONTAINER) 5%	500	ML	FC	IV	ML	500	ML	0.002	04/14/2006	99/99/9999						
00409-7922-48		J7060		04/14/2006	11/01/2013	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (VISIV CONTAINER) 5%	1000	ML	FC	IV	ML	500	ML	0.002	04/14/2006	11/01/2013						
00409-7922-53		J7060		09/01/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE, 24X250ML) 5%	250	ML	FC	IV	ML	500	ML	0.002	09/01/2005	99/99/9999						
00409-7922-55		J7060		10/31/2006	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (18X500ML,LATEX-FREE) 5%	500	ML	FC	IV	ML	500	ML	0.002	10/31/2006	99/99/9999						
00409-7922-61		J7060		08/05/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE, 32X150ML) 5%	150	ML	FC	IV	ML	500	ML	0.002	08/05/2005	99/99/9999						
00409-7923-13		J7060		06/09/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (48X50ML,LATEX-FREE) 5%	50	ML	FC	IV	ML	500	ML	0.002	06/09/2005	99/99/9999						
00409-7923-20		J7060		06/17/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE, 48X25ML) 5%	25	ML	FC	IV	ML	500	ML	0.002	06/17/2005	99/99/9999						
00409-7923-23		J7060		07/15/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (48X100ML,LATEX-FREE) 5%	100	ML	FC	IV	ML	500	ML	0.002	07/15/2005	99/99/9999						
00409-7923-36		J7060		04/05/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE,QUAD PACK) 5%	50	ML	FC	IV	ML	500	ML	0.002	04/05/2005	99/99/9999						
00409-7923-37		J7060		03/16/2005	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (LIFECARE,80X100ML) 5%	100	ML	FC	IV	ML	500	ML	0.002	03/16/2005	99/99/9999						
00409-7924-02		J7799		07/28/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (24X250ML,LATEX-FREE) 5%-0.225%	250	ML	FC	IV	ML	1	EA	1	07/28/2005	99/99/9999						
00409-7924-03		J7799		07/28/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (LIFECARE/PLASTIC) 5%-0.225%	500	ML	FC	IV	ML	1	EA	1	07/28/2005	99/99/9999						
00409-7924-09		J7799		12/21/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (LIFECARE, PLASTIC) 5%-0.225%	1000	ML	FC	IV	ML	1	EA	1	12/21/2005	99/99/9999						
00409-7925-03		J7799		09/16/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (LIFECARE,PLASTIC) 5%-0.3%	500	ML	FC	IV	ML	1	EA	1	09/16/2005	99/99/9999						
00409-7925-09		J7799		03/17/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (12X1000ML) 5%-0.3%	1000	ML	FC	IV	ML	1	EA	1	03/17/2006	99/99/9999						
00409-7926-02		J7799		08/30/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (LIFECARE/PLASTIC) 5%-0.45%	250	ML	FC	IV	ML	1	EA	1	08/30/2005	99/99/9999						
00409-7926-03		J7799		06/07/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (24X500ML,LATEX-FREE) 5%-0.45%	500	ML	FC	IV	ML	1	EA	1	06/07/2005	99/99/9999						
00409-7926-09		J7799		08/25/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (12X1000ML, LIFECARE) 5%-0.45%	1000	ML	FC	IV	ML	1	EA	1	08/25/2005	99/99/9999						
00409-7926-30		J7799		04/14/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (VISIV CONTAINER) 5%-0.45%	500	ML	FC	IV	ML	1	EA	1	04/14/2006	99/99/9999						
00409-7926-48		J7799		04/14/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (VISIV CONTAINER) 5%-0.45%	1000	ML	FC	IV	ML	1	EA	1	04/14/2006	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-7929-03	J7120			06/09/2005	12/31/2015	RINGERS LACTATE INFUSION, UP TO 1000 CC	DEXTRROSE 5% IN RINGERS (LATEX-FREE)	500	ML	FC	IV	ML	1000	ML	0.0005	06/09/2005	12/31/2015						
00409-7929-09	J7120			02/07/2005	12/31/2015	RINGERS LACTATE INFUSION, UP TO 1000 CC	DEXTRROSE 5% IN RINGERS (LIFECARE,LATEX-FREE)	1000	ML	FC	IV	ML	1000	ML	0.0005	02/07/2005	12/31/2015						
00409-7930-02	J7799			07/05/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (24X250ML,LIFECARE) 10%	250	ML	FC	IV	ML	1	EA	1	07/05/2005	99/99/9999						
00409-7930-03	J7799			01/12/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (LIFECARE,LATEX-FREE) 10%	500	ML	FC	IV	ML	1	EA	1	01/12/2005	99/99/9999						
00409-7930-09	J7799			03/16/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (LIFECARE,LATEX-FREE) 10%	1000	ML	FC	IV	ML	1	EA	1	03/16/2005	99/99/9999						
00409-7931-24	J2001			05/18/2005	06/01/2013	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTRROSE/LIDOCAINE HCL (LIFECARE,24X500ML) 5%-0.4%	500	ML	PC	IV	ML	10	MG	0.4	05/18/2005	06/01/2013						
00409-7931-32	J2001			09/16/2005	11/01/2012	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTRROSE/LIDOCAINE HCL (LIFECARE,12X250ML) 5%-0.4%	250	ML	FC	IV	ML	10	MG	0.4	09/16/2005	11/01/2012						
00409-7935-19	J7799			09/12/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (1000ML CONTAINER) 20%	500	ML	FC	IV	ML	1	EA	1	09/12/2005	99/99/9999						
00409-7936-19	J7799			06/24/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (12X500ML,LATEX-FREE) 50%	500	ML	PC	IV	ML	1	EA	1	06/24/2005	99/99/9999						
00409-7936-29	J7799			10/28/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (2000ML BAG,6X1000ML) 50%	1000	ML	FC	IV	ML	1	EA	1	10/28/2005	99/99/9999						
00409-7937-19	J7799			08/24/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (12X500ML,LATEX-FREE) 40%	500	ML	FC	IV	ML	1	EA	1	08/24/2005	99/99/9999						
00409-7938-19	J7799			09/29/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTRROSE (1000ML CONTAINER) 10%	500	ML	PC	IV	ML	1	EA	1	09/29/2005	99/99/9999						
00409-7939-32	J2001			01/11/2006	05/01/2012	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	DEXTRROSE/LIDOCAINE HCL (LIFECARE,LATEX-FREE) 5%-0.8%	250	ML	FC	IV	ML	10	MG	0.8	01/11/2006	05/01/2012						
00409-7941-02	J7042			05/27/2006	99/99/9999	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTRROSE AND SODIUM CHLORIDE (250MLX2,USP,LATEX-FREE) 5%-0.9%	250	ML	FC	IV	ML	5	%	0.002	05/27/2006	99/99/9999						
00409-7941-03	J7042			09/20/2005	99/99/9999	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTRROSE/SODIUM CHLORIDE (24X500ML,LATEX-FREE) 5%-0.9%	500	ML	FC	IV	ML	5	%	0.002	09/20/2005	99/99/9999						
00409-7941-09	J7042			08/08/2005	99/99/9999	5% DEXTROSE/NORMAL SALINE (500 ML = 1 UNIT)	DEXTRROSE/SODIUM CHLORIDE (LIFECARE,12X1000ML) 5%-0.9%	1000	ML	FC	IV	ML	5	%	0.002	08/08/2005	99/99/9999						
00409-7953-02	J7120			03/09/2005	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (LIFECARE,LATEX-FREE)	250	ML	FC	IV	ML	1000	ML	0.001	03/09/2005	99/99/9999						
00409-7953-03	J7120			05/20/2005	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (LIFECARE,24X500ML)	500	ML	PC	IV	ML	1000	ML	0.001	05/20/2005	99/99/9999						
00409-7953-09	J7120			05/18/2005	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (LIFECARE,LATEX-FREE)	1000	ML	PC	IV	ML	1000	ML	0.001	05/18/2005	99/99/9999						
00409-7953-30	J7120			04/14/2006	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (VISIV CONTAINER)	500	ML	FC	IV	ML	1000	ML	0.001	04/14/2006	99/99/9999						
00409-7953-48	J7120			04/14/2006	99/99/9999	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGER'S (VISIV CONTAINER)	1000	ML	FC	IV	ML	1000	ML	0.001	04/14/2006	99/99/9999						
00409-7972-05	A4217			09/01/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (FLEXIBLE CONTAINER,PF) 0.9%	1000	ML	FC	IR	ML	500	ML	0.002	09/01/2005	99/99/9999						
00409-7972-07	A4217			04/05/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (FLEX CONTAINER,6X2000ML) 0.9%	2000	ML	FC	IR	ML	500	ML	0.002	04/05/2005	99/99/9999						
00409-7972-08	A4217			05/18/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (FLEX CONTAINER-4X3000ML) 0.9%	3000	ML	PC	IR	ML	500	ML	0.002	05/18/2005	99/99/9999						
00409-7973-05	A4217			03/16/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (FLEXIBLE CONTAINER,PF)	1000	ML	FC	IR	ML	500	ML	0.002	03/16/2005	99/99/9999						
00409-7973-07	A4217			08/09/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (FLEXIBLE, CONTAINER,PF)	2000	ML	FC	IR	ML	500	ML	0.002	08/09/2005	99/99/9999						
00409-7973-08	A4217			07/14/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION (4X3000ML,PF,LATEX-FREE)	3000	ML	FC	IR	ML	500	ML	0.002	07/14/2005	99/99/9999						
00409-7975-07	A4217			04/26/2006	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (USP,6X2000ML) 0.45%	2000	ML	FC	IR	ML	500	ML	0.002	04/26/2006	99/99/9999						
00409-7983-02	J7050			07/01/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (LIFECARE,24X250ML,PF) 0.9%	250	ML	FC	IV	ML	250	ML	0.004	07/01/2005	99/99/9999						
00409-7983-03	J7040			01/05/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION, STERILE (500 ML=1 UNIT)	SODIUM CHLORIDE (LIFECARE,P.C.,24X500ML) 0.9%	500	ML	FC	IV	ML	500	ML	0.002	01/05/2005	99/99/9999						
00409-7983-09	J7030			02/07/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 1000 CC	SODIUM CHLORIDE (LIFECARE,P.C.,12X1000ML) 0.9%	1000	ML	FC	IV	ML	1000	ML	0.001	02/07/2005	99/99/9999						
00409-7983-30	J7040			04/14/2006	10/16/2014	INFUSION, NORMAL SALINE SOLUTION, STERILE (500 ML=1 UNIT)	SODIUM CHLORIDE (VISIV CONTAINER) 0.9%	500	ML	FC	IV	ML	500	ML	0.002	04/14/2006	10/16/2014						
00409-7983-48	J7030			04/14/2006	10/16/2014	INFUSION, NORMAL SALINE SOLUTION , 1000 CC	SODIUM CHLORIDE (VISIV CONTAINER) 0.9%	1000	ML	FC	IV	ML	1000	ML	0.001	04/14/2006	10/16/2014						
00409-7983-53	J7050			09/30/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (LIFECARE,2 PORTS,PC,LF) 0.9%	250	ML	FC	IV	ML	250	ML	0.004	09/30/2005	99/99/9999						
00409-7983-55	J7040			04/11/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION, STERILE (500 ML=1 UNIT)	SODIUM CHLORIDE (LIFECARE,2 PORTS,PC,LF) 0.9%	500	ML	FC	IV	ML	500	ML	0.002	04/11/2005	99/99/9999						
00409-7983-61	J7050			06/17/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (LIFECARE,P.C.,32X150ML) 0.9%	150	ML	FC	IV	ML	250	ML	0.004	06/17/2005	99/99/9999						
00409-7984-13	A4216			06/20/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (48X50ML,PF,LATEX-FREE) 0.9%	50	ML	FC	IV	ML	10	ML	0.1	06/20/2005	99/99/9999						
00409-7984-20	A4216			06/17/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (LIFECARE,QUAD PACK,LF) 0.9%	25	ML	FC	IV	ML	10	ML	0.1	06/17/2005	99/99/9999						
00409-7984-23	J7050			05/18/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (LIFECARE SINGLE-PF) 0.9%	100	ML	PC	IV	ML	250	ML	0.004	05/18/2005	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-7984-36	A4216			07/14/2005	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (LFCARE,QUAD,LF,80X50ML) 0.9%	50 ML	FC	IV	ML		10 ML		0.1	07/14/2005	99/99/9999						
00409-7984-37	J7050			07/15/2005	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (LFCARE,QUAD,LF,80X100ML) 0.9%	100 ML	FC	IV	ML		250 ML		0.004	07/15/2005	99/99/9999						
00409-7985-02	J7799			04/06/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (24X250ML,LATEX-FREE) 0.45%	250 ML	FC	IV	ML		1 EA		1	04/06/2005	99/99/9999						
00409-7985-03	J7799			04/06/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (LIFECARE,24X500ML) 0.45%	500 ML	FC	IV	ML		1 EA		1	04/06/2005	99/99/9999						
00409-7985-09	J7799			11/24/2004	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (LIFECARE,12X1000ML) 0.45%	1000 ML	FC	IV	ML		1 EA		1	11/24/2004	99/99/9999						
00409-7990-09	A4217			09/02/2005	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION (LIFECARE,PF,LATEX-FREE)	1000 ML	FC	IV	ML		500 ML		0.002	09/02/2005	99/99/9999						
00409-8004-15	J7799			08/01/2005	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (12X500ML,LATEX-FREE) 30%	500 ML	FC	IV	ML		1 EA		1	08/01/2005	99/99/9999						
00409-9093-32	J3010			11/14/2005	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (10X2ML,LATEX-FREE) 0.05 MG/ML	2 ML	AM	IJ	ML		0.1 MG		0.5	11/14/2005	99/99/9999						
00409-9093-35	J3010			12/13/2005	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (AMP,LATEX-FREE) 0.05 MG/ML	5 ML	AM	IJ	ML		0.1 MG		0.5	12/13/2005	99/99/9999						
00409-9093-38	J3010			03/03/2006	09/01/2017	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (5X20ML) 0.05 MG/ML	20 ML	AM	IJ	ML		0.1 MG		0.5	03/03/2006	09/01/2017						
00409-9094-22	J3010			10/12/2005	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (FTV,25X2ML,LATEX-FREE) 0.05 MG/ML	2 ML	VL	IJ	ML		0.1 MG		0.5	10/12/2005	99/99/9999						
00409-9094-25	J3010			11/07/2005	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (VIAL,FLIPTOP,LATEX-FREE) 0.05 MG/ML	5 ML	VL	IJ	ML		0.1 MG		0.5	11/07/2005	99/99/9999						
00409-9094-28	J3010			02/14/2006	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (25X10ML,FTV) 0.05 MG/ML	10 ML	VL	IJ	ML		0.1 MG		0.5	02/14/2006	99/99/9999						
00409-9094-31	J3010			09/23/2005	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (FTV,LATEX-FREE) 0.05 MG/ML	20 ML	VL	IJ	ML		0.1 MG		0.5	09/23/2005	99/99/9999						
25021-0828-50	J0640			09/04/2018	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (SDV,PF,LATEX-FREE) 500 MG	1 EA	VL	IJ	EA		50 MG		10	09/04/2018	99/99/9999						
00409-9094-61	J3010			12/30/2005	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (VIAL, FLIPTOP) 0.05 MG/ML	50 ML	VL	IJ	ML		0.1 MG		0.5	12/30/2005	99/99/9999						
00409-9104-20	J1265			01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HCL (25X10ML) 40 MG/ML	10 ML	VL	IV	ML		40 MG		1	01/01/2006	99/99/9999						
00409-9137-05	J2001			06/30/2005	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (ANSYR,10X5ML,LATEX-FREE) 1%	5 ML	SR	EP	ML		10 MG		1	06/30/2005	99/99/9999						
00409-9631-04	J1940			04/21/2006	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (PF) 10 MG/ML	4 ML	SR	IJ	ML		20 MG		0.5	04/21/2006	99/99/9999						
00463-1015-30	J3420			01/01/2002	02/03/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	VITAMIN B12 (VIAL) 1000 MCG/ML	30 ML	VL	IM	ML		1000 MCG		1	01/01/2002	02/03/2016						
00463-1019-30	J2650			01/01/2002	02/03/2016	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	COTOLONE (VIAL) 25 MG/ML	30 ML	VL	IJ	ML		1 ML		1	01/01/2002	02/03/2016						
00463-1020-10	J2650			01/01/2002	02/03/2016	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	COTOLONE (VIAL) 50 MG/ML	10 ML	VL	IJ	ML		1 ML		1	01/01/2002	02/03/2016						
00463-1021-30	J3420			01/01/2002	02/03/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	VITAMIN B12 (VIAL) 100 MCG/ML	30 ML	VL	IM	ML		1000 MCG		0.1	01/01/2002	02/03/2016						
00463-1029-30	J1435			01/01/2002	01/28/2016	INJECTION, ESTRONE, PER 1 MG	ESTRONE (VIAL, AQUEOUS) 5 MG/ML	30 ML	EA	IM	ML		1 MG		5	01/01/2002	01/28/2016						
00463-1036-10	J1700			01/01/2002	02/03/2016	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (VIAL) 25 MG/ML	10 ML	VL	IJ	ML		25 MG		1	01/01/2002	02/03/2016						
00463-1069-10	J3140			01/01/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTO AQ (VIAL) 100 MG/ML	10 ML	VL	IM	ML		50 MG		2	01/01/2002	12/31/2014						
00463-1073-10	J3150			01/01/2002	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (VIAL) 100 MG/ML	10 ML	VL	IM	ML		100 MG		1	01/01/2002	12/31/2014						
00463-1074-30	J3411			01/01/2004	02/03/2016	INJECTION, THIAMINE HCL, 100 MG	THIAMINE HCL (VIAL) 100 MG/ML	30 ML	VL	IJ	ML		100 MG		1	01/01/2004	02/03/2016						
00463-1080-30	J1200			01/01/2002	02/03/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	TRUXADRYL (VIAL) 10 MG/ML	30 ML	VL	IJ	ML		50 MG		0.2	01/01/2002	02/03/2016						
00463-1086-10	J1240			01/01/2002	09/30/2013	INJECTION, DIMENHYDRINATE, UP TO 50 MG	DIMENHYDRINATE (VIAL) 50 MG/ML	10 ML	VL	IJ	ML		50 MG		1	01/01/2002	09/30/2013						
00463-1091-05	J3302			01/01/2002	02/03/2016	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCOT (VIAL) 40 MG/ML	5 ML	VL	IJ	ML		5 MG		8	01/01/2002	02/03/2016						
00463-1092-10	J2360			01/01/2002	01/28/2016	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORFRO (VIAL) 30 MG/ML	10 ML	VL	IJ	ML		60 MG		0.5	01/01/2002	01/28/2016						
00463-1094-30	J3420			01/01/2002	01/01/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	HYDROXOCOBALAMIN (VIAL) 1000 MCG/ML	30 ML	VL	IM	ML		1000 MCG		1	01/01/2002	01/01/2016						
00463-1101-10	J3410			01/01/2002	02/03/2016	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	VISTACOT (VIAL) 50 MG/ML	10 ML	VL	IM	ML		25 MG		2	01/01/2002	02/03/2016						
00463-1104-10	J0500			01/01/2002	01/01/2016	INJECTION, DICYCLIMINE HCL, UP TO 20 MG	DICYCLOCOT (VIAL) 10 MG/ML	10 ML	VL	IM	ML		20 MG		0.5	01/01/2002	01/01/2016						
00463-1108-20	J3250			01/01/2002	01/01/2016	INJECTION, TRIMETHOBENZAMIDE HCL, UP TO 200 MG	BENZACOT (VIAL) 100 MG/ML	20 ML	VL	IM	ML		200 MG		0.5	01/01/2002	01/01/2016						
00463-6071-10	J7510			01/01/2002	02/03/2016	PREDNISOLONE ORAL, PER 5 MG	COTOLONE 5 MG	1000 EA	NA	PO	EA		5 MG		1	01/01/2002	02/03/2016						
00463-6140-10	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNICOT 10 MG	1000 EA	NA	PO	EA		5 MG		2	01/01/2002	12/31/2015						
00463-6141-10	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNICOT 20 MG	1000 EA	NA	PO	EA		5 MG		4	01/01/2002	12/31/2015						
00463-6155-10	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNICOT 5 MG	1000 EA	NA	PO	EA		5 MG		1	01/01/2002	12/31/2015						
00463-6156-10	Q0170			01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMACOT 25 MG	1000 EA	NA	PO	EA		25 MG		1	01/01/2002	12/31/2013						
00469-0607-73	J7507			01/01/2002	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	PROGRAF 0.5 MG	100 EA	BO	PO	EA		1 MG		0.5	01/01/2002	99/99/9999						
00469-0617-11	J7507			01/01/2002	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	PROGRAF (10X10,BLISTER PACK) 1 MG	100 EA	BX	PO	EA		1 MG		1	01/01/2002	99/99/9999						
00469-0617-73	J7507			02/13/2002	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	PROGRAF 1 MG	100 EA	BO	PO	EA		1 MG		1	02/13/2002	99/99/9999						
00469-0657-11	J7507			01/01/2004	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	PROGRAF (10X10,BLISTER PACK) 5 MG	100 EA	BX	PO	EA		1 MG		5	01/01/2004	99/99/9999						
00469-0657-73	J7507			01/01/2004	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	PROGRAF 5 MG	100 EA	BO	PO	EA		1 MG		5	01/01/2004	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00469-0871-20		J0152		01/01/2004	12/31/2013	INJECTION, ADENOSINE FOR DIAGNOSTIC USE, 30 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOSCAN (S.D.V.,PF) 3 MG/ML	20	ML	VL	IV	ML	30	MG	0.1	01/01/2004	12/31/2013							
00469-0871-30		J0152		01/01/2004	12/31/2013	INJECTION, ADENOSINE FOR DIAGNOSTIC USE, 30 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOSCAN (S.D.V.,PF) 3 MG/ML	30	ML	VL	IV	ML	30	MG	0.1	01/01/2004	12/31/2013							
00469-3016-01		J7525		01/01/2002	99/99/9999	TACROLIMUS, PARENTERAL, 5 MG	PROGRAF (AMP,PF) 5 MG/ML	1	ML	AM	IV	ML	5	MG	1	01/01/2002	99/99/9999							
00469-3051-30		J0289		01/01/2003	99/99/9999	INJECTION, AMPHOTERICIN B LIPOSOME, 10 MG	AMBISOME 50 MG	1	EA	VL	IV	EA	10	MG	5	01/01/2003	99/99/9999							
00469-3211-10		J2248		01/01/2007	99/99/9999	INJECTION, MICAFUNGIN SODIUM, 1 MG	MYCAMINE (W/RED FLIP-OFF CAP) 100 MG	1	EA	VL	IV	EA	1	MG	100	01/01/2007	99/99/9999							
00469-3250-10		J2248		01/01/2007	99/99/9999	INJECTION, MICAFUNGIN SODIUM, 1 MG	MYCAMINE (PF) 50 MG	1	EA	VL	IV	EA	1	MG	50	01/01/2007	99/99/9999							
00469-8234-12		J0150		06/14/2002	12/31/2014	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOCARD (ANSYR,LUER LOK) 3 MG/ML	2	ML	SR	IV	ML	6	MG	0.5	06/14/2002	12/31/2014							
00469-8234-14		J0150		06/14/2002	12/31/2014	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOCARD (ANSYR,LUER LOK) 3 MG/ML	4	ML	SR	IV	ML	6	MG	0.5	06/14/2002	12/31/2014							
00472-0082-16		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG/5 ML	480	ML	BO	PO	ML	1	EA	1	01/01/2002	99/99/9999							
00487-0201-01		J7620		01/01/2008	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	30	ML	PC	IH	ML	3	MG	0.33333	01/01/2008	99/99/9999							
00487-0201-02		J7620		01/01/2008	07/21/2016	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML, ROBOT READY) 3 MG/3 ML-0.5 MG/3 ML	30	ML	PC	IH	ML	3	MG	0.33333	01/01/2008	07/21/2016							
00487-0201-60		J7620		01/01/2008	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	60	ML	PC	IH	ML	3	MG	0.33333	01/01/2008	99/99/9999							
00487-9301-02		A4216		01/01/2006	07/21/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (ROBOT READY,30X3ML) 0.9%	3	ML	PC	IH	ML	10	ML	0.1	01/01/2006	07/21/2016							
00487-9301-03		A4216		01/01/2006	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (VIAL) 0.9%	3	ML	PC	IH	ML	10	ML	0.1	01/01/2006	99/99/9999							
00487-9301-33		A4216		01/01/2006	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE 0.9%	3	ML	PC	IH	ML	10	ML	0.1	01/01/2006	99/99/9999							
00487-9501-01		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9501-01	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9501-02		J7613		04/01/2008	07/21/2016	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (ROBOT READY,PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	07/21/2016							
00487-9501-02	KO	J7613	KO	04/01/2008	07/21/2016	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (ROBOT READY,PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	07/21/2016							
00487-9501-03		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9501-03	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9501-25		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9501-25	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9501-60		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9501-60	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	99/99/9999							
00487-9801-01		J7644		01/03/2003	99/99/9999	MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/03/2003	99/99/9999							
00487-9801-01	KO	J7644	KO	01/03/2003	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/03/2003	99/99/9999							
00487-9801-02		J7644		07/20/2005	07/21/2016	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (ROBOT READY,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	07/20/2005	07/21/2016							
00487-9801-02	KO	J7644	KO	07/20/2005	07/21/2016	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (ROBOT READY,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	07/20/2005	07/21/2016							
00487-9801-25		J7644		10/11/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	10/11/2002	99/99/9999							
00487-9801-25	KO	J7644	KO	10/11/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	10/11/2002	99/99/9999							
00487-9801-30		J7644		01/03/2003	99/99/9999	MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/03/2003	99/99/9999							
00487-9801-30	KO	J7644	KO	01/03/2003	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/03/2003	99/99/9999							
00487-9801-60		J7644		01/03/2003	99/99/9999	MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/03/2003	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00487-9801-60	KO	J7644	KO	01/03/2003	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/03/2003	99/99/9999						
00487-9901-02		J7611		04/01/2008	07/21/2016	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE (UNIT OF USE,ROBOT READY) 0.5%	0.5	ML	PC	IH	ML	1	MG	5	04/01/2008	07/21/2016						
00487-9901-30		J7611		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE (UNIT OF USE,PF) 0.5%	0.5	ML	PC	IH	ML	1	MG	5	04/01/2008	99/99/9999						
00487-9904-01		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.042%	3	ML	PC	IH	ML	1	MG	0.42	04/01/2008	99/99/9999						
00487-9904-01	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (PF) 0.042%	3	ML	PC	IH	ML	1	MG	0.42	04/01/2008	99/99/9999						
00487-9904-02		J7613		04/01/2008	07/21/2016	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (ROBOT READY,LDPE VIAL) 0.042%	3	ML	PC	IH	ML	1	MG	0.42	04/01/2008	07/21/2016						
00487-9904-02	KO	J7613	KO	04/01/2008	07/21/2016	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (ROBOT READY,LDPE VIAL) 0.042%	3	ML	PC	IH	ML	1	MG	0.42	04/01/2008	07/21/2016						
00487-9904-25		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (LDPE VIAL) 0.042%	3	ML	VL	IH	ML	1	MG	0.42	04/01/2008	99/99/9999						
00487-9904-25	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (LDPE VIAL) 0.042%	3	ML	VL	IH	ML	1	MG	0.42	04/01/2008	99/99/9999						
00490-0091-00		Q0175		01/01/2007	01/31/2014	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	100	EA	BO	PO	EA	4	MG	1	01/01/2007	01/31/2014						
00490-0091-30		Q0175		01/01/2007	01/31/2014	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	30	EA	BO	PO	EA	4	MG	1	01/01/2007	01/31/2014						
00490-0091-60		Q0175		01/01/2007	01/31/2014	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	60	EA	BO	PO	EA	4	MG	1	01/01/2007	01/31/2014						
00490-0091-90		Q0175		01/01/2007	01/31/2014	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	90	EA	BO	PO	EA	4	MG	1	01/01/2007	01/31/2014						
00517-0031-25		J3420		01/01/2002	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN 1000 MCG/ML	1	ML	VL	IM	ML	1000	MCG	1	01/01/2002	99/99/9999						
00517-0032-25		J3420		01/01/2002	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (M.D.V.) 1000 MCG/ML	10	ML	VL	IM	ML	1000	MCG	1	01/01/2002	99/99/9999						
00517-0033-25		J2710		01/15/2003	11/07/2013	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (M.D.V.) 1 MG/ML	10	ML	VL	IJ	ML	0.5	MG	2	01/15/2003	11/07/2013						
00517-0034-25		J2710		01/15/2003	11/07/2013	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (M.D.V.) 0.5 MG/ML	10	ML	VL	IJ	ML	0.5	MG	1	01/15/2003	11/07/2013						
00517-0130-05		J3420		05/29/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (M.D.V.) 1000 MCG/ML	30	ML	VL	IM	ML	1000	MCG	1	05/29/2003	99/99/9999						
00517-0132-25		J0636		03/14/2005	02/28/2013	INJECTION, CALCITRIOL, 0.1 MCG	CALCITRIOL 1 MCG/ML	1	ML	AM	IV	ML	0.1	MCG	10	03/14/2005	02/28/2013						
00517-0299-25		J2370		01/01/2002	07/31/2013	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	PHENYLEPHRINE HCL (S.D.V.) 10 MG/ML	1	ML	VL	IJ	ML	1	ML	1	01/01/2002	07/31/2013						
00517-0405-25		J2370		01/01/2002	06/30/2013	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	PHENYLEPHRINE HCL (VIAL) 10 MG/ML	5	ML	VL	IJ	ML	1	ML	1	01/01/2002	06/30/2013						
00517-0901-25		J0360		01/01/2002	02/28/2013	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HYDROCHLORIDE (S.D.V.) 20 MG/ML	1	ML	VL	IJ	ML	20	MG	1	01/01/2002	02/28/2013						
00517-1045-25		J1955		01/01/2002	12/31/2013	INJECTION, LEVOCARNITINE, PER 1 GM	LEVOCARNITINE (S.D.V.) 200 MG/ML	5	ML	VL	IV	ML	1	GM	0.2	01/01/2002	12/31/2013						
00517-1305-25		J1265		01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HCL (S.D.V.) 160 MG/ML	5	ML	VL	IV	ML	40	MG	4	01/01/2006	99/99/9999						
00517-1805-25		J1265		01/01/2006	12/31/2013	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HCL (S.D.V.) 40 MG/ML	5	ML	VL	IV	ML	40	MG	1	01/01/2006	12/31/2013						
00517-1905-25		J1265		01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HCL (S.D.V.) 80 MG/ML	5	ML	VL	IV	ML	40	MG	2	01/01/2006	99/99/9999						
00517-2310-05		J1756		05/01/2007	99/99/9999	INJECTION, IRON SUCROSE, 1 MG	VENOFER (5X10ML,SDV,USP,PF) 20 MG/ML	10	ML	VL	IV	ML	1	MG	20	05/01/2007	99/99/9999						
00517-2340-10		J1756		01/01/2003	99/99/9999	INJECTION, IRON SUCROSE, 1 MG	VENOFER (S.D.V.,PF) 20 MG/ML	5	ML	VL	IV	ML	1	MG	20	01/01/2003	99/99/9999						
00517-2340-25		J1756		10/01/2006	99/99/9999	INJECTION, IRON SUCROSE, 1 MG	VENOFER (25X5ML SDV,PF) 20 MG/ML	5	ML	VL	IV	ML	1	MG	20	10/01/2006	99/99/9999						
00517-2602-25		J3475		01/01/2002	03/31/2013	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.,PF) 500 MG/ML	2	ML	VL	IJ	ML	500	MG	1	01/01/2002	03/31/2013						
00517-2610-25		J3475		01/01/2002	03/31/2013	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.,PF) 500 MG/ML	10	ML	VL	IJ	ML	500	MG	1	01/01/2002	03/31/2013						
00517-2650-25		J3475		01/01/2002	08/31/2012	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.,PF) 500 MG/ML	50	ML	VL	IJ	ML	500	MG	1	01/01/2002	08/31/2012						
00517-2810-25		A4216		01/01/2004	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (S.D.V.,PF) 0.9% SODIUM CHLORIDE CONCENTRATE (S.D.V.) 23.4%	10	ML	VL	IV	ML	10	ML	0.1	01/01/2004	02/03/2016						
00517-2930-25		J7799		01/01/2002	02/28/2013	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	WATER FOR INJECTION (S.D.V.)	30	ML	VL	IV	ML	1	EA	1	01/01/2002	02/28/2013						
00517-3005-25		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.)	5	ML	VL	IV	ML	10	ML	0.1	01/01/2004	99/99/9999						
00517-3010-25		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.)	10	ML	VL	IV	ML	10	ML	0.1	01/01/2004	99/99/9999						
00517-3020-25		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.)	20	ML	VL	IV	ML	10	ML	0.1	01/01/2004	99/99/9999						
00517-3900-25		J0610		01/01/2002	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (VIAL,PF) 100 MG/ML	100	ML	VL	IV	ML	10	ML	0.1	01/01/2002	99/99/9999						
00517-3950-25		J0610		01/01/2002	01/31/2014	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (S.D.V.,PF) 100 MG/ML	50	ML	VL	IV	ML	10	ML	0.1	01/01/2002	01/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00517-4002-25		J2440		09/15/2003	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (S.D.V.) 30 MG/ML	2 ML	VL	IJ	ML		60 MG		0.5	09/15/2003	99/99/9999						
00517-4010-01		J2440		01/01/2002	04/03/2014	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (M.D.V.) 30 MG/ML	10 ML	VL	IJ	ML		60 MG		0.5	01/01/2002	04/03/2014						
00517-4050-25		J2150		01/01/2002	03/31/2014	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (S.D.V.,PF) 25%	50 ML	VL	IV	ML		50 ML		0.02	01/01/2002	03/31/2014						
00517-4201-25		J3410		01/01/2002	99/99/9999	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (S.D.V.) 25 MG/ML	1 ML	VL	IM	ML		25 MG		1	01/01/2002	99/99/9999						
00517-4601-25		J7643		01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	1 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-4601-25	KO	J7643	KO	01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	1 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-4602-25		J7643		01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	2 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-4602-25	KO	J7643	KO	01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	2 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-4605-25		J7643		01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	5 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-4605-25	KO	J7643	KO	01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	5 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-4620-25		J7643		01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	20 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-4620-25	KO	J7643	KO	01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	20 ML	VL	IJ	ML		1 MG		0.2	01/01/2002	99/99/9999						
00517-5601-25		J3410		01/01/2002	99/99/9999	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (S.D.V.) 50 MG/ML	1 ML	VL	IM	ML		25 MG		2	01/01/2002	99/99/9999						
00517-5602-25		J3410		01/01/2002	99/99/9999	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (S.D.V.) 50 MG/ML	2 ML	VL	IM	ML		25 MG		2	01/01/2002	99/99/9999						
00517-5610-25		J3410		01/01/2002	99/99/9999	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (M.D.V.) 50 MG/ML	10 ML	VL	IM	ML		25 MG		2	01/01/2002	99/99/9999						
00517-5702-25		J1940		01/01/2002	11/30/2013	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (S.D.V.) 10 MG/ML	2 ML	VL	IJ	ML		20 MG		0.5	01/01/2002	11/30/2013						
00517-5704-25		J1940		01/01/2002	12/31/2013	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (S.D.V.) 10 MG/ML	4 ML	VL	IJ	ML		20 MG		0.5	01/01/2002	12/31/2013						
00517-5710-25		J1940		01/01/2002	12/31/2013	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (S.D.V.) 10 MG/ML	10 ML	VL	IJ	ML		20 MG		0.5	01/01/2002	12/31/2013						
00517-7504-25		J7608		01/24/2003	99/99/9999	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 10%	4 ML	VL	IH	ML		1 GM		0.1	01/24/2003	99/99/9999						
00517-7504-25	KO	J7608	KO	01/24/2003	99/99/9999	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 10%	4 ML	VL	IH	ML		1 GM		0.1	01/24/2003	99/99/9999						
00517-7510-03		J7608		01/01/2002	99/99/9999	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 10%	10 ML	VL	IH	ML		1 GM		0.1	01/01/2002	99/99/9999						
00517-7510-03	KO	J7608	KO	01/01/2002	99/99/9999	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 10%	10 ML	VL	IH	ML		1 GM		0.1	01/01/2002	99/99/9999						
00517-7604-25		J7608		01/29/2003	99/99/9999	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 20%	4 ML	VL	IH	ML		1 GM		0.2	01/29/2003	99/99/9999						
00517-7604-25	KO	J7608	KO	01/29/2003	99/99/9999	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 20%	4 ML	VL	IH	ML		1 GM		0.2	01/29/2003	99/99/9999						
00517-7610-03		J7608		01/01/2002	05/31/2013	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 20%	10 ML	VL	IH	ML		1 GM		0.2	01/01/2002	05/31/2013						
00517-7610-03	KO	J7608	KO	01/01/2002	05/31/2013	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 20%	10 ML	VL	IH	ML		1 GM		0.2	01/01/2002	05/31/2013						
00517-7630-03		J7608		01/01/2002	04/30/2013	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 20%	30 ML	VL	IH	ML		1 GM		0.2	01/01/2002	04/30/2013						
00517-7630-03	KO	J7608	KO	01/01/2002	04/30/2013	FORM, PER GRAM ACETYLCHOLINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE	ACETYLCHOLINE (PF) 20%	30 ML	VL	IH	ML		1 GM		0.2	01/01/2002	04/30/2013						
00517-8905-10		J0210		02/26/2003	99/99/9999	INJECTION, METHYLDOPATE HCL, UP TO 250 MG	METHYLDOPATE HCL (S.D.V.) 50 MG/ML	5 ML	VL	IV	ML		250 MG		0.2	02/26/2003	99/99/9999						
00517-9120-25		J3490		03/12/2003	01/31/2014	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (M.D.V.) 250 MG/ML	20 ML	VL	IV	ML		1 EA		1	03/12/2003	01/31/2014						
00517-9702-25		J1790		01/01/2002	99/99/9999	INJECTION, DROPERIDOL, UP TO 5 MG	DROPERIDOL (S.D.V.) 2.5 MG/ML	2 ML	VL	IJ	ML		5 MG		0.5	01/01/2002	99/99/9999						
00536-0770-85		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHIST 12.5 MG/5 ML	480 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
00536-0770-97		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHIST 12.5 MG/5 ML	120 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
00536-3594-01		Q0163		01/01/2002	01/28/2015	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHIST 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/28/2015						
00536-3597-01		Q0163		01/01/2002	01/14/2015	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHIST (CAPTAB) 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/14/2015						
00536-3772-06		Q0163		01/01/2002	01/22/2015	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG NALOXONE HCL (21GX1 1/2' MINIJET,PF) 1 MG/ML	50 EA	BO	PO	EA		50 MG		1	01/01/2002	01/22/2015						
00548-1469-00		J2310		01/01/2002	04/03/2012	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (21GX1 1/2' MINIJET,PF) 1 MG/ML	2 ML	SR	IJ	ML		1 MG		1	01/01/2002	04/03/2012						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00548-1911-25		J2270		01/01/2002	08/31/2015	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (SRN,PREFILLED,PUMP-JET) 1 MG/ML	30	ML	SR	IJ	ML	10	MG	0.1	01/01/2002	08/31/2015							
00548-3301-00		J7799		01/01/2002	11/28/2012	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE (SRN,PREFILLED,LUER-JET) 50%	50	ML	SR	IV	ML	1	EA	1	01/01/2002	11/28/2012							
00548-3369-00		J2310		01/15/2002	04/12/2012	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (SRN,PREFILL,LUERJET,PF) 1 MG/ML	2	ML	SR	IJ	ML	1	MG	1	01/15/2002	04/12/2012							
00548-3390-00		J2001		01/01/2004	11/19/2012	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (SRN,PREFILLED,LUER-JET) 2%	5	ML	SR	IV	ML	10	MG	2	01/01/2004	11/19/2012							
00555-0059-02		Q0163		01/01/2002	08/19/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	08/19/2013							
00555-0059-05		Q0163		01/01/2002	08/19/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	1000	EA	BO	PO	EA	50	MG	1	01/01/2002	08/19/2013							
00555-0302-02		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013							
00555-0302-04		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013							
00555-0323-02		Q0177		01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	99/99/9999							
00555-0323-04		Q0177		01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	500	EA	BO	PO	EA	25	MG	1	01/01/2002	99/99/9999							
00555-0324-02		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	100	EA	BO	PO	EA	50	MG	2	01/01/2002	12/31/2013							
00555-0572-02	None			01/01/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	100	EA	BO	PO	EA	2.5	MG	1	01/01/1994	99/99/9999							
00555-0572-35	None			01/01/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	36	EA	BO	PO	EA	2.5	MG	1	01/01/1994	99/99/9999							
00555-0606-02	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 20 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
00555-0607-02	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
00555-0607-04	J8999			01/01/2002	09/27/2013	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	500	EA	BO	PO	EA	1	EA	1	01/01/2002	09/27/2013							
00555-0882-02	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDROXYUREA 500 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
00555-1131-11	J0895			09/05/2007	09/27/2013	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE 2 GM	1	EA	VL	IJ	EA	500	MG	4	09/05/2007	09/27/2013							
00555-1132-12	J0895			09/05/2007	02/05/2013	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE 500 MG	1	EA	VL	IJ	EA	500	MG	1	09/05/2007	02/05/2013							
00562-7805-01	J2790			09/01/2007	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, FULL DOSE, 300 MICROGRAMS (1500 I.U.)	RHOGAM ULTRA-FILTERED PLUS (PF,LATEX-FREE) 300 MCG	1	EA	SR	IM	EA	300	MCG	1	09/01/2007	99/99/9999							
00562-7805-05	J2790			09/01/2007	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, FULL DOSE, 300 MICROGRAMS (1500 I.U.)	RHOGAM ULTRA-FILTERED PLUS (PF,LATEX-FREE) 300 MCG	5	EA	SR	IM	EA	300	MCG	1	09/01/2007	99/99/9999							
00562-7805-25	J2790			09/01/2007	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, FULL DOSE, 300 MICROGRAMS (1500 I.U.)	RHOGAM ULTRA-FILTERED PLUS (PF,LATEX-FREE) 300 MCG	25	EA	SR	IM	EA	300	MCG	1	09/01/2007	99/99/9999							
00562-7806-01	J2788			09/01/2007	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, MINIDOSE, 50 MICROGRAMS (250 I.U.)	MICRHOGAM ULTRA-FILTERED PLUS (PF,LATEX-FREE) 50 MCG	1	EA	SR	IM	EA	50	MCG	1	09/01/2007	99/99/9999							
00562-7806-05	J2788			09/01/2007	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, MINIDOSE, 50 MICROGRAMS (250 I.U.)	MICRHOGAM ULTRA-FILTERED PLUS (PF,LATEX-FREE) 50 MCG	5	EA	SR	IM	EA	50	MCG	1	09/01/2007	99/99/9999							
00562-7806-25	J2788			09/01/2007	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, MINIDOSE, 50 MICROGRAMS (250 I.U.)	MICRHOGAM ULTRA-FILTERED PLUS (PF,LATEX-FREE) 50 MCG	25	EA	SR	IM	EA	50	MCG	1	09/01/2007	99/99/9999							
00574-0421-25	J1700			01/01/2002	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P. MICRONIZED)	1	EA	BO	NA	GM	25	MG	40	01/01/2002	99/99/9999							
00574-0820-01	J1080			12/21/2007	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (1X1ML,USP) 200 MG/ML	1	ML	VL	IM	ML	200	MG	1	12/21/2007	12/31/2014							
54879-0022-01	None			05/08/2018	99/99/9999	CYCLOPHOSPHAMIDE, 50 MG, ORAL	CYCLOPHOSPHAMIDE 50 MG	100	EA	BO	PO	EA	50	MG	1	05/08/2018	99/99/9999							
00574-0820-10	J1080			12/21/2007	11/11/2013	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (1X10ML,USP) 200 MG/ML	10	ML	VL	IM	ML	200	MG	1	12/21/2007	11/11/2013							
00574-0823-01	J0706			09/21/2006	04/21/2014	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATE (USP,PF) 20 MG/ML	3	ML	VL	IV	ML	5	MG	4	09/21/2006	04/21/2014							
00574-0823-81	J0706			09/28/2007	09/18/2014	INJECTION, CAFFEINE CITRATE, 5MG	NOVAPLUS CAFFEINE CITRATE (USP,10X3ML,PF) 20 MG/ML	3	ML	VL	IV	ML	5	MG	4	09/28/2007	09/18/2014							
00574-0850-05	J1110			08/04/2003	99/99/9999	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG	DIHYDROERGOTAMINE MESYLATE (AMP) 1 MG/ML	1	ML	AM	IJ	ML	1	MG	1	08/04/2003	99/99/9999							
50742-0494-17	J0641			09/01/2018	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (PF) 10 MG/1 ML	17.5	ML	VL	IV	ML	0.5	MG	20	09/01/2018	99/99/9999							
50742-0495-25	J0641			09/01/2018	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (PF) 10 MG/1 ML	25	ML	VL	IV	ML	0.5	MG	20	09/01/2018	99/99/9999							
00574-0850-10	J1110			03/15/2004	99/99/9999	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG	DIHYDROERGOTAMINE MESYLATE (AMP) 1 MG/ML	1	ML	AM	IJ	ML	1	MG	1	03/15/2004	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00574-0858-01	J0770			03/11/2005	06/30/2018	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE SODIUM (VIAL,STERILE) 150 MG	1 EA	VL	U	EA		150 MG		1	03/11/2005	06/30/2018						
00574-7226-12	J8498			01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	COMPRO 25 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
67877-0266-05	J7517			08/01/2013	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	500 EA	BO	PO	EA		250 MG		1	08/01/2013	99/99/9999						
42291-0166-60	None			05/14/2018	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM COATED) 150 MG	60 EA	BO	PO	EA		150 MG		1	05/14/2018	99/99/9999						
00069-1306-10	Q5106			01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS HYDROXYZYNE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	RETACRIT (PF) 3000 U/1 ML	1 ML	VL	U	ML		1000 U		3	01/01/2019	99/99/9999						
00591-0800-01	Q0177			09/18/2006	99/99/9999	HYDROXYZYNE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZYNE PAMOATE (USP) 25 MG	100 EA	BO	PO	EA		25 MG		1	09/18/2006	99/99/9999						
00591-0800-05	Q0177			09/18/2006	99/99/9999	HYDROXYZYNE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZYNE PAMOATE (USP) 25 MG	500 EA	BO	PO	EA		25 MG		1	09/18/2006	99/99/9999						
00591-0801-01	Q0178			01/01/2002	12/31/2013	HYDROXYZYNE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZYNE PAMOATE 50 MG	100 EA	BO	PO	EA		50 MG		1	09/18/2006	12/31/2013	01/01/2002	08/17/2005	1			
00591-0801-05	Q0178			01/01/2002	12/31/2013	HYDROXYZYNE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZYNE PAMOATE 50 MG	500 EA	BO	PO	EA		50 MG		1	09/18/2006	12/31/2013	01/01/2002	08/09/2005	1			
00591-3128-79	J2675			12/17/2002	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE IN SESAME OIL (VIAL) 50 MG/ML	10 ML	VL	IM	ML		50 MG		1	12/17/2002	99/99/9999						
00591-3221-26	J3130			03/09/2004	12/31/2014	INJECTION, TESTOSTERONE ENANTHATE, UP TO 200 MG	TESTOSTERONE ENANTHATE 200 MG/ML	5 ML	VL	IM	ML		200 MG		1	03/09/2004	12/31/2014						
00591-3222-47	J2360			09/07/2004	11/05/2018	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORPHENADRINE CITRATE 30 MG/ML	2 ML	AM	U	ML		60 MG		0.5	09/07/2004	11/05/2018						
00591-3223-79	J1080			03/29/2004	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (M.D.V.) 200 MG/ML	10 ML	VL	IM	ML		200 MG		1	03/29/2004	12/31/2014						
00591-3433-30	J7620			01/02/2008	05/12/2013	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	30 ML	PC	IH	ML		3 MG	0.33333	01/02/2008	05/12/2013							
00591-3433-60	J7620			01/02/2008	05/12/2013	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	60 ML	PC	IH	ML		3 MG	0.33333	01/02/2008	05/12/2013							
00591-3467-53	J7613			04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 0.021%	3 ML	PC	IH	ML		1 MG	0.21	04/01/2008	99/99/9999							
00591-3467-53	KO J7613	KO		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 0.021%	3 ML	PC	IH	ML		1 MG	0.21	04/01/2008	99/99/9999							
00591-3468-53	J7613			04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 0.042%	3 ML	PC	IH	ML		1 MG	0.42	04/01/2008	99/99/9999							
00591-3468-53	KO J7613	KO		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 0.042%	3 ML	PC	IH	ML		1 MG	0.42	04/01/2008	99/99/9999							
00591-5052-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	100 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
00591-5052-10	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	1000 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
00591-5307-01	Q0170			04/15/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100 EA	BO	PO	EA		25 MG		1	04/15/2002	12/31/2013						
00591-5307-10	Q0170			04/15/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	1000 EA	BO	PO	EA		25 MG		1	04/15/2002	12/31/2013						
00591-5319-01	Q0170			04/15/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	100 EA	BO	PO	EA		25 MG		2	04/15/2002	12/31/2013						
00591-5442-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
00591-5442-05	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	500 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
00591-5442-10	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	1000 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
00591-5443-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	100 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
00591-5443-05	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	500 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
00591-5443-10	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	1000 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
00603-0241-18	Q0163			06/05/2007	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	Q-DRYL 25 MG	24 EA	BO	PO	EA		50 MG		0.5	06/05/2007	06/30/2017						
00603-0823-54	Q0163			01/01/2002	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	Q-DRYL (AF,CHERRY) 12.5 MG/5 ML	120 ML	BO	PO	ML		50 MG		0.05	01/01/2002	06/30/2017						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00603-0823-58		Q0163		01/01/2002	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	Q-DRYL 12.5 MG/5 ML	473	ML	BO	PO	ML	50	MG	0.05	01/01/2002	06/30/2017						
00603-0823-81		Q0163		07/25/2002	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	Q-DRYL 12.5 MG/5 ML	240	ML	BO	PO	ML	50	MG	0.05	07/25/2002	06/30/2017						
00603-0823-94		Q0163		01/01/2002	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	Q-DRYL (UNBOXED,AF,CHERRY) 12.5 MG/5 ML	120	ML	BO	PO	ML	50	MG	0.05	01/01/2002	06/30/2017						
00603-0860-54		Q0163		01/01/2002	08/31/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	QUENALIN 12.5 MG/5 ML	120	ML	BO	PO	ML	50	MG	0.05	01/01/2002	08/31/2016						
00603-1584-54		Q0170		05/12/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE PLAIN (USP) 6.25 MG/5 ML	118	ML	BO	PO	ML	25	MG	0.05	05/12/2006	12/31/2013						
00603-1584-58		Q0170		05/12/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE PLAIN (USP) 6.25 MG/5 ML	473	ML	BO	PO	ML	25	MG	0.05	05/12/2006	12/31/2013						
00603-3339-21		Q0163		05/24/2007	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE (USP) 25 MG	100	EA	BO	PO	EA	50	MG	0.5	05/24/2007	06/30/2017						
00603-3339-32		Q0163		06/05/2007	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE (USP) 25 MG	1000	EA	BO	PO	EA	50	MG	0.5	06/05/2007	06/30/2017						
00603-3340-21		Q0163		04/03/2007	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE (USP) 50 MG	100	EA	BO	PO	EA	50	MG	1	04/03/2007	06/30/2017						
00603-3340-32		Q0163		04/03/2007	06/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE (USP) 50 MG	1000	EA	BO	PO	EA	50	MG	1	04/03/2007	06/30/2017						
00603-4593-15		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (DOSE PACK) 4 MG	21	EA	DP	PO	EA	4	MG	1	01/01/2002	99/99/9999						
00603-4593-21		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	100	EA	BO	PO	EA	4	MG	1	01/01/2002	99/99/9999						
00603-5090-21		Q0175		01/01/2002	07/15/2012	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 2 MG	100	EA	BO	PO	EA	4	MG	0.5	07/02/2009	07/15/2012	01/01/2002	09/19/2008	0.5			
00603-5090-28		Q0175		01/01/2002	04/16/2012	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 2 MG	500	EA	BO	PO	EA	4	MG	0.5	07/02/2009	04/16/2012	01/01/2002	09/19/2008	0.5			
00603-5091-21		Q0175		01/01/2002	04/02/2012	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	100	EA	BO	PO	EA	4	MG	1	07/02/2009	04/02/2012	01/01/2002	09/19/2008	1			
00603-5092-21		Q0176		01/01/2002	05/07/2012	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	100	EA	BO	PO	EA	8	MG	1	07/02/2009	05/07/2012	01/01/2002	09/19/2008	1			
00603-5092-28		Q0176		01/01/2002	04/30/2012	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	500	EA	BO	PO	EA	8	MG	1	07/02/2009	04/30/2012	01/01/2002	09/19/2008	1			
00603-5093-21		Q0176		01/01/2002	05/14/2012	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 16 MG	100	EA	BO	PO	EA	8	MG	2	07/02/2009	05/14/2012	01/01/2002	09/19/2008	2			
00603-5335-21		J7506		01/03/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 1 MG	100	EA	BO	PO	EA	5	MG	0.2	01/03/2005	12/31/2015						
00603-5335-32		J7506		01/03/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 1 MG	1000	EA	BO	PO	EA	5	MG	0.2	01/03/2005	12/31/2015						
00603-5336-21		J7506		01/03/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 2.5 MG	100	EA	BO	PO	EA	5	MG	0.5	01/03/2005	12/31/2015						
00603-5337-15		J7506		08/20/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (DOSE PACK) 5 MG	21	EA	DP	PO	EA	5	MG	1	08/20/2003	12/31/2015						
00603-5337-21		J7506		01/16/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	100	EA	BO	PO	EA	5	MG	1	01/16/2003	12/31/2015						
00603-5337-31		J7506		08/20/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (DOSE PACK) 5 MG	48	EA	DP	PO	EA	5	MG	1	08/20/2003	12/31/2015						
00603-5337-32		J7506		01/16/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	1000	EA	BO	PO	EA	5	MG	1	01/16/2003	12/31/2015						
00603-5338-15		J7506		03/06/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (DOSE PACK) 10 MG	21	EA	DP	PO	EA	5	MG	2	03/06/2003	12/31/2015						
00603-5338-21		J7506		01/30/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/30/2003	12/31/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00603-5338-28	J7506			01/30/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	500	EA	BO	PO	EA	5	MG	2	01/30/2003	12/31/2015						
00603-5338-31	J7506			04/02/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (DOSE PACK) 10 MG	48	EA	DP	PO	EA	5	MG	2	04/02/2003	12/31/2015						
00603-5338-32	J7506			01/30/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	1000	EA	BO	PO	EA	5	MG	2	01/30/2003	12/31/2015						
00603-5339-21	J7506			09/10/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	100	EA	BO	PO	EA	5	MG	4	09/10/2003	12/31/2015						
00603-5339-28	J7506			09/10/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	500	EA	BO	PO	EA	5	MG	4	09/10/2003	12/31/2015						
00603-5339-32	J7506			09/10/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	1000	EA	BO	PO	EA	5	MG	4	09/10/2003	12/31/2015						
00603-5437-21	Q0169			08/25/2006	01/09/2017	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 12.5 MG	100	EA	BO	PO	EA	12.5	MG	1	08/25/2006	01/09/2017						
00603-5438-21	Q0170			08/25/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	100	EA	BO	PO	EA	25	MG	1	08/25/2006	12/31/2013						
00603-5438-32	Q0170			08/25/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	1000	EA	BO	PO	EA	25	MG	1	08/25/2006	12/31/2013						
00603-5439-21	Q0170			08/25/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 50 MG	100	EA	BO	PO	EA	25	MG	2	08/25/2006	12/31/2013						
00641-0121-21	J1170			12/08/2004	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (VIAL, DOSETTE) 2 MG/ML	1	ML	VL	IJ	ML	4	MG	0.5	12/08/2004	99/99/9999						
00641-0121-25	J1170			01/01/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (VIAL, DOSETTE) 2 MG/ML	1	ML	VL	IJ	ML	4	MG	0.5	01/01/2002	99/99/9999						
00641-0367-21	J1100			12/08/2004	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (VIAL, DOSETTE) 10 MG/ML	1	ML	VL	IJ	ML	1	MG	10	12/08/2004	99/99/9999						
00641-0376-21	J1200			12/08/2004	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (DOSETTE VIAL) 50 MG/ML	1	ML	VL	IJ	ML	50	MG	1	12/08/2004	99/99/9999						
00641-0476-21	J2560			12/08/2004	99/99/9999	INJECTION, PHENOBARBITAL SODIUM, UP TO 120 MG	PHENOBARBITAL SODIUM (VIAL, DOSETTE) 65 MG/ML	1	ML	VL	IJ	ML	120	MG	0.54166	12/08/2004	99/99/9999						
00641-0477-21	J2560			12/08/2004	99/99/9999	INJECTION, PHENOBARBITAL SODIUM, UP TO 120 MG	PHENOBARBITAL SODIUM (DOSETTE VIAL) 130 MG/ML	1	ML	VL	IJ	ML	120	MG	1.08333	12/08/2004	99/99/9999						
00641-0493-21	J1165			12/08/2004	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (DOSETTE,VIAL) 50 MG/ML	2	ML	VL	IV	ML	50	MG	1	12/08/2004	99/99/9999						
00641-0928-21	J2550			12/08/2004	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (DOSETTE,VIAL) 25 MG/ML	1	ML	VL	IJ	ML	50	MG	0.5	12/08/2004	99/99/9999						
00641-0929-21	J2550			12/08/2004	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (DOSETTE,VIAL) 50 MG/ML	1	ML	VL	IJ	ML	50	MG	1	12/08/2004	99/99/9999						
00641-0929-25	J2550			12/27/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (DOSETTE,VIAL) 50 MG/ML	1	ML	VL	IJ	ML	50	MG	1	12/27/2002	99/99/9999						
00641-0948-31	J2550			12/08/2004	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL NOVAPLUS (AMP,DOSETTE) 25 MG/ML	1	ML	AM	IJ	ML	50	MG	0.5	12/08/2004	99/99/9999						
00641-0949-31	J2550			05/05/2007	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL NOVAPLUS (DOSETTE) 50 MG/ML	1	ML	AM	IJ	ML	50	MG	1	05/05/2007	99/99/9999						
00641-0955-21	J2550			05/05/2007	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL NOVAPLUS (DOSETTE) 25 MG/ML	1	ML	VL	IJ	ML	50	MG	0.5	05/05/2007	99/99/9999						
00641-0956-21	J2550			05/05/2007	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL NOVAPLUS (DOSETTE) 50 MG/ML	1	ML	VL	IJ	ML	50	MG	1	05/05/2007	99/99/9999						
00641-1397-31	J3230			05/05/2007	99/99/9999	INJECTION, CHLORPROMAZINE HCL, UP TO 50 MG	CHLORPROMAZINE HCL (USP) 25 MG/ML	1	ML	AM	IJ	ML	50	MG	0.5	05/05/2007	99/99/9999						
00641-1398-35	J3230			01/01/2002	99/99/9999	INJECTION, CHLORPROMAZINE HCL, UP TO 50 MG	CHLORPROMAZINE HCL (AMP, DOSETTE) 25 MG/ML	2	ML	AM	IJ	ML	50	MG	0.5	01/01/2002	99/99/9999						
00641-1410-31	J1160			05/05/2007	99/99/9999	INJECTION, DIGOXIN, UP TO 0.5 MG	DIGOXIN (USP) 0.25 MG/ML	2	ML	AM	IV	ML	0.5	MG	0.5	05/05/2007	99/99/9999						
00641-1495-31	J2550			05/05/2007	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (USP) 25 MG/ML	1	ML	AM	IJ	ML	50	MG	0.5	05/05/2007	99/99/9999						
00641-1496-31	J2550			05/05/2007	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (USP) 50 MG/ML	1	ML	AM	IJ	ML	50	MG	1	05/05/2007	99/99/9999						
00641-2341-39	J1170			05/05/2007	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (USP) 2 MG/ML	1	ML	NA	IJ	ML	4	MG	0.5	05/05/2007	99/99/9999						
00641-2341-41	J1170			01/01/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (M.D.V.) 2 MG/ML	20	ML	VL	IJ	ML	4	MG	0.5	01/01/2002	99/99/9999						
00641-2555-41	J1165			05/05/2007	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (USP) 50 MG/ML	1	ML	VL	IV	ML	50	MG	1	05/05/2007	99/99/9999						
00641-2569-41	J1245			05/05/2007	99/99/9999	INJECTION, DIPYRIDAMOLE, PER 10 MG	DIPYRIDAMOLE (SDV) 5 MG/ML	10	ML	VL	IV	ML	10	MG	0.5	05/05/2007	99/99/9999						
00703-0031-01	J1030			03/09/2005	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (SDV) 40 MG/ML	1	ML	VL	IJ	ML	40	MG	1	03/09/2005	99/99/9999						
00703-0031-04	J1030			03/09/2005	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (SDV) 40 MG/ML	1	ML	VL	IJ	ML	40	MG	1	03/09/2005	99/99/9999						
00703-0043-01	J1030			10/31/2006	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (MDV,USP) 40 MG/ML	5	ML	VL	IJ	ML	40	MG	1	10/31/2006	99/99/9999						
00703-0045-01	J1030			10/31/2006	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (MDV,USP) 40 MG/ML	10	ML	VL	IJ	ML	40	MG	1	10/31/2006	99/99/9999						
00703-0051-01	J1040			03/09/2005	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 80 MG	METHYLPREDNISOLONE ACETATE (SDV) 80 MG/ML	1	ML	VL	IJ	ML	80	MG	1	03/09/2005	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00703-0051-04	J1040			03/09/2005	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 80 MG	METHYLPREDNISOLONE ACETATE (SDV) 80 MG/ML	1 ML	VL	IJ	ML		80 MG		1	03/09/2005	99/99/9999						
00703-0063-01	J1040			10/31/2006	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 80 MG	METHYLPREDNISOLONE ACETATE (MDV,USP) 80 MG/ML	5 ML	VL	IJ	ML		80 MG		1	10/31/2006	99/99/9999						
00703-0346-03	J0696			12/21/2007	10/12/2012	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP,SINGLE-DOSE) 2 GM MORPHINE SULFATE (PF-LATEX-FREE) 10 MG/1 ML	1 EA	VL	IJ	EA		250 MG		8	12/21/2007	10/12/2012						
63323-0451-01	J2270			05/23/2018	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	CEFTRIAXONE (USP,PHARMACY BULK PCKGE) 10 GM	1 EA	VL	IV	EA		250 MG		40	12/21/2007	01/17/2013						
00703-0359-01	J0696			12/21/2007	01/17/2013	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	LEVOCARNITINE (VIAL) 200 MG/ML	5 ML	VL	IV	ML		1 GM		0.2	01/01/2002	05/02/2017						
00703-0404-02	J1955			01/01/2002	05/02/2017	INJECTION, LEVOCARNITINE, PER 1 GM	LEVOCARNITINE (VIAL) 200 MG/ML	12.5 ML	VL	IV	ML		1 GM		0.2	01/01/2002	05/02/2017						
00703-0405-02	J1955			01/01/2002	05/02/2017	INJECTION, LEVOCARNITINE, PER 1 GM	FLUCONAZOLE IV 400 MG/200 ML	200 ML	VL	IV	ML		200 MG		0.01	08/02/2004	09/05/2013						
00703-1010-09	J1450			08/02/2004	09/05/2013	INJECTION, FLUCONAZOLE, 200 MG	FLUCONAZOLE IV 200 MG/100 ML	100 ML	VL	IV	ML		200 MG		0.01	08/02/2004	09/05/2013						
00703-1019-09	J1450			08/02/2004	09/05/2013	INJECTION, FLUCONAZOLE, 200 MG	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	1 ML	VL	IV	ML		1.25 MCG		400	01/01/2002	99/99/9999						
00703-1501-02	J0270			01/01/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	EPOPROSTENOL SODIUM 0.5 MG	1 EA	VL	IV	EA		0.5 MG		1	04/23/2008	99/99/9999						
00703-1985-01	J1325			04/23/2008	99/99/9999	INJECTION, EPOPROSTENOL, 0.5 MG	EPOPROSTENOL SODIUM 1.5 MG	1 EA	VL	IV	EA		0.5 MG		3	04/23/2008	99/99/9999						
00703-1995-01	J1325			04/23/2008	99/99/9999	INJECTION, EPOPROSTENOL, 0.5 MG	PROMETHAZINE HCL 25 MG/ML	1 ML	VL	IJ	ML		50 MG		0.5	09/30/2002	99/99/9999						
00703-2191-04	J2550			09/30/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL 50 MG/ML	1 ML	VL	IJ	ML		50 MG		1	09/30/2002	99/99/9999						
00703-2201-04	J2550			09/30/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	ADRUCIL (S.D.V.) 50 MG/ML	10 ML	VL	IV	ML		500 MG		0.1	09/02/2003	99/99/9999						
00703-3015-13	J9190			09/02/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	ADRUCIL (PHARMACY BULK PACKAGE) 50 MG/ML	50 ML	VL	IV	ML		500 MG		0.1	09/02/2003	99/99/9999						
00703-3018-12	J9190			09/02/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	ADRUCIL (PHARMACY BULK PACKAGE) 50 MG/ML	100 ML	VL	IV	ML		500 MG		0.1	09/02/2003	99/99/9999						
00703-3019-12	J9190			09/02/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	ADRUCIL (PHARMACY BULK PACKAGE) 50 MG/ML	100 ML	VL	IV	ML		500 MG		0.1	09/02/2003	99/99/9999						
00703-3067-11	J9178			08/09/2007	11/30/2017	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (SDV,PF) 2 MG/ML	25 ML	VL	IV	ML		2 MG		1	08/09/2007	11/30/2017						
00703-3069-11	J9178			08/09/2007	03/31/2017	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (SDV,PF) 2 MG/ML	100 ML	VL	IV	ML		2 MG		1	08/09/2007	03/31/2017						
00703-3154-01	J9040			01/01/2002	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN SULFATE (S.D.V.) 15 U	1 EA	VL	IJ	EA		15 U		1	01/01/2002	99/99/9999						
00703-3155-01	J9040			01/01/2002	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN SULFATE (S.D.V.) 30 U	1 EA	VL	IJ	EA		15 U		2	01/01/2002	99/99/9999						
00703-3246-11	J9045			06/24/2004	10/17/2016	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (M.D.V.) 10 MG/ML	15 ML	VL	IV	ML		50 MG		0.2	06/24/2004	10/17/2016						
00703-3249-11	J9045			11/17/2005	05/24/2016	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (AQUEOUS SOLUTION) 10 MG/ML	60 ML	VL	IV	ML		50 MG		0.2	11/17/2005	05/24/2016						
00703-3264-01	J9045			06/24/2004	10/17/2016	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN 50 MG	1 EA	VL	IV	EA		50 MG		1	06/24/2004	10/17/2016						
00703-3266-01	J9045			06/24/2004	10/17/2016	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (VIAL) 150 MG	1 EA	VL	IV	EA		50 MG		3	06/24/2004	10/17/2016						
00703-3268-71	J9045			05/01/2006	10/17/2016	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN 450 MG	1 EA	VL	IV	EA		50 MG		9	05/01/2006	10/17/2016						
00703-3301-04	J2354			11/14/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (1MLX25 VIALS) 50 MCG/ML	1 ML	VL	IJ	ML		25 MCG		2	11/14/2005	99/99/9999						
00703-3311-04	J2354			11/14/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (1MLX25 VIALS) 100 MCG/ML	1 ML	VL	IJ	ML		25 MCG		4	11/14/2005	99/99/9999						
00703-3321-04	J2354			11/14/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (1MLX25 VIALS) 500 MCG/ML	1 ML	VL	IJ	ML		25 MCG		20	11/14/2005	99/99/9999						
00703-3333-01	J2354			11/23/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE 200 MCG/ML	5 ML	VL	IJ	ML		25 MCG		8	11/23/2005	99/99/9999						
00703-3343-01	J2354			11/23/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE 1000 MCG/ML	5 ML	VL	IJ	ML		25 MCG		40	11/23/2005	99/99/9999						
00703-3427-11	J9208			07/26/2007	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE 1 GM	1 EA	VL	IV	EA		1 GM		1	07/26/2007	99/99/9999						
00703-3429-11	J9208			07/26/2007	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE 3 GM	1 EA	VL	IV	EA		1 GM		3	07/26/2007	99/99/9999						
00703-4014-19	J9218			01/01/2002	99/99/9999	LEUPROLIDE ACETATE, PER 1 MG	LEUPROLIDE ACETATE (M.D.V.) 5 MG/ML	2.8 ML	VL	SC	ML		1 MG		5	01/01/2002	99/99/9999						
00703-4075-59	J2430			11/08/2005	03/26/2015	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (S.D.V.) 3 MG/ML	10 ML	VL	IV	ML		30 MG		0.1	11/08/2005	03/26/2015						
00703-4085-51	J2430			11/08/2005	99/99/9999	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM 9 MG/ML	10 ML	VL	IV	ML		30 MG		0.3	11/08/2005	99/99/9999						
00703-4100-48	J9999			04/08/2002	01/03/2017	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	IFOSFAMIDE/MESNA (COMBO-PACK) 5 GM-3 GM	1 EA	BX	IV	EA		1 EA		1	04/08/2002	01/03/2017						
00703-4100-58	J9999			04/08/2002	01/03/2017	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	IFOSFAMIDE/MESNA (COMBO-PACK) 10 GM-10 GM	1 EA	BX	IV	EA		1 EA		1	04/08/2002	01/03/2017						
00703-4100-68	J9999			04/08/2002	01/03/2017	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	IFOSFAMIDE/MESNA (COMBO-PACK) 6 GM-6 GM	1 EA	BX	IV	EA		1 EA		1	04/08/2002	01/03/2017						
00703-4154-11	J9211			09/24/2002	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (S.D.V.) 1 MG/ML	5 ML	VL	IV	ML		5 MG		0.2	09/24/2002	99/99/9999						
00703-4155-11	J9211			09/24/2002	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (S.D.V.) 1 MG/ML	10 ML	VL	IV	ML		5 MG		0.2	09/24/2002	99/99/9999						
00703-4156-11	J9211			09/24/2002	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (S.D.V.) 1 MG/ML	20 ML	VL	IV	ML		5 MG		0.2	09/24/2002	99/99/9999						
00703-4182-01	J9390			02/10/2003	11/30/2012	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (S.D.V.,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	02/10/2003	11/30/2012						
00703-4182-91	J9390			05/01/2006	04/03/2013	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (NOV,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	05/01/2006	04/03/2013						
00703-4183-01	J9390			02/10/2003	04/30/2013	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (S.D.V.,PF) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	02/10/2003	04/30/2013						

NDC	NDC Mod	HPCS	HPCS Mod	Relationship Start Date	Relationship End Date	HPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HPCS Amount #1	HPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00703-4183-91	J9390			05/01/2006	04/03/2013	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (NOV,PF) 10 MG/ML	5 ML	VL	IV	ML	10 MG	1		05/01/2006	04/03/2013							
00703-4244-01	J9045			05/01/2006	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (1X5ML) 10 MG/ML	5 ML	VL	IV	ML	50 MG	0.2		05/01/2006	99/99/9999							
00703-4246-01	J9045			05/01/2006	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (1X15ML) 10 MG/ML	15 ML	VL	IV	ML	50 MG	0.2		05/01/2006	99/99/9999							
00703-4248-01	J9045			02/01/2006	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN 10 MG/ML	45 ML	VL	IV	ML	50 MG	0.2		02/01/2006	99/99/9999							
00703-4402-11	J9370			01/01/2002	99/99/9999	VINCRIStINE SULFATE, 1 MG	VINCRIStINE SULFATE (S.D.V.) 1 MG/ML	1 ML	VL	IV	ML	1 MG	1		01/01/2002	99/99/9999							
00703-4412-11	J9370			01/01/2002	99/99/9999	VINCRIStINE SULFATE, 1 MG	VINCRIStINE SULFATE (S.D.V.) 1 MG/ML	2 ML	VL	IV	ML	1 MG	1		01/01/2002	99/99/9999							
00703-4432-11	J9206			02/28/2008	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML SINGLE DOSE) 20 MG/ML	2 ML	VL	IV	ML	20 MG	1		02/28/2008	99/99/9999							
00703-4636-01	J9320			12/03/2003	99/99/9999	INJECTION, STREPTOZOcIN, 1 GRAM	ZANOSAR 1 GM	1 EA	VL	IV	EA	1 GM	1		12/03/2003	99/99/9999							
00703-4680-01	J9293			04/11/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (MDV,PF) 2 MG/ML	12.5 ML	VL	IV	ML	5 MG	0.4		04/11/2006	99/99/9999							
00703-4685-01	J9293			04/11/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (MDV,PF) 2 MG/ML	10 ML	VL	IV	ML	5 MG	0.4		04/11/2006	99/99/9999							
00703-4686-01	J9293			04/11/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (MDV,PF) 2 MG/ML	15 ML	VL	IV	ML	5 MG	0.4		04/11/2006	99/99/9999							
00703-4805-03	J9209			02/22/2002	04/27/2015	INJECTION, MESNA, 200 MG	MESNA (M.D.V.) 100 MG/ML	10 ML	VL	IV	ML	200 MG	0.5		02/22/2002	04/27/2015							
00703-4852-11	J9185			05/02/2007	99/99/9999	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE (SDV) 25 MG/ML	2 ML	VL	IV	ML	50 MG	0.5		05/02/2007	99/99/9999							
00703-5051-03	J2597			01/01/2002	99/99/9999	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DESMOPRESSIN ACETATE (VIAL) 4 MCG/ML	1 ML	VL	IJ	ML	1 MCG	4		01/01/2002	99/99/9999							
00703-5054-01	J2597			01/01/2002	99/99/9999	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DESMOPRESSIN ACETATE (M.D.V.) 4 MCG/ML	10 ML	VL	IJ	ML	1 MCG	4		01/01/2002	99/99/9999							
00703-5140-01	J0640			01/01/2002	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (VIAL,PF) 100 MG	1 EA	VL	IJ	EA	50 MG	2		01/01/2002	99/99/9999							
00703-5145-01	J0640			01/01/2002	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (PF) 350 MG	1 EA	VL	IJ	EA	50 MG	7		01/01/2002	99/99/9999							
00703-5233-13	J9150			01/27/2003	99/99/9999	INJECTION, DAUNORUBICIN, 10 MG	DAUNORUBICIN HCL (S.D.V.,PF) 5 MG/ML	4 ML	VL	IV	ML	10 MG	0.5		01/27/2003	99/99/9999							
00703-5653-01	J9181			01/01/2002	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V. POLYMER) 20 MG/ML	5 ML	VL	IV	ML	10 MG	2		01/01/2002	99/99/9999							
00703-5656-01	J9181			01/01/2002	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V. POLYMER) 20 MG/ML	25 ML	VL	IV	ML	10 MG	2		01/01/2002	99/99/9999							
00703-5657-01	J9181			01/01/2002	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V.) 20 MG/ML	50 ML	VL	IV	ML	10 MG	2		01/01/2002	99/99/9999							
00703-5854-01	J9185			09/12/2003	99/99/9999	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE 50 MG	1 EA	VL	IV	EA	50 MG	1		09/12/2003	99/99/9999							
00703-6121-01	J1080			04/16/2007	10/19/2012	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (USP,MDV) 200 MG/ML	1 ML	VL	IM	ML	200 MG	1		04/16/2007	10/19/2012							
00703-6125-01	J1080			04/16/2007	10/19/2012	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (USP,MDV) 200 MG/ML	10 ML	VL	IM	ML	200 MG	1		04/16/2007	10/19/2012							
00703-6801-01	J1055			09/13/2004	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	MEDROXYPROGESTERONE ACETATE (ODOR-FREE) 150 MG/ML	1 ML	VL	IM	ML	150 MG	1		09/13/2004	12/31/2012							
00703-6801-04	J1055			09/13/2004	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	MEDROXYPROGESTERONE ACETATE (ODOR-FREE) 150 MG/ML	1 ML	VL	IM	ML	150 MG	1		09/13/2004	12/31/2012							
00703-7011-03	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (VIAL) 50 MG/ML	1 ML	VL	IM	ML	50 MG	1		01/01/2002	99/99/9999							
00703-7013-01	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (M.D.V.) 50 MG/ML	5 ML	VL	IM	ML	50 MG	1		01/01/2002	99/99/9999							
00703-7021-03	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (VIAL) 100 MG/ML	1 ML	VL	IM	ML	50 MG	2		01/01/2002	99/99/9999							
00703-7023-01	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (M.D.V.) 100 MG/ML	5 ML	VL	IM	ML	50 MG	2		01/01/2002	99/99/9999							
00703-7041-03	J1630			01/18/2002	05/09/2012	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL LACTATE (S.D.V.) 5 MG/ML	1 ML	VL	IM	ML	5 MG	1		01/18/2002	05/09/2012							
00703-7221-04	J2405			11/22/2006	10/08/2018	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (SDV,USP,25X2ML) 2 MG/ML	2 ML	VL	IJ	ML	1 MG	2		11/22/2006	10/08/2018							
00703-7226-01	J2405			11/22/2006	10/08/2018	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV,USP) 2 MG/ML	20 ML	VL	IJ	ML	1 MG	2		11/22/2006	10/08/2018							
00703-7226-03	J2405			11/22/2006	10/08/2018	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV,USP,10X20ML) 2 MG/ML	20 ML	VL	IJ	ML	1 MG	2		11/22/2006	10/08/2018							
00703-7239-39	J2405			11/22/2006	11/29/2012	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (SINGLE DOSE,6X50ML,PF) 32 MG/50 ML	50 ML	FC	IV	ML	1 MG	0.64		11/22/2006	11/29/2012							
00703-9032-03	J0278			01/01/2006	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (S.D.V.) 250 MG/ML	2 ML	VL	IJ	ML	100 MG	2.5		01/01/2006	99/99/9999							
00703-9040-03	J0278			01/01/2006	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (VIAL) 250 MG/ML	4 ML	VL	IJ	ML	100 MG	2.5		01/01/2006	99/99/9999							
00703-9402-04	J3260			01/01/2002	12/18/2017	INJECTION, TOBRAMYcIN SULFATE, UP TO 80 MG	TOBRAMYcIN SULFATE (M.D.V.) 40 MG/ML	2 ML	VL	IJ	ML	80 MG	0.5		01/01/2002	12/18/2017							
00703-4094-01	J2469			03/23/2018	99/99/9999	INJECTION, PALONOsETRON HCL, 25 MCG	PALONOsETRON HCL (S.D.V.) 0.05 MG/1 ML	5 ML	VL	IV	ML	25 MCG	2		03/23/2018	99/99/9999							
00781-3312-75	J2469			03/23/2018	99/99/9999	INJECTION, PALONOsETRON HCL, 25 MCG	PALONOsETRON HCL 0.05 MG/1 ML	5 ML	VL	IV	ML	25 MCG	2		03/23/2018	99/99/9999							
16729-0259-38	J1327			02/01/2018	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE 0.75 MG/1 ML	100 ML	VL	IV	ML	5 MG	0.15		02/01/2018	99/99/9999							
00703-9416-01	J3260			01/01/2002	06/25/2018	INJECTION, TOBRAMYcIN SULFATE, UP TO 80 MG	TOBRAMYcIN SULFATE (M.D.V.) 40 MG/ML	30 ML	VL	IJ	ML	80 MG	0.5		01/01/2002	06/25/2018							
00703-9503-03	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	SMZ-TMP CONCENTRATE (S.D.V.) 80 MG/ML-16 MG/ML	5 ML	VL	IV	ML	1 EA	1		01/01/2002	99/99/9999							
00703-9514-03	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	SMZ-TMP CONCENTRATE (M.D.V.) 80 MG/ML-16 MG/ML	10 ML	VL	IV	ML	1 EA	1		01/01/2002	99/99/9999							
00703-9526-01	J3490			01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	SMZ-TMP (M.D.V.) 80 MG/ML-16 MG/ML	30 ML	VL	IV	ML	1 EA	1		01/01/2002	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00713-0135-12		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
00713-0526-12		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHEGAN 25 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
00713-0536-12		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHEGAN 12.5 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
00761-0914-20		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ANTI-HIST 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
00781-1046-01		Q0175		01/01/2002	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 2 MG	100 EA	BO	PO	EA		4 MG		0.5	01/01/2002	99/99/9999						
00781-1046-10		Q0175		01/01/2002	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 2 MG	1000 EA	BO	PO	EA		4 MG		0.5	05/16/2008	99/99/9999	01/01/2002	12/01/2004	0.5			
00781-1046-13		Q0175		01/01/2002	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 2 MG	100 EA	BX	PO	EA		4 MG		0.5	01/01/2002	99/99/9999						
00781-1047-01		Q0175		01/01/2002	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	100 EA	BO	PO	EA		4 MG		1	01/01/2002	99/99/9999						
00781-1047-13		Q0175		01/01/2002	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	100 EA	BX	PO	EA		4 MG		1	01/01/2002	99/99/9999						
00781-1048-01		Q0176		01/01/2002	12/31/2013	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	100 EA	BO	PO	EA		8 MG		1	01/01/2002	12/31/2013						
00781-1048-13		Q0176		01/01/2002	12/31/2013	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	100 EA	BX	PO	EA		8 MG		1	01/01/2002	12/31/2013						
00781-1049-01		Q0176		01/01/2002	12/31/2013	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 16 MG	100 EA	BO	PO	EA		8 MG		2	01/01/2002	12/31/2013						
00781-1496-31		Q0144		01/09/2006	05/15/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	30 EA	BO	PO	EA		1 GM		0.25	01/09/2006	05/15/2017						
00781-1496-68		Q0144		11/14/2005	09/07/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (3X6,UNIT OF USE) 250 MG	3 EA	DP	PO	EA		1 GM		0.25	11/14/2005	09/07/2017						
00781-1496-69		Q0144		11/14/2005	06/13/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	50 EA	BX	PO	EA		1 GM		0.25	11/14/2005	06/13/2017						
00781-1497-31		Q0144		11/14/2005	10/29/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 600 MG	30 EA	BO	PO	EA		1 GM		0.6	11/14/2005	10/29/2017						
00781-1830-01		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100 EA	BO	PO	EA		25 MG		1	01/01/2002	12/31/2013						
00781-1830-10		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	1000 EA	BO	PO	EA		25 MG		1	01/20/2005	12/31/2013	01/01/2002	08/25/2003	1			
00781-1832-01		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	100 EA	BO	PO	EA		25 MG		2	01/01/2002	12/31/2013						
00781-1941-31		Q0144		11/16/2005	09/25/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	30 EA	BO	PO	EA		1 GM		0.5	11/16/2005	09/25/2017						
00781-1941-33		Q0144		11/16/2005	09/07/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (3X3,UNIT OF USE) 500 MG	3 EA	DP	PO	EA		1 GM		0.5	11/16/2005	09/07/2017						
00781-3001-07		J2941		03/12/2008	99/99/9999	INJECTION, SOMATROPIN, 1 MG	OMNITROPE (1X1.5ML/W/DILUENT) 5 MG/1.5 ML	1.5 ML	CT	SC	ML		1 MG		3.33333	03/12/2008	99/99/9999						
00781-3001-26		J2941		03/12/2008	99/99/9999	INJECTION, SOMATROPIN, 1 MG	OMNITROPE (5X1.5ML/W/DILUENT) 5 MG/1.5 ML	1.5 ML	CT	SC	ML		1 MG		3.33333	03/12/2008	99/99/9999						
00781-3009-95		J0330		04/15/2005	09/28/2015	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	ANECTINE (MDV,10MLX10VIALS) 20 MG/ML	10 ML	VL	IV	ML		20 MG		1	04/15/2005	09/28/2015						
00781-3032-95		J0295		09/05/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP) 1 GM-0.5 GM	1 EA	VL	IJ	EA		1.5 GM		1	09/05/2006	99/99/9999						
00781-3033-95		J0295		09/05/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP) 2 GM-1 GM	1 EA	VL	IJ	EA		1.5 GM		2	09/05/2006	99/99/9999						
00781-3034-46		J0295		09/05/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP) 10 GM-5 GM	1 EA	VL	IV	EA		1.5 GM		10	09/05/2006	99/99/9999						
00781-3059-95		J1160		07/21/2006	99/99/9999	INJECTION, DIGOXIN, UP TO 0.5 MG	DIGOXIN (USP,10X2ML) 0.25 MG/ML	2 ML	AM	IJ	ML		0.5 MG		0.5	07/21/2006	99/99/9999						
00781-3073-70		J1070		10/17/2006	11/30/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (USP,MDV) 100 MG/ML	10 ML	VL	IM	ML		100 MG		1	10/17/2006	11/30/2014						
00781-3074-70		J1080		10/17/2006	05/30/2013	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (USP,MDV) 200 MG/ML	10 ML	VL	IM	ML		200 MG		1	10/17/2006	05/30/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00781-3074-71		J1080		10/17/2006	05/30/2013	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (USP,MDV) 200 MG/ML	1 ML	VL	IM	ML		200 MG		1	10/17/2006	05/30/2013						
00781-3084-75		J3303		01/29/2007	08/29/2013	INJECTION, TRIAMCINOLONE HEXACETONIDE, PER 5MG	ARISTOSPAN 5 MG/ML	5 ML	VL	IJ	ML		5 MG		1	01/29/2007	08/29/2013						
00781-3094-15		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN (USP,ADD-VANTAGE VIAL) 1 GM	1 EA	VL	IV	EA		250 MG		4	03/19/2008	99/99/9999						
00781-3094-92		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN (1X10,USP,ADD-VANTAGE) 1 GM	1 EA	VL	IV	EA		250 MG		4	03/19/2008	99/99/9999						
00781-3095-80		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN (USP,ADD-VANTAGE VIAL) 2 GM	1 EA	VL	IV	EA		250 MG		8	03/19/2008	99/99/9999						
00781-3095-92		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN (1X10,USP,ADD-VANTAGE) 2 GM	1 EA	VL	IV	EA		250 MG		8	03/19/2008	99/99/9999						
00781-3099-95		J2700		02/08/2005	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN SODIUM 1 GM	1 EA	VL	IJ	EA		250 MG		4	02/08/2005	99/99/9999						
00781-3101-80		J2700		02/01/2007	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN (USP) 2 GM	1 EA	VL	IJ	EA		250 MG		8	02/01/2007	99/99/9999						
00781-3101-95		J2700		07/02/2004	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN SODIUM (VIAL,PIGGYBACK) 2 GM	1 EA	VL	IJ	EA		250 MG		8	07/02/2004	99/99/9999						
00781-3103-95		J2700		08/31/2004	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN SODIUM (PHARMACY BULK PACKAGE) 10 GM	1 EA	VL	IJ	EA		250 MG		40	08/31/2004	99/99/9999						
00781-3124-85		J3490		09/09/2005	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM 1 GM	1 EA	VL	IJ	EA		1 EA		1	09/09/2005	99/99/9999						
00781-3124-95		J3490		04/27/2004	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM (VIAL) 1 GM	1 EA	VL	IJ	EA		1 EA		1	04/27/2004	99/99/9999						
00781-3125-85		J3490		09/09/2005	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM 2 GM	1 EA	VL	IJ	EA		1 EA		1	09/09/2005	99/99/9999						
00781-3125-92		J3490		02/23/2005	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM (ADD-VANTAGE VIAL) 2 GM	1 EA	VL	IJ	EA		1 EA		1	02/23/2005	99/99/9999						
00781-3125-95		J3490		04/27/2004	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM (VIAL) 2 GM	1 EA	VL	IJ	EA		1 EA		1	04/27/2004	99/99/9999						
00781-3126-46		J3490		09/09/2005	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM 10 GM	1 EA	VL	IJ	EA		1 EA		1	09/09/2005	99/99/9999						
00781-3126-95		J3490		04/27/2004	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM (VIAL,PHARMACY BULK) 10 GM	1 EA	VL	IJ	EA		1 EA		1	04/27/2004	99/99/9999						
00781-3128-92		J3490		04/17/2006	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN (USP,ADD-VANTAGE VIAL) 1 GM	1 EA	VL	IV	EA		1 EA		1	04/17/2006	99/99/9999						
00781-3129-92		J3490		02/22/2006	99/99/9999	UNCLASSIFIED DRUGS	NAFCILLIN SODIUM (2GMX10, ADD-VANTAGE) 2 GM	1 EA	VL	IV	EA		1 EA		1	02/22/2006	99/99/9999						
00781-3177-96		J0713		02/23/2007	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	CEFTAZIDIME (USP) 1 GM	1 EA	VL	IJ	EA		500 MG		2	02/23/2007	99/99/9999						
00781-3178-95		J0713		02/23/2007	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	CEFTAZIDIME (USP) 2 GM	1 EA	VL	IV	EA		500 MG		4	02/23/2007	99/99/9999						
00781-3179-86		J0713		02/23/2007	99/99/9999	INJECTION, CEFTAZIDIME, PER 500 MG	CEFTAZIDIME (USP,PHARMACY BULK PKG) 6 GM	1 EA	VL	IV	EA		500 MG		12	02/23/2007	99/99/9999						
00781-3182-73		J1451		04/02/2008	99/99/9999	INJECTION, FOMEPIZOLE, 15 MG	FOMEPIZOLE (1X1.5ML,PF) 1 GM/ML	1.5 ML	VL	IV	ML		15 MG	66.66666		04/02/2008	99/99/9999						
00781-3182-84		J1451		04/02/2008	99/99/9999	INJECTION, FOMEPIZOLE, 15 MG	FOMEPIZOLE (4X1.5ML,PF) 1 GM/ML	1.5 ML	VL	IV	ML		15 MG	66.66666		04/02/2008	99/99/9999						
00781-3206-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 250 MG	1 EA	VL	IJ	EA		250 MG		1	07/19/2005	99/99/9999						
00781-3207-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 500 MG	1 EA	VL	IJ	EA		250 MG		2	07/19/2005	99/99/9999						
00781-3208-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/19/2005	99/99/9999						
00781-3209-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 2 GM	1 EA	VL	IJ	EA		250 MG		8	07/19/2005	99/99/9999						
00781-3210-46		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 10 GM	1 EA	VL	IJ	EA		250 MG		40	07/19/2005	99/99/9999						
00781-3222-80		J0692		04/14/2008	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME HYDROCHLORIDE (S.D.V,USP) 1 GM	1 EA	VL	IJ	EA		500 MG		2	04/14/2008	99/99/9999						
00781-3222-95		J0692		04/14/2008	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME HYDROCHLORIDE (USP) 1 GM	1 EA	VL	IJ	EA		500 MG		2	04/14/2008	99/99/9999						
00781-3223-91		J0692		04/14/2008	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME HYDROCHLORIDE (S.D.V,USP) 2 GM	1 EA	VL	IJ	EA		500 MG		4	04/14/2008	99/99/9999						
00781-3223-95		J0692		04/14/2008	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME HYDROCHLORIDE (USP) 2 GM	1 EA	VL	IJ	EA		500 MG		4	04/14/2008	99/99/9999						
00781-3239-09		J0744		03/18/2008	99/99/9999	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	CIPROFLOXACIN (24X100ML,USP,LATEX-FREE) 200 MG/100 ML	100 ML	FC	IV	ML		200 MG		0.01	03/18/2008	99/99/9999						
00781-3240-09		J0744		03/18/2008	99/99/9999	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	CIPROFLOXACIN (24X200ML,USP,LATEX-FREE) 400 MG/200 ML	200 ML	FC	IV	ML		200 MG		0.01	03/18/2008	99/99/9999						
00781-3338-70		J0690		08/23/2004	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (1X10ML VIAL) 500 MG	1 EA	VL	IJ	EA		500 MG		1	08/23/2004	99/99/9999						
00781-3400-95		J0290		05/12/2004	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM 125 MG	1 EA	VL	IJ	EA		500 MG		0.25	05/12/2004	99/99/9999						
00781-3402-95		J0290		12/01/2005	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (U.S.P.) 250 MG	1 EA	VL	IJ	EA		500 MG		0.5	12/01/2005	99/99/9999						
00781-3404-95		J0290		12/01/2005	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (U.S.P.) 1 GM	1 EA	VL	IJ	EA		500 MG		2	12/01/2005	99/99/9999						
00781-3407-95		J0290		12/01/2005	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (U.S.P.) 500 MG	1 EA	VL	IJ	EA		500 MG		1	12/01/2005	99/99/9999						
00781-3408-95		J0290		12/01/2005	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (U.S.P.) 2 GM	1 EA	VL	IJ	EA		500 MG		4	12/01/2005	99/99/9999						
00781-3409-95		J0290		05/12/2004	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM 10 GM	1 EA	VL	IJ	EA		500 MG		20	05/12/2004	99/99/9999						
00781-3412-92		J0290		03/20/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (ADD-VANTAGE,USP) 1 GM	1 EA	VL	IJ	EA		500 MG		2	03/20/2007	99/99/9999						
00781-3413-92		J0290		03/20/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (ADD-VANTAGE,ADD-VANTAGE) 2 GM	1 EA	VL	IJ	EA		500 MG		4	03/20/2007	99/99/9999						
00781-3450-95		J0690		11/08/2006	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (USP) 500 MG	1 EA	VL	IJ	EA		500 MG		1	11/08/2006	99/99/9999						
00781-3451-96		J0690		09/13/2006	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN (USP) 1 GM	1 EA	VL	IJ	EA		500 MG		2	09/13/2006	99/99/9999						
00781-3452-95		J0690		09/13/2006	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN (USP) 10 GM	1 EA	VL	IV	EA		500 MG		20	09/13/2006	99/99/9999						
00781-3777-95		J1800		02/15/2007	11/30/2013	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HYDROCHLORIDE (USP,10X1ML) 1 MG/ML	1 ML	VL	IV	ML		1 MG		1	02/15/2007	11/30/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00781-4004-36		J2941		01/15/2007	99/99/9999	INJECTION, SOMATROPIN, 1 MG	OMNITROPE (W/ 8 VIALS OF DILUENT) 5.8 MG	1 EA	VL	SC	EA		1 MG		5.8	01/15/2007	99/99/9999						
00781-5020-01		Q0164		01/01/2002	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	100 EA	BO	PO	EA		5 MG		1	01/01/2002	99/99/9999						
00781-5021-01		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100 EA	BO	PO	EA		10 MG		1	01/01/2002	12/31/2013						
00781-5022-01		J7509		04/04/2003	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	100 EA	BO	PO	EA		4 MG		1	04/04/2003	99/99/9999						
00781-5022-07		J7509		04/04/2003	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (UNIT OF USE) 4 MG	21 EA	DP	PO	EA		4 MG		1	04/04/2003	99/99/9999						
00781-6135-95		J2540		11/25/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PENICILLIN G POTASSIUM 5 Million U	1 EA	VL	IV	EA		600000 U		8.33333	11/25/2002	99/99/9999						
55150-0266-05		J3489		09/27/2018	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SINGLE-USE,LATEX-FREE) 4 MG/5 ML	5 ML	VL	IV	ML		1 MG		0.8	09/27/2018	99/99/9999						
00781-6136-94		J2540		11/25/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PENICILLIN G POTASSIUM 20 Million U	1 EA	VL	IV	EA		600000 U		33.33333	11/25/2002	99/99/9999						
00781-6153-95		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	PENICILLIN G SODIUM (VIAL) 5 Million U	1 EA	VL	IV	EA		1 EA		1	01/01/2002	99/99/9999						
00781-9109-85		J2700		02/01/2007	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN 1 GM	1 EA	VL	IJ	EA		250 MG		4	02/01/2007	99/99/9999						
00781-9109-95		J2700		03/01/2006	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN (USP) 1 GM	1 EA	VL	IJ	EA		250 MG		4	03/01/2006	99/99/9999						
00781-9110-15		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN (USP,ADD-VANTAGE VIAL) 1 GM	1 EA	VL	IV	EA		250 MG		4	03/19/2008	99/99/9999						
00781-9110-92		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN (1X10,USP,ADD-VANTAGE) 1 GM	1 EA	VL	IV	EA		250 MG		4	03/19/2008	99/99/9999						
00781-9111-80		J2700		02/01/2007	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN 2 GM	1 EA	VL	IJ	EA		250 MG		8	02/01/2007	99/99/9999						
00781-9111-95		J2700		05/04/2006	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN (USP) 2 GM	1 EA	VL	IJ	EA		250 MG		8	05/04/2006	99/99/9999						
00781-9112-20		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN (USP,ADD-VANTAGE VIAL) 2 GM	1 EA	VL	IV	EA		250 MG		8	03/19/2008	99/99/9999						
00781-9112-92		J2700		03/19/2008	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN (1X10,USP,ADD-VANTAGE) 2 GM	1 EA	VL	IV	EA		250 MG		8	03/19/2008	99/99/9999						
00781-9113-46		J2700		02/01/2007	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN 10 GM	1 EA	VL	IJ	EA		250 MG		40	02/01/2007	99/99/9999						
00781-9113-95		J2700		05/03/2006	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	NOVAPLUS OXACILLIN 10 GM	1 EA	VL	IJ	EA		250 MG		40	05/03/2006	99/99/9999						
00781-9124-85		J3490		02/01/2007	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN 1 GM	1 EA	VL	IJ	EA		1 EA		1	02/01/2007	99/99/9999						
00781-9124-95		J3490		02/01/2006	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN 1 GM	1 EA	VL	IJ	EA		1 EA		1	02/01/2006	99/99/9999						
00781-9125-85		J3490		02/01/2007	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN 2 GM	1 EA	VL	IJ	EA		1 EA		1	02/01/2007	99/99/9999						
00781-9125-95		J3490		02/01/2006	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN 2 GM	1 EA	VL	IJ	EA		1 EA		1	02/01/2006	99/99/9999						
00781-9126-46		J3490		03/31/2007	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN 10 GM	1 EA	VL	IJ	EA		1 EA		1	03/31/2007	99/99/9999						
00781-9126-95		J3490		02/01/2006	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN (BULK PACKAGE) 10 GM	1 EA	VL	IJ	EA		1 EA		1	02/01/2006	99/99/9999						
00781-9164-75		J2354		04/07/2005	03/28/2013	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE NOVAPLUS (M.D.V.) 1000 MCG/ML	5 ML	VL	IJ	ML		25 MCG		40	04/07/2005	03/28/2013						
00781-9165-75		J2354		04/07/2005	03/28/2013	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE NOVAPLUS (M.D.V.) 200 MCG/ML	5 ML	VL	IJ	ML		25 MCG		8	04/07/2005	03/28/2013						
00781-9166-95		J2354		04/07/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE NOVAPLUS (M.D.V.) 50 MCG/ML	1 ML	AM	IJ	ML		25 MCG		2	04/07/2005	99/99/9999						
00781-9167-95		J2354		04/07/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE NOVAPLUS (M.D.V.) 100 MCG/ML	1 ML	AM	IJ	ML		25 MCG		4	04/07/2005	99/99/9999						
00781-9168-95		J2354		04/07/2005	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE NOVAPLUS (M.D.V.) 500 MCG/ML	1 ML	AM	IJ	ML		25 MCG		20	04/07/2005	99/99/9999						
00781-9224-15		J3490		02/01/2007	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN (ADD-VANTAGE) 1 GM	1 EA	VL	IV	EA		1 EA		1	02/01/2007	99/99/9999						
00781-9224-92		J3490		09/18/2006	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN (USP,ADD-VANTAGE) 1 GM	1 EA	VL	IV	EA		1 EA		1	09/18/2006	99/99/9999						
00781-9225-20		J3490		02/01/2007	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN (ADD-VANTAGE) 2 GM	1 EA	VL	IV	EA		1 EA		1	02/01/2007	99/99/9999						
00781-9225-92		J3490		09/18/2006	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS NAFACILLIN (USP,ADD-VANTAGE) 2 GM	1 EA	VL	IV	EA		1 EA		1	09/18/2006	99/99/9999						
00781-9326-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS 250 MG	1 EA	VL	IJ	EA		250 MG		1	07/19/2005	99/99/9999						
00781-9327-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS 500 MG	1 EA	VL	IJ	EA		250 MG		2	07/19/2005	99/99/9999						
00781-9328-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/19/2005	99/99/9999						
00781-9329-90		J0696		03/31/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS 2 GM	1 EA	VL	IJ	EA		250 MG		8	03/31/2007	99/99/9999						
00781-9329-95		J0696		07/19/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS 2 GM	1 EA	VL	IJ	EA		250 MG		8	07/19/2005	99/99/9999						
00781-9330-46		J0696		07/19/2005	06/30/2015	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS 10 GM	1 EA	VL	IJ	EA		250 MG		40	07/19/2005	06/30/2015						
00069-1307-10		Q5106		01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 4000 U/1 ML	1 ML	VL	IJ	ML		1000 U		4	01/01/2019	99/99/9999						
00069-1308-10		Q5106		01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 10000 U/1 ML	1 ML	VL	IJ	ML		1000 U		10	01/01/2019	99/99/9999						
00781-9338-85		J0690		02/27/2006	99/99/9999	INJECTION, CEFZAZOLIN SODIUM, 500 MG	NOVAPLUS CEFZAZOLIN 500 MG	1 EA	VL	IJ	EA		500 MG		1	02/27/2006	99/99/9999						
00781-9338-95		J0690		02/27/2006	99/99/9999	INJECTION, CEFZAZOLIN SODIUM, 500 MG	NOVAPLUS CEFZAZOLIN (USP) 500 MG	1 EA	VL	IJ	EA		500 MG		1	02/27/2006	99/99/9999						
00781-9339-85		J0690		05/15/2007	05/02/2012	INJECTION, CEFZAZOLIN SODIUM, 500 MG	NOVAPLUS CEFZAZOLIN 1 GM	1 EA	VL	IJ	EA		500 MG		2	05/15/2007	05/02/2012						
00781-9401-78		J0290		02/01/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN 125 MG	1 EA	VL	IJ	EA		500 MG		0.25	02/01/2007	99/99/9999						
00781-9401-95		J0290		02/01/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (USP) 125 MG	1 EA	VL	IJ	EA		500 MG		0.25	02/01/2006	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00069-1309-04		Q5106		01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 40000 U/1 ML	1	ML	VL	IJ	ML	1000 U		40	01/01/2019	99/99/9999						
00781-9402-78	J0290			01/24/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN 250 MG	1	EA	VL	IJ	EA	500 MG		0.5	01/24/2006	99/99/9999						
00781-9402-95	J0290			02/01/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (USP) 250 MG	1	EA	VL	IJ	EA	500 MG		0.5	02/01/2006	99/99/9999						
00781-9404-85	J0290			01/24/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN 1 GM	1	EA	VL	IJ	EA	500 MG		2	01/24/2006	99/99/9999						
00781-9404-95	J0290			02/01/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (USP) 1 GM	1	EA	VL	IJ	EA	500 MG		2	02/01/2006	99/99/9999						
00781-9407-78	J0290			01/24/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN 500 MG	1	EA	VL	IJ	EA	500 MG		1	01/24/2006	99/99/9999						
00781-9407-95	J0290			02/01/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (USP) 500 MG	1	EA	VL	IJ	EA	500 MG		1	02/01/2006	99/99/9999						
00781-9408-80	J0290			01/24/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN 2 GM	1	EA	VL	IJ	EA	500 MG		4	01/24/2006	99/99/9999						
00781-9408-95	J0290			02/01/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (ADD-VANTAGE) 2 GM	1	EA	VL	IJ	EA	500 MG		4	02/01/2007	99/99/9999						
00781-9408-95	J0290			02/01/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (USP) 2 GM	1	EA	VL	IJ	EA	500 MG		4	02/01/2006	99/99/9999						
00781-9409-95	J0290			02/01/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (USP) 10 GM	1	EA	VL	IJ	EA	500 MG		20	02/01/2006	99/99/9999						
00781-9412-15	J0290			02/01/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (ADD-VANTAGE) 1 GM	1	EA	VL	IJ	EA	500 MG		2	02/01/2007	99/99/9999						
00781-9412-92	J0290			03/20/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (ADD-VANTAGE) 1 GM	1	EA	VL	IJ	EA	500 MG		2	03/20/2007	99/99/9999						
00781-9413-92	J0290			03/20/2007	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	NOVAPLUS AMPICILLIN (ADD-VANTAGE) 2 GM	1	EA	VL	IJ	EA	500 MG		4	03/20/2007	99/99/9999						
00781-9452-95	J0690			01/09/2007	02/09/2013	INJECTION, CEFAZOLIN SODIUM, 500 MG	NOVAPLUS CEFAZOLIN (USP) 10 GM	1	EA	VL	IV	EA	500 MG		20	01/09/2007	02/09/2013						
00904-1228-00	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	BANOPHEN (AF) 12.5 MG/5 ML	120	ML	BO	PO	ML	50 MG		0.05	01/01/2002	99/99/9999						
00904-1228-20	Q0163			01/01/2002	07/30/2015	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	BANOPHEN (BOXED) 12.5 MG/5 ML	120	ML	BO	PO	ML	50 MG		0.05	01/01/2002	07/30/2015						
00904-2035-24	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	BANOPHEN 25 MG	24	EA	BX	PO	EA	50 MG		0.5	01/01/2002	99/99/9999						
00904-2035-59	Q0163			01/01/2002	06/28/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	BANOPHEN 25 MG	100	EA	BO	PO	EA	50 MG		0.5	01/01/2002	06/28/2013						
00904-2056-61	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (10X10) 50 MG	100	EA	BX	PO	EA	50 MG		1	01/01/2002	99/99/9999						
00904-3571-61	J8999			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (10X10) 40 MG	100	EA	BX	PO	EA	1 EA		1	01/01/2002	99/99/9999						
00904-4274-51	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SLEEP TABS 25 MG	50	EA	BO	PO	EA	50 MG		0.5	01/01/2002	99/99/9999						
00904-5174-16	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	BANOPHEN 12.5 MG/5 ML	480	ML	BO	PO	ML	50 MG		0.05	01/01/2002	99/99/9999						
00904-5306-60	Q0163			01/01/2002	08/09/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100	EA	BO	PO	EA	50 MG		0.5	01/01/2002	08/09/2012						
00904-5306-61	Q0163			05/12/2003	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (10X10) 25 MG	100	EA	BX	PO	EA	50 MG		0.5	05/12/2003	99/99/9999						
00904-5306-80	Q0163			01/01/2002	08/09/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	1000	EA	BO	PO	EA	50 MG		0.5	01/01/2002	08/09/2012						
00904-5307-60	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50 MG		1	01/01/2002	99/99/9999						
00904-5307-80	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	1000	EA	BO	PO	EA	50 MG		1	01/01/2002	99/99/9999						
00904-5551-59	Q0163			08/13/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	BANOPHEN (MINI TABS,MINI TAB) 25 MG	100	EA	BX	PO	EA	50 MG		0.5	08/13/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00904-5840-61		Q0170		05/06/2008	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	100	EA	BX	PO	EA	25	MG	1	05/06/2008	12/31/2013						
00927-0221-24		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALLERMAX 50 MG	24	EA	BX	PO	EA	50	MG	1	01/01/2002	02/03/2016						
00927-0616-34		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TWILITE 50 MG ZOMACTON WITH VIAL ADAPTER (LYOPHILIZED) 10 MG	20	EA	BX	PO	EA	50	MG	1	01/01/2002	99/99/9999						
55566-1902-01		J2941		09/26/2018	99/99/9999	INJECTION, SOMATROPIN, 1 MG		1	EA	VL	SC	EA	1	MG	10	09/26/2018	99/99/9999						
00927-0617-12		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALLERMAX 12.5 MG/5 ML	120	ML	BO	PO	ML	50	MG	0.05	01/01/2002	99/99/9999						
00944-2620-02		J1566		01/01/2006	05/25/2013	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	GAMMAGARD S/D 2.5 GM	1	EA	VL	IV	EA	500	MG	5	01/01/2006	05/25/2013						
00944-2620-03		J1566		01/01/2006	04/11/2014	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	GAMMAGARD S/D 5 GM	1	EA	VL	IV	EA	500	MG	10	01/01/2006	04/11/2014						
00944-2620-04		J1566		01/01/2006	06/21/2014	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	GAMMAGARD S/D 10 GM	1	EA	VL	IV	EA	500	MG	20	01/01/2006	06/21/2014						
00944-2655-03		J1566		06/01/2007	01/03/2015	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	GAMMAGARD S/D (W/TRANSFER SET) 5 GM	1	EA	VL	IV	EA	500	MG	10	06/01/2007	01/03/2015						
00944-2655-04		J1566		06/01/2007	01/03/2015	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	GAMMAGARD S/D (W/TRANSFER SET) 10 GM	1	EA	VL	IV	EA	500	MG	20	06/01/2007	01/03/2015						
00944-2700-02		J1569		01/01/2008	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAGARD LIQUID), NON-LYOPHILIZED,(E.G. LIQUID), 500 MG	GAMMAGARD LIQUID (PF.LATEX-FREE) 100 MG/ML	10	ML	VL	IV	ML	500	MG	0.2	01/01/2008	99/99/9999						
00944-2700-03		J1569		01/01/2008	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAGARD LIQUID), NON-LYOPHILIZED,(E.G. LIQUID), 500 MG	GAMMAGARD LIQUID (PF.LATEX-FREE) 100 MG/ML	25	ML	VL	IV	ML	500	MG	0.2	01/01/2008	99/99/9999						
00944-2700-04		J1569		01/01/2008	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAGARD LIQUID), NON-LYOPHILIZED,(E.G. LIQUID), 500 MG	GAMMAGARD LIQUID (PF.LATEX-FREE) 100 MG/ML	50	ML	VL	IV	ML	500	MG	0.2	01/01/2008	99/99/9999						
00944-2700-05		J1569		01/01/2008	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAGARD LIQUID), NON-LYOPHILIZED,(E.G. LIQUID), 500 MG	GAMMAGARD LIQUID (PF.LATEX-FREE) 100 MG/ML	100	ML	VL	IV	ML	500	MG	0.2	01/01/2008	99/99/9999						
00944-2700-06		J1569		01/01/2008	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAGARD LIQUID), NON-LYOPHILIZED,(E.G. LIQUID), 500 MG	GAMMAGARD LIQUID (PF.LATEX-FREE) 100 MG/ML	200	ML	VL	IV	ML	500	MG	0.2	01/01/2008	99/99/9999						
00944-2967-03		J2792		03/01/2006	09/30/2012	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT DETERGENT, 100 IU	WINRHO SDF (SDV,PF) 1500 IU	1.3	ML	VL	IV	ML	100	IU	11.53846	03/01/2006	09/30/2012						
00944-2967-05		J2792		03/01/2006	09/30/2012	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT DETERGENT, 100 IU	WINRHO SDF (SDV,PF) 5000 IU	4.4	ML	VL	IV	ML	100	IU	11.36363	03/01/2006	09/30/2012						
00944-2967-07		J2792		03/01/2006	10/31/2012	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT DETERGENT, 100 IU	WINRHO SDF (SDV,PF) 2500 IU	2.2	ML	VL	IV	ML	100	IU	11.36363	03/01/2006	10/31/2012						
00944-2967-09		J2792		03/01/2006	10/31/2012	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT DETERGENT, 100 IU	WINRHO SDF (SDV,PF) 15000 IU	13	ML	VL	IV	ML	100	IU	11.53846	03/01/2006	10/31/2012						
00944-4175-05		J2724		01/01/2008	06/30/2015	INJECTION, PROTEIN C CONCENTRATE, INTRAVENOUS, HUMAN, 10 IU	CEPROTIN (400-600IU) 1 IU	600	IU	VL	IV	EA	10	IU	0.1	01/01/2008	06/30/2015						
00944-4175-10		J2724		01/01/2008	06/30/2015	INJECTION, PROTEIN C CONCENTRATE, INTRAVENOUS, HUMAN, 10 IU	CEPROTIN (800-1200IU) 1 IU	1200	IU	VL	IV	EA	10	IU	0.1	01/01/2008	06/30/2015						
03221-0208-11		J3490		01/01/2008	99/99/9999	UNCLASSIFIED DRUGS	VERITAS COLLAGEN MATRIX (2CMX8CM)	1	EA	NA	IP	EA	1	EA	1	01/01/2008	99/99/9999						
03221-0407-11		J3490		01/01/2008	99/99/9999	UNCLASSIFIED DRUGS	VERITAS COLLAGEN MATRIX (4CMX7CM)	1	EA	NA	IP	EA	1	EA	1	01/01/2008	99/99/9999						
03221-0415-11		J3490		01/01/2008	99/99/9999	UNCLASSIFIED DRUGS	VERITAS COLLAGEN MATRIX (4CMX15CM)	1	EA	NA	IP	EA	1	EA	1	01/01/2008	99/99/9999						
03221-0608-11		J3490		01/01/2008	99/99/9999	UNCLASSIFIED DRUGS	VERITAS COLLAGEN MATRIX (6CMX8CM)	1	EA	NA	IP	EA	1	EA	1	01/01/2008	99/99/9999						
03221-0814-11		J3490		01/01/2008	99/99/9999	UNCLASSIFIED DRUGS	VERITAS COLLAGEN MATRIX (8CMX14CM)	1	EA	NA	IP	EA	1	EA	1	01/01/2008	99/99/9999						
03221-1016-11		J3490		01/01/2008	99/99/9999	UNCLASSIFIED DRUGS	VERITAS COLLAGEN MATRIX (10CMX16CM)	1	EA	NA	IP	EA	1	EA	1	01/01/2008	99/99/9999						
03221-1225-11		J3490		01/01/2008	99/99/9999	UNCLASSIFIED DRUGS	VERITAS COLLAGEN MATRIX (12CMX25CM)	1	EA	NA	IP	EA	1	EA	1	01/01/2008	99/99/9999						
08080-1000-00		A4217		03/01/2006	99/99/9999	STERILE WATER/SALINE, 500 ML	CURITY STERILE WATER	100	ML	NA	IR	ML	500	ML	0.002	03/01/2006	99/99/9999						
08080-1020-00		A4217		03/01/2006	99/99/9999	STERILE WATER/SALINE, 500 ML	CURITY STERILE SALINE (100MLX48) 0.9%	100	ML	NA	IR	ML	500	ML	0.002	03/01/2006	99/99/9999						
08080-1022-00		A4217		03/01/2006	99/99/9999	STERILE WATER/SALINE, 500 ML	CURITY STERILE SALINE (100MLX48) 0.9%	100	ML	NA	IR	ML	500	ML	0.002	03/01/2006	99/99/9999						
08166-1100-03		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	VASCEZE HEPARIN LOCK FLUSH (LUER SLIP NOZZLE) 100 U/ML	3	ML	NA	IV	ML	10	U	10	01/01/2002	99/99/9999						
08166-1100-05		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	VASCEZE HEPARIN LOCK FLUSH (LUER SLIP NOZZLE) 100 U/ML	5	ML	NA	IV	ML	10	U	10	01/01/2002	02/03/2016						
08166-1109-03		A4216		01/01/2007	09/19/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	VASCEZE SODIUM CHLORIDE (LUER SLIP NOZZLE) 0.9%	3	ML	NA	IV	ML	10	ML	0.1	01/01/2007	09/19/2016						
08166-1109-05		A4216		01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	VASCEZE SODIUM CHLORIDE (LUER SLIP NOZZLE) 0.9%	5	ML	NA	IV	ML	10	ML	0.1	01/01/2007	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
08166-1109-10		A4216		01/01/2004	09/19/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	VASCEZE SODIUM CHLORIDE (LUER SLIP NOZZLE) 0.9%	10	ML	NA	IV	ML	10	ML	0.1	01/01/2004	09/19/2016							
08166-1110-03		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	VASCEZE HEPARIN LOCK FLUSH (LUER SLIP NOZZLE.PF) 10 U/ML	3	ML	NA	IV	ML	10	U	1	01/01/2002	99/99/9999							
08166-1110-05		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	VASCEZE HEPARIN LOCK FLUSH (LUER SLIP NOZZLE.PF) 10 U/ML	5	ML	NA	IV	ML	10	U	1	01/01/2002	02/03/2016							
08290-0310-02		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,3 ML,PF) 0.9%	2	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0310-03		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,3 ML,PF) 0.9%	3	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0311-03		A4216		01/01/2004	10/17/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,3 ML W/CANNULA,PF) 0.9%	3	ML	SR	IV	ML	10	ML	0.1	01/01/2004	10/17/2016							
08290-0320-03		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,6 ML,PF) 0.9%	3	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0320-05		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,6 ML,PF) 0.9%	5	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0321-05		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,6 ML W/CANNULA,PF) 0.9%	5	ML	SR	IV	ML	10	ML	0.1	01/01/2004	99/99/9999							
08290-0330-03		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,12 ML,PF) 0.9%	3	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0330-05		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,12 ML,PF) 0.9%	5	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0330-10		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,12 ML,PF) 0.9%	10	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0331-05		A4216		01/01/2004	10/17/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN, 12 ML W/ CANN,PF) 0.9%	5	ML	SR	IV	ML	10	ML	0.1	01/01/2004	10/17/2016							
08290-0331-10		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN, 12 ML W/CANN,PF) 0.9%	10	ML	SR	IV	ML	10	ML	0.1	01/01/2004	99/99/9999							
08290-0910-02		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,2ML,PF) 0.9%	2	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08290-0911-02		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN, W/CANNULA,PF) 0.9%	2	ML	SR	IV	ML	10	ML	0.1	01/01/2004	99/99/9999							
08290-0930-10		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN, 10ML,PF) 0.9%	10	ML	SR	IV	ML	10	ML	0.1	01/01/2007	99/99/9999							
08881-5701-28		A4216		07/01/2006	01/01/2017	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	MONOJECT PREFILL ADVANCED (60X10ML,PF,LATEX-FREE) 0.9%	10	ML	SR	IV	ML	10	ML	0.1	07/01/2006	01/01/2017							
08881-5701-29		A4216		07/01/2006	01/01/2017	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	MONOJECT PREFILL ADVANCED (120X10ML,PF,LATEX-FREE) 0.9%	10	ML	SR	IV	ML	10	ML	0.1	07/01/2006	01/01/2017							
10019-0016-02		J7643		09/28/2005	10/09/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	20	ML	VL	IJ	ML	1	MG	0.2	09/28/2005	10/09/2012							
10019-0016-02	KO	J7643	KO	09/28/2005	10/09/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	20	ML	VL	IJ	ML	1	MG	0.2	09/28/2005	10/09/2012							
10019-0016-17		J7643		01/01/2002	10/09/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	2	ML	VL	IJ	ML	1	MG	0.2	01/01/2002	10/09/2012							
10019-0016-17	KO	J7643	KO	01/01/2002	10/09/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	2	ML	VL	IJ	ML	1	MG	0.2	01/01/2002	10/09/2012							
10019-0016-29		J7643		05/05/2007	04/30/2014	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (MDV) 0.2 MG/ML	20	ML	VL	IJ	ML	1	MG	0.2	05/05/2007	04/30/2014							
10019-0016-29	KO	J7643	KO	05/05/2007	04/30/2014	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (MDV) 0.2 MG/ML	20	ML	VL	IJ	ML	1	MG	0.2	05/05/2007	04/30/2014							
10019-0016-54		J7643		01/01/2002	07/24/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	5	ML	VL	IJ	ML	1	MG	0.2	01/01/2002	07/24/2012							
10019-0016-54	KO	J7643	KO	01/01/2002	07/24/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (M.D.V.) 0.2 MG/ML	5	ML	VL	IJ	ML	1	MG	0.2	01/01/2002	07/24/2012							
10019-0016-81		J7643		01/01/2002	09/06/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	1	ML	VL	IJ	ML	1	MG	0.2	01/01/2002	09/06/2012							
10019-0016-81	KO	J7643	KO	01/01/2002	09/06/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (S.D.V.) 0.2 MG/ML	1	ML	VL	IJ	ML	1	MG	0.2	01/01/2002	09/06/2012							
10019-0027-39		J2250		05/05/2007	10/17/2016	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL 5 MG/ML	10	ML	VL	IJ	ML	1	MG	5	05/05/2007	10/17/2016							
16729-0260-03		J1327		02/01/2018	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE 2 MG/1 ML	10	ML	VL	IV	ML	5	MG	0.4	02/01/2018	99/99/9999							
16729-0260-38		J1327		02/01/2018	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE 2 MG/1 ML	100	ML	VL	IV	ML	5	MG	0.4	02/01/2018	99/99/9999							
10019-0028-37		J2250		05/05/2007	02/03/2016	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL 1 MG/ML	5	ML	VL	IJ	ML	1	MG	1	05/05/2007	02/03/2016							
10019-0028-39		J2250		05/05/2007	02/03/2016	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL 1 MG/ML	10	ML	VL	IJ	ML	1	MG	1	05/05/2007	02/03/2016							
10019-0029-02		J1885		07/21/2004	07/24/2012	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (1X25) 15 MG/ML	1	ML	VL	IJ	ML	15	MG	1	07/21/2004	07/24/2012							
10019-0029-12		J1885		05/05/2007	07/25/2012	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 15 MG/ML	1	ML	VL	IJ	ML	15	MG	1	05/05/2007	07/25/2012							
10019-0030-03		J1885		07/21/2004	07/24/2012	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (1X25) 30 MG/ML	1	ML	VL	IJ	ML	15	MG	2	07/21/2004	07/24/2012							
10019-0030-04		J1885		07/21/2004	07/24/2012	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (1X25) 30 MG/ML	2	ML	VL	IJ	ML	15	MG	2	07/21/2004	07/24/2012							
10019-0030-12		J1885		05/05/2007	10/17/2016	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP) 30 MG/ML	1	ML	VL	IJ	ML	15	MG	2	05/05/2007	10/17/2016							
10019-0030-17		J1885		05/05/2007	10/31/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/ML	2	ML	VL	IJ	ML	15	MG	2	05/05/2007	10/31/2013							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
10019-0033-72		J3010		01/01/2002	11/12/2012	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (AMP,PF) 0.05 MG/ML	5 ML	AM	IJ	ML		0.1 MG		0.5	01/01/2002	11/12/2012						
10019-0035-74		J3010		01/01/2002	10/09/2012	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (AMP,PF) 0.05 MG/ML	20 ML	AM	IJ	ML		0.1 MG		0.5	01/01/2002	10/09/2012						
10019-0037-83		J3010		01/01/2002	07/24/2012	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (S.D.V.,PF) 0.05 MG/ML	50 ML	VL	IJ	ML		0.1 MG		0.5	01/01/2002	07/24/2012						
10019-0038-67		J3010		01/01/2002	10/09/2012	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (AMP,PF) 0.05 MG/ML	2 ML	AM	IJ	ML		0.1 MG		0.5	01/01/2002	10/09/2012						
10019-0045-02		J3490		11/01/2003	05/03/2012	UNCLASSIFIED DRUGS	FAMOTIDINE (S.D.V.,PF) 10 MG/ML	2 ML	VL	IV	ML	1 EA	1		11/01/2003	05/03/2012							
10019-0045-17		J3490		05/05/2007	03/31/2014	UNCLASSIFIED DRUGS	FAMOTIDINE (SDV,PF) 10 MG/ML	2 ML	VL	IV	ML	1 EA	1		05/05/2007	03/31/2014							
10019-0046-03		J3490		01/01/2002	11/12/2012	UNCLASSIFIED DRUGS	FAMOTIDINE (M.D.V.) 10 MG/ML	20 ML	VL	IV	ML	1 EA	1		01/01/2002	11/12/2012							
10019-0046-04		J3490		11/01/2003	11/12/2012	UNCLASSIFIED DRUGS	FAMOTIDINE (M.D.V.) 10 MG/ML	4 ML	VL	IV	ML	1 EA	1		11/01/2003	11/12/2012							
10019-0046-14		J3490		05/05/2007	02/03/2016	UNCLASSIFIED DRUGS	FAMOTIDINE (MDV) 10 MG/ML	4 ML	VL	IV	ML	1 EA	1		05/05/2007	02/03/2016							
10019-0046-63		J3490		05/05/2007	02/03/2016	UNCLASSIFIED DRUGS	FAMOTIDINE (MDV) 10 MG/ML	20 ML	VL	IV	ML	1 EA	1		05/05/2007	02/03/2016							
10019-0050-06		J3490		01/01/2002	05/03/2012	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (AMP) 50 MCG/ML	5 ML	AM	IJ	ML		1 EA		1	01/01/2002	05/03/2012						
10019-0050-21		J3490		01/01/2002	05/03/2012	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (AMP) 50 MCG/ML	2 ML	AM	IJ	ML		1 EA		1	01/01/2002	05/03/2012						
10019-0050-36		J3490		05/05/2007	02/03/2016	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE 50 MCG/ML	5 ML	AM	IJ	ML		1 EA		1	05/05/2007	02/03/2016						
10019-0050-37		J3490		05/05/2007	02/03/2016	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE 50 MCG/ML	2 ML	AM	IJ	ML		1 EA		1	05/05/2007	02/03/2016						
10019-0050-39		J3490		05/05/2007	02/03/2016	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE 50 MCG/ML	1 ML	AM	IJ	ML		1 EA		1	05/05/2007	02/03/2016						
10019-0050-43		J3490		01/01/2002	05/03/2012	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (AMP) 50 MCG/ML	1 ML	AM	IJ	ML		1 EA		1	01/01/2002	05/03/2012						
10019-0063-03		J0150		06/17/2004	05/03/2012	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS, INSTEAD USE A9270)	ADENOSINE (PF) 3 MG/ML	2 ML	VL	IV	ML		6 MG		0.5	06/17/2004	05/03/2012						
10019-0070-10		J2260		05/05/2007	10/17/2016	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (SDV) 1 MG/ML	10 ML	VL	IV	ML		5 MG		0.2	05/05/2007	10/17/2016						
10019-0070-20		J2260		05/05/2007	10/17/2016	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (SDV) 1 MG/ML	20 ML	VL	IV	ML		5 MG		0.2	05/05/2007	10/17/2016						
10019-0097-44		J2550		05/05/2007	10/17/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL AMERINET CHOICE 25 MG/ML	1 ML	AM	IJ	ML		50 MG		0.5	05/05/2007	10/17/2016						
10019-0102-01		J2060		01/01/2002	04/05/2012	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (S.D.V.) 2 MG/ML	1 ML	VL	IJ	ML		2 MG		1	01/01/2002	04/05/2012						
00078-0422-20		J7527		10/29/2018	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (6X10) 1 MG	60 EA	ST	PO	EA		0.25 MG		4	10/29/2018	99/99/9999						
10019-0102-10		J2060		01/01/2002	04/05/2012	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (M.D.V.) 2 MG/ML	10 ML	VL	IJ	ML		2 MG		1	01/01/2002	04/05/2012						
10019-0102-37		J2060		05/05/2007	02/03/2016	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM 2 MG/ML	10 ML	VL	IJ	ML		2 MG		1	05/05/2007	02/03/2016						
10019-0102-39		J2060		05/05/2007	04/06/2012	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM 2 MG/ML	1 ML	VL	IJ	ML		2 MG		1	05/05/2007	04/06/2012						
10019-0103-01		J2060		01/01/2002	04/05/2012	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (S.D.V.) 4 MG/ML	1 ML	VL	IJ	ML		2 MG		2	01/01/2002	04/05/2012						
10019-0103-10		J2060		01/01/2002	04/05/2012	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (M.D.V.) 4 MG/ML	10 ML	VL	IJ	ML		2 MG		2	01/01/2002	04/05/2012						
10019-0103-37		J2060		05/05/2007	01/31/2014	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM 4 MG/ML	10 ML	VL	IJ	ML		2 MG		2	05/05/2007	01/31/2014						
10019-0103-39		J2060		05/05/2007	01/31/2014	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM 4 MG/ML	25 ML	VL	IJ	ML		2 MG		2	05/05/2007	01/31/2014						
10019-0105-01		J2060		05/03/2006	04/05/2012	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM (25X1ML,SDV) 2 MG/ML	1 ML	VL	IJ	ML		2 MG		1	05/03/2006	04/05/2012						
10019-0105-02		J2060		05/03/2006	04/05/2012	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM (10X10ML,MDV) 2 MG/ML	10 ML	VL	IJ	ML		2 MG		1	05/03/2006	04/05/2012						
10019-0105-44		J2060		05/05/2007	02/03/2016	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM (USP) 2 MG/ML	1 ML	VL	IJ	ML		2 MG		1	05/05/2007	02/03/2016						
10019-0105-71		J2060		05/05/2007	02/03/2016	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM (USP) 2 MG/ML	1 ML	VL	IJ	ML		2 MG		1	05/05/2007	02/03/2016						
10019-0106-01		J2060		05/03/2006	04/05/2012	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM (25X1ML,SDV) 4 MG/ML	1 ML	VL	IJ	ML		2 MG		2	05/03/2006	04/05/2012						
10019-0106-02		J2060		05/03/2006	04/05/2012	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM (10X10ML,MDV) 4 MG/ML	10 ML	VL	IJ	ML		2 MG		2	05/03/2006	04/05/2012						
10019-0106-44		J2060		05/05/2007	02/03/2016	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM 4 MG/ML	1 ML	VL	IJ	ML		2 MG		2	05/05/2007	02/03/2016						
10019-0106-71		J2060		05/05/2007	02/03/2016	INJECTION, LORAZEPAM, 2 MG	NOVAPLUS LORAZEPAM 4 MG/ML	1 ML	VL	IJ	ML		2 MG		2	05/05/2007	02/03/2016						
10019-0159-44		J2175		05/05/2007	10/17/2016	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL 25 MG/ML	1 ML	VL	IJ	ML		100 MG		0.25	05/05/2007	10/17/2016						
60505-6193-01		J2469		09/19/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL 0.05 MG/1 ML	5 ML	VL	IV	ML		25 MCG		2	09/19/2018	99/99/9999						
10019-0160-44		J2175		05/05/2007	10/17/2016	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL 50 MG/ML	1 ML	VL	IJ	ML		100 MG		0.5	05/05/2007	10/17/2016						
10019-0162-44		J2175		05/05/2007	10/17/2016	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL 100 MG/ML	1 ML	VL	IJ	ML		100 MG		1	05/05/2007	10/17/2016						
16729-0223-61		J9330		08/13/2018	99/99/9999	INJECTION, TEMSIRILIMUS, 1 MG	TEMSIRILIMUS (WITH DILUENT) 25 MG/1 ML	1 ML	VL	IV	ML		1 MG		25	08/13/2018	99/99/9999						
00078-0422-61		J7527		10/29/2018	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (1X1) 1 MG	1 EA	ST	PO	EA		0.25 MG		4	10/29/2018	99/99/9999						
10019-0177-37		J2270		05/05/2007	10/17/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE 8 MG/ML	1 ML	AM	IJ	ML		10 MG		0.8	05/05/2007	10/17/2016						
10019-0178-36		J2270		05/05/2007	02/03/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (MDV) 10 MG/ML	10 ML	NA	IJ	ML		10 MG		1	05/05/2007	02/03/2016						
10019-0178-39		J2270		05/05/2007	10/17/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE 10 MG/ML	1 ML	VL	IJ	ML		10 MG		1	05/05/2007	10/17/2016						
10019-0179-36		J2271		05/05/2007	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (MDV) 15 MG/ML	20 ML	NA	IJ	ML		100 MG		0.15	05/05/2007	12/31/2014						
10019-0270-10		J2710		01/01/2002	09/06/2012	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (M.D.V.) 1 MG/ML	10 ML	VL	IJ	ML		0.5 MG		2	01/01/2002	09/06/2012						
10019-0271-10		J2710		01/01/2002	10/09/2012	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (M.D.V.) 0.5 MG/ML	10 ML	VL	IJ	ML		0.5 MG		1	01/01/2002	10/09/2012						
10019-0291-02		J2590		05/07/2007	05/04/2012	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN (1MLX25,SDV,USP) 10 U/ML	1 ML	VL	IJ	ML		10 U		1	05/07/2007	05/04/2012						
10019-0291-04		J2590		05/07/2007	05/03/2012	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN (10MLX25,MDV,USP) 10 U/ML	10 ML	VL	IJ	ML		10 U		1	05/07/2007	05/03/2012						
10019-0291-12		J2590		05/29/2007	02/27/2013	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN (SDV,USP) 10 U/ML	1 ML	VL	IJ	ML		10 U		1	05/29/2007	02/27/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
10019-0291-71		J2590		05/29/2007	02/27/2013	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN (MDV,USP) 10 U/ML	10	ML	VL	IJ	ML	10 U		1	05/29/2007	02/27/2013							
10019-0450-39		J2765		05/05/2007	04/30/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL 5 MG/ML	2	ML	VL	IV	ML	10 MG		0.5	05/05/2007	04/30/2013							
10019-0506-10		J3490		01/01/2002	04/02/2012	UNCLASSIFIED DRUGS	BUMETANIDE (M.D.V.) 0.25 MG/ML	10	ML	VL	IJ	ML	1 EA		1	01/01/2002	04/02/2012							
10019-0506-45		J3490		01/01/2002	04/02/2012	UNCLASSIFIED DRUGS	BUMETANIDE (S.D.V.) 0.25 MG/ML	4	ML	VL	IJ	ML	1 EA		1	01/01/2002	04/02/2012							
00409-2504-10		J2469		11/15/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL (PF,LATEX-FREE) 0.05 MG/1 ML	5	ML	VL	IV	ML	25 MCG		2	11/15/2018	99/99/9999							
10019-0630-33		J0295		05/05/2007	10/31/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN/SULBACTAM 2 GM-1 GM	1	EA	VL	IJ	EA	1.5 GM		2	05/05/2007	10/31/2013							
10019-0631-31		J0295		05/05/2007	10/31/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN/SULBACTAM 1 GM-0.5 GM	1	EA	VL	IJ	EA	1.5 GM		1	05/05/2007	10/31/2013							
10019-0633-02		J0295		03/10/2006	05/15/2012	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMERINET CHOICE AMPICILLIN AND SULBACTAM 2 GM-1 GM	1	EA	VL	IJ	EA	1.5 GM		2	03/10/2006	05/15/2012							
10019-0633-33		J0295		05/05/2007	07/30/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMERINET CHOICE AMPICILLIN AND SULBACTAM 2 GM-1 GM	1	EA	VL	IJ	EA	1.5 GM		2	05/05/2007	07/30/2013							
10019-0634-01		J0295		03/10/2006	02/01/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMERINET CHOICE AMPICILLIN AND SULBACTAM (10X10MLVIALS) 1 GM-0.5 GM	1	EA	VL	IJ	EA	1.5 GM		1	03/10/2006	02/01/2013							
10019-0634-31		J0295		05/05/2007	10/17/2016	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMERINET CHOICE AMPICILLIN AND SULBACTAM 1 GM-0.5 GM	1	EA	VL	IJ	EA	1.5 GM		1	05/05/2007	10/17/2016							
10019-0635-03		J0295		12/14/2005	02/01/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMERINET CHOICE AMPICILLIN AND SULBACTAM (PHARMACY BULK) 10 GM-5 GM	1	EA	VL	IJ	EA	1.5 GM		10	12/14/2005	02/01/2013							
10019-0636-31		J0295		05/05/2007	02/03/2016	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	NOVAPLUS AMPICILLIN AND SULBACTAM 1 GM-0.5 GM	1	EA	VL	IJ	EA	1.5 GM		1	05/05/2007	02/03/2016							
10019-0637-33		J0295		05/05/2007	02/03/2016	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	NOVAPLUS AMPICILLIN AND SULBACTAM 2 GM-1 GM	1	EA	VL	IJ	EA	1.5 GM		2	05/05/2007	02/03/2016							
10019-0638-03		J0295		05/02/2006	05/15/2012	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	NOVAPLUS AMPICILLIN AND SULBACTAM (USP,PHARMACYBULK) 10 GM-5 GM	1	EA	VL	IV	EA	1.5 GM		10	05/02/2006	05/15/2012							
69097-0319-87	KO	J7626	KO	11/14/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.5 MG/2 ML	2	ML	AM	IH	ML	0.5 MG		0.5	11/14/2017	99/99/9999							
10019-0688-04		J0696		07/05/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 2 GM	1	EA	VL	IJ	EA	250 MG		8	07/05/2005	99/99/9999							
10019-0688-27		J0696		05/05/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 2 GM	1	EA	VL	IJ	EA	250 MG		8	05/05/2007	99/99/9999							
10019-0689-05		J0696		10/05/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP,PHARMACY BULK) 10 GM	1	EA	VL	IV	EA	250 MG		40	10/05/2006	99/99/9999							
10019-0905-17		J2405		05/05/2007	10/17/2016	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (LATEX-FREE) 2 MG/ML	1	ML	VL	IJ	ML	1 MG		2	05/05/2007	10/17/2016							
60505-6113-06		J9201		02/23/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 38 MG/1 ML	5.26	ML	VL	IV	ML	200 MG		0.19	02/23/2018	99/99/9999							
10019-0906-63		J2405		05/05/2007	10/17/2016	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (LATEX-FREE) 2 MG/ML	1	ML	NA	IJ	ML	1 MG		2	05/05/2007	10/17/2016							
10019-0925-01		J9208		09/12/2005	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (SDV,30ML VIAL) 1 GM	1	EA	VL	IV	EA	1 GM		1	09/12/2005	99/99/9999							
10019-0925-82		J9208		05/05/2007	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (SDV,30ML) 1 GM	1	EA	VL	IV	EA	1 GM		1	05/05/2007	99/99/9999							
10019-0926-02		J9208		09/12/2005	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (SDV,75ML VIAL) 3 GM	1	EA	VL	IV	EA	1 GM		3	09/12/2005	99/99/9999							
10019-0926-16		J9208		05/05/2007	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (SDV,75ML) 3 GM	1	EA	VL	IV	EA	1 GM		3	05/05/2007	99/99/9999							
10019-0934-01		J9206		02/21/2008	02/03/2016	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML,SDV,AMBER GLASS) 20 MG/ML	2	ML	VL	IV	ML	20 MG		1	02/21/2008	02/03/2016							
10019-0934-02		J9206		02/21/2008	02/03/2016	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML,SDV,AMBER GLASS) 20 MG/ML	5	ML	VL	IV	ML	20 MG		1	02/21/2008	02/03/2016							
10019-0934-17		J9206		02/21/2008	02/03/2016	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML,SDV,INNER NDC) 20 MG/ML	2	ML	VL	IV	ML	20 MG		1	02/21/2008	02/03/2016							
10019-0934-79		J9206		02/21/2008	02/03/2016	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML,SDV,INNER NDC) 20 MG/ML	5	ML	VL	IV	ML	20 MG		1	02/21/2008	02/03/2016							
10019-0953-01		J9209		03/15/2004	99/99/9999	INJECTION, MESNA, 200 MG	MESNA (S.D.V.) 100 MG/ML	10	ML	VL	IV	ML	200 MG		0.5	03/15/2004	99/99/9999							
10019-0953-02		J9209		03/15/2004	99/99/9999	INJECTION, MESNA, 200 MG	MESNA (S.D.V.) 100 MG/ML	10	ML	VL	IV	ML	200 MG		0.5	03/15/2004	99/99/9999							
10019-0953-62		J9209		05/05/2007	99/99/9999	INJECTION, MESNA, 200 MG	MESNA 100 MG/ML	1	ML	VL	IV	ML	200 MG		0.5	05/05/2007	99/99/9999							
10106-0061-01		J9017		01/01/2002	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (A.C.S., REAGENT)	1	EA	NA	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
10106-0061-04		J9017		01/01/2002	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (A.C.S., REAGENT)	1	EA	NA	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
10106-0062-01		J9017		01/01/2002	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (REAGENT)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
10106-0062-04		J9017		01/01/2002	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (REAGENT)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
10106-1080-01		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BENZOCAINE (FINE, U.S.P.)	1	EA	BO	NA	GM	1 EA		1	01/01/2002	99/99/9999							
10106-1649-01		J0706		01/01/2002	10/17/2016	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATED (PURIFIED)	1	EA	BO	NA	GM	5 MG		200	01/01/2002	10/17/2016							
10106-1649-04		J0706		01/01/2002	10/17/2016	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATED (PURIFIED)	1	EA	BO	NA	GM	5 MG		200	01/01/2002	10/17/2016							
10106-2506-01		J3475		01/01/2002	10/17/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE ANHYDROUS (REAGENT)	1	EA	BO	NA	GM	500 MG		2	01/01/2002	10/17/2016							
10106-2506-05		J3475		01/01/2002	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE ANHYDROUS (REAGENT)	1	EA	BO	NA	GM	500 MG		2	01/01/2002	99/99/9999							
10106-2555-05		J2150		01/01/2002	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	1	EA	BO	NA	GM	50 ML		0.08	01/01/2002	99/99/9999							
10106-3046-01		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P., F.C.C.)	1	EA	BO	NA	GM	2 MEQ		6.71141	01/01/2002	99/99/9999							
10106-3046-05		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P., F.C.C.)	1	EA	BO	NA	GM	2 MEQ		6.71141	01/01/2002	99/99/9999							
10106-3052-01		J3480		01/01/2002	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P., F.C.C.)	1	EA	BO	NA	GM	2 MEQ		6.71141	01/01/2002	10/17/2016							
10106-3052-05		J3480		01/01/2002	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P., F.C.C.)	1	EA	BO	NA	GM	2 MEQ		6.71141	01/01/2002	10/17/2016							
10106-3343-01		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P., F.C.C.)	1	EA	BO	NA	GM	100 MG		10	01/01/2004	99/99/9999							
10106-4206-01		J3350																						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
10106-4206-05		J3350		01/01/2002	99/99/9999	INJECTION, UREA, UP TO 40 MG	UREA (U.S.P.)	1	EA	BO	NA	GM	40	GM	0.025	01/01/2002	99/99/9999						
10106-8994-01		J3520		01/01/2002	99/99/9999	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM (U.S.P.)	1	EA	BO	NA	GM	150	MG	6.666666	01/01/2002	99/99/9999						
10106-9224-01		J1212		01/01/2002	99/99/9999	INJECTION, DMSO, DIMETHYL SULFOXIDE, 50%, 50 ML	DIMETHYL SULFOXIDE (A.C.S., REAGENT)	500	ML	EA	NA	ML	50	%	0.02	01/01/2002	99/99/9999						
10135-0149-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0149-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	1000	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0149-24		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	24	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0149-61		Q0163		11/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	10	EA	BO	PO	EA	50	MG	0.5	11/01/2002	99/99/9999						
10135-0151-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0151-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 25 MG	1000	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0151-24		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 25 MG	24	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0151-50		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 25 MG	50	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0151-52		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (BOXED,CAPLET) 25 MG	24	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0151-57		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (BOXED,CAPLET) 25 MG	100	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10135-0156-01		Q0163		11/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50	MG	1	11/01/2002	99/99/9999						
10135-0156-10		Q0163		11/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	1000	EA	BO	PO	EA	50	MG	1	11/01/2002	99/99/9999						
10135-0156-13		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BX	PO	EA	50	MG	1	01/01/2002	99/99/9999						
10135-0166-13		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (BLISTER PACK,CAPLET) 25 MG	100	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
10139-0062-02		J9250		07/02/2007	02/14/2013	METHOTREXATE SODIUM, 5 MG	METHOTREXATE (USP,SDV,PF) 25 MG/ML	2	ML	VL	IJ	ML	5	MG	5	07/02/2007	02/14/2013						
10139-0062-10		J9250		06/07/2007	08/04/2013	METHOTREXATE SODIUM, 5 MG	METHOTREXATE (USP,SDV,PF) 25 MG/ML	10	ML	VL	IJ	ML	5	MG	5	06/07/2007	08/04/2013						
10139-0062-40		J9250		06/07/2007	02/06/2013	METHOTREXATE SODIUM, 5 MG	METHOTREXATE (USP,SDV,PF) 25 MG/ML	40	ML	VL	IJ	ML	5	MG	5	06/07/2007	02/06/2013						
10139-0063-01		J9190		07/02/2007	06/30/2014	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (USP,BULK) 50 MG/ML	100	ML	VL	IV	ML	500	MG	0.1	07/02/2007	06/30/2014						
10139-0063-11		J9190		06/11/2007	06/30/2014	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (USP,SDV,10MLX10) 50 MG/ML	10	ML	VL	IV	ML	500	MG	0.1	06/11/2007	06/30/2014						
10139-0063-12		J9190		06/11/2007	06/30/2014	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (USP,SDV,20MLX10) 50 MG/ML	20	ML	VL	IV	ML	500	MG	0.1	06/11/2007	06/30/2014						
10139-0063-50		J9190		06/07/2007	06/30/2014	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (USP) 50 MG/ML	50	ML	VL	IV	ML	500	MG	0.1	06/07/2007	06/30/2014						
10139-0070-11		J0295		07/03/2007	04/29/2013	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP) 1 GM-0.5 GM	1	EA	VL	IJ	EA	1.5	GM	1	07/03/2007	04/29/2013						
10139-0071-10		J0295		07/03/2007	11/12/2012	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP) 2 GM-1 GM	1	EA	VL	IJ	EA	1.5	GM	2	07/03/2007	11/12/2012						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
10158-0042-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	NYTOL QUICKGELS MAXIMUM STRENGTH (SOFTGEL) 50 MG	8 EA	BX	PO	EA		50 MG		1	01/01/2002	99/99/9999						
10158-0043-02		Q0163		01/01/2002	09/30/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	NYTOL QUICKCAPS 25 MG	16 EA	BX	PO	EA		50 MG		0.5	01/01/2002	09/30/2017						
10158-0043-04		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	NYTOL QUICKCAPS 25 MG	32 EA	BX	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
10158-0043-06		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	NYTOL QUICKCAPS 25 MG	72 EA	BX	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
10267-0835-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
10267-0835-04		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	1000 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
10267-0836-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100 EA	BO	PO	EA		50 MG		1	01/01/2002	99/99/9999						
10267-0836-04		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	1000 EA	BO	PO	EA		50 MG		1	01/01/2002	99/99/9999						
10454-0710-10		J0587		08/01/2005	99/99/9999	INJECTION, RIMABOTULINUMTOXINB, 100 UNITS	MYOBLOC (PF) 2500 U/0.5 ML	0.5 ML	VL	IM	ML		100 U		50	08/01/2005	99/99/9999						
10454-0711-10		J0587		08/01/2005	99/99/9999	INJECTION, RIMABOTULINUMTOXINB, 100 UNITS	MYOBLOC (PF) 5000 U/ML	1 ML	VL	IM	ML		100 U		50	08/01/2005	99/99/9999						
10454-0712-10		J0587		06/30/2006	99/99/9999	INJECTION, RIMABOTULINUMTOXINB, 100 UNITS	MYOBLOC 5000 U/ML	2 ML	VL	IM	ML		100 U		50	06/30/2006	99/99/9999						
10702-0002-01		Q0169		05/10/2007	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 12.5 MG	100 EA	BO	PO	EA		12.5 MG		1	05/10/2007	99/99/9999						
10702-0003-01		Q0170		01/16/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	100 EA	BO	PO	EA		25 MG		1	01/16/2007	12/31/2013						
10702-0003-10		Q0170		01/16/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	1000 EA	BO	PO	EA		25 MG		1	01/16/2007	12/31/2013						
10702-0004-01		Q0170		01/16/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 50 MG	100 EA	BO	PO	EA		25 MG		2	01/16/2007	12/31/2013						
10892-0112-65		Q0163		01/01/2002	02/08/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DYTUSS 12.5 MG/5 ML	480 ML	BO	PO	ML		50 MG		0.05	01/01/2002	02/08/2013						
10956-0750-24		Q0163		11/02/2004	06/18/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SLEEP-ETTES D 50 MG	24 EA	NA	PO	EA		50 MG		1	11/02/2004	06/18/2013						
10956-0750-48		Q0163		11/02/2004	06/18/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SLEEP-ETTES D 50 MG	48 EA	BO	PO	EA		50 MG		1	11/02/2004	06/18/2013						
10956-0751-24		Q0163		11/02/2004	06/18/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALER-DRYL 50 MG	24 EA	BX	PO	EA		50 MG		1	11/02/2004	06/18/2013						
10956-0751-48		Q0163		11/02/2004	06/18/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALER-DRYL 50 MG	48 EA	BO	PO	EA		50 MG		1	11/02/2004	06/18/2013						
11098-0526-03		J0470		01/01/2002	06/13/2012	INJECTION, DIMERCAPROL, PER 100 MG	BAL IN OIL (AMP) 10% HEPARIN SODIUM (HEMOCHRON RXDX,VIAL) 1000 U/ML	3 ML	AM	IM	ML		100 MG		1	01/01/2002	06/13/2012						
11743-0210-02		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (HEMOCHRON RXDX,VIAL) 1000 U/ML	10 ML	VL	IJ	ML		1000 U		1	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
11822-0527-10		Q0163		05/02/2006	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	RITE AID ALLERGY (AF,SF,DYE-FREE) 12.5 MG/5 ML	118	ML	NA	PO	ML	50	MG	0.05	05/02/2006	99/99/9999						
11845-0896-01		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALLERGY RELIEF MEDICINE 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	02/03/2016						
16729-0365-66		J2469		03/23/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL 0.05 MG/1 ML	5	ML	VL	IV	ML	25	MCG									
13411-0131-01		Q0144		08/23/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	10	EA	BO	PO	EA	1	GM	0.25	08/23/2006	99/99/9999						
13411-0131-03		Q0144		06/01/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	30	EA	BO	PO	EA	1	GM	0.25	06/01/2005	99/99/9999						
13411-0131-06		Q0144		08/23/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	60	EA	BO	PO	EA	1	GM	0.25	08/23/2006	99/99/9999						
13411-0131-09		Q0144		08/23/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	90	EA	BO	PO	EA	1	GM	0.25	08/23/2006	99/99/9999						
13411-0131-15		Q0144		08/23/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	15	EA	BO	PO	EA	1	GM	0.25	08/23/2006	99/99/9999						
13411-0182-01		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	10	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0182-03		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0182-06		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	60	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0182-09		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	90	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0182-10		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0183-01		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	10	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0183-06		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0183-09		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	60	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0183-10		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	90	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13411-0183-10		J8499		08/23/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100	EA	BO	PO	EA	1	EA	1	08/23/2006	99/99/9999						
13533-0631-02		J2790		12/21/2005	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, FULL DOSE, 300 MICROGRAMS (1500 I.U.)	HYPERRHO S/D (FULL DOSE,PF)	1	EA	SR	IM	EA	300	MCG	1	12/21/2005	99/99/9999						
13533-0631-06		J2792		12/21/2005	10/31/2013	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT DETERGENT, 100 IU	HYPERRHO S/D (MINI-DOSE)	0.17	ML	SR	IM	ML	100	IU	12.5	12/21/2005	10/31/2013						
13533-0634-02		J1670		10/14/2006	99/99/9999	INJECTION, TETANUS IMMUNE GLOBULIN, HUMAN, UP TO 250 UNITS	HYPERTET S/D (PF) 250 U	1	ML	SR	IM	ML	250	U	1	10/14/2006	99/99/9999						
13533-0635-04		J1460		10/04/2005	99/99/9999	INJECTION, GAMMA GLOBULIN, INTRAMUSCULAR, 1 CC	GAMASTAN S/D (S.D.V.,PF)	2	ML	VL	IM	ML	1	ML	1	10/04/2005	99/99/9999						
13533-0635-12		J1460		10/04/2005	99/99/9999	INJECTION, GAMMA GLOBULIN, INTRAMUSCULAR, 1 CC	GAMASTAN S/D (S.D.V.,PF)	10	ML	VL	IM	ML	1	ML	1	10/04/2005	99/99/9999						
13533-0645-12		J1561		01/01/2008	03/24/2013	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX (PF) 100 MG/ML	10	ML	VL	IV	ML	500	MG	0.2	01/01/2008	03/24/2013						
13533-0645-15		J1561		01/01/2008	04/19/2013	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX (PF) 100 MG/ML	25	ML	VL	IV	ML	500	MG	0.2	01/01/2008	04/19/2013						
13533-0645-20		J1561		01/01/2008	06/26/2014	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX (PF) 100 MG/ML	50	ML	VL	IV	ML	500	MG	0.2	01/01/2008	06/26/2014						
13533-0645-24		J1561		01/01/2008	10/17/2013	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX (PF) 100 MG/ML	200	ML	VL	IV	ML	500	MG	0.2	01/01/2008	10/17/2013						
13533-0645-71		J1561		01/01/2008	10/22/2013	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX (PF) 100 MG/ML	100	ML	VL	IV	ML	500	MG	0.2	01/01/2008	10/22/2013						
15054-1040-05		J2170		01/01/2007	99/99/9999	INJECTION, MECASERMIN, 1 MG	INCRELEX (10X4ML,M.D.V.) 10 MG/ML	4	ML	VL	SC	ML	1	MG	10	01/01/2007	99/99/9999						
49502-0500-02		J0171		05/02/2001	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPIPEN AUTO-INJECTOR (W/TRAINER DEVICE) 0.3 MG/0.3 ML	2	EA	PG	IJ	EA	0.1	MG	3	05/02/2001	99/99/9999						
00378-9691-52	KO	J7614	KO	07/23/2018	99/99/9999	INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.63 MG/3 ML	3	ML	VL	IJ	ML	0.5	MG	0.42	07/23/2018	99/99/9999						
15927-3220-00		J7799		09/08/2003	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	EPINEPHRINE (BASE) MORPHINE SULFATE (PF,LATEX-FREE) 2 MG/1 ML	1	EA	BO	NA	GM	1	EA	1	09/08/2003	99/99/9999						
63323-0452-01		J2270		05/23/2018	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (PF,LATEX-FREE) 4 MG/1 ML	1	ML	VL	IJ	ML	10	MG	0.2	05/23/2018	99/99/9999						
63323-0454-01		J2270		05/23/2018	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (PF,LATEX-FREE) 4 MG/1 ML	1	ML	VL	IJ	ML	10	MG	0.4	05/23/2018	99/99/9999						
60710-0015-50		J3480		09/05/2018	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE PROAMP 2 MEQ/1 ML	10	ML	AM	IV	ML	2	MEQ	1	09/05/2018	99/99/9999						
63323-0637-10		J9017		09/19/2018	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (10X10 SDV,PF,LATEX-FREE) 1 MG/1 ML	10	ML	VL	IV	ML	1	MG	1	09/19/2018	99/99/9999						
63323-0673-89		J2469		09/07/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	SIMPLIST PALONOSETRON HCL 0.05 MG/1 ML	5	ML	SR	IV	ML	25	MCG	2	09/07/2018	99/99/9999						
16252-0536-08		J8515		05/01/2008	07/29/2014	CABERGOLINE, ORAL, 0.25 MG	CABERGOLINE 0.5 MG	8	EA	BO	PO	EA	0.25	MG	2	05/01/2008	07/29/2014						
69097-0321-87	KO	J7626	KO	11/14/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 1 MG/2 ML	2	ML	AM	IH	ML	0.5	MG	1	11/14/2017	99/99/9999						
69097-0173-64		J7620		07/01/2015	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (60X3ML 5 VIALS/POUCH) 3 MG/3 ML-0.5 MG/3 ML	3	ML	VL	IH	ML	3	MG	0.33333	07/01/2015	99/99/9999						
63323-0455-01		J2270		05/23/2018	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (PF,LATEX-FREE) 5 MG/1 ML	1	ML	VL	IJ	ML	10	MG	0.5	05/23/2018	99/99/9999						
63323-0458-01		J2270		05/23/2018	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (PF,LATEX-FREE) 8 MG/1 ML	1	ML	VL	IJ	ML	10	MG	0.8	05/23/2018	99/99/9999						
16252-0547-33		J7620		12/31/2007	07/02/2013	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE/ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	30	ML	PC	IH	ML	3	MG	0.33333	12/31/2007	07/02/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
16252-0547-66		J7620		12/31/2007	05/12/2013	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE/ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	60	ML	PC	IH	ML	3	MG	0.33333	12/31/2007	05/12/2013						
16477-0510-08		J8499		04/30/2008	07/14/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	MILLIPRED (1X237ML AF.DYE-FREE) 10 MG/5 ML	237	ML	BO	PO	ML	1	EA	1	04/30/2008	07/14/2014						
16590-0003-30		J8499		02/01/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	02/01/2006	06/01/2014						
16590-0003-60		J8499		02/01/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	60	EA	BO	PO	EA	1	EA	1	02/01/2006	06/01/2014						
16590-0078-20		Q0163		02/01/2006	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	20	EA	BO	PO	EA	50	MG	0.5	02/01/2006	06/01/2014						
16590-0079-20		Q0163		02/01/2006	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 50 MG	20	EA	BO	PO	EA	50	MG	1	02/01/2006	06/01/2014						
16590-0149-21		J7509		01/01/2006	06/01/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPRED-DP 4 MG	21	EA	DP	PO	EA	4	MG	1	01/01/2006	06/01/2014						
16590-0191-10		Q0170		04/01/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10	EA	BO	PO	EA	25	MG	1	04/01/2007	12/31/2013						
16590-0191-15		Q0170		02/01/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	15	EA	BO	PO	EA	25	MG	1	02/01/2006	12/31/2013						
16590-0191-20		Q0170		06/01/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	20	EA	BO	PO	EA	25	MG	1	06/01/2006	12/31/2013						
16590-0191-30		Q0170		02/01/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	30	EA	BO	PO	EA	25	MG	1	02/01/2006	12/31/2013						
16590-0191-60		Q0170		02/01/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	60	EA	BO	PO	EA	25	MG	1	02/01/2006	12/31/2013						
16590-0191-90		Q0170		02/01/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	90	EA	BO	PO	EA	25	MG	1	02/01/2006	12/31/2013						
16590-0248-06		Q0144		02/01/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX Z-PAK 250 MG	6	EA	DP	PO	EA	1	GM	0.25	02/01/2006	06/01/2014						
16590-0326-10		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	10	EA	BO	PO	EA	5	MG	4	06/01/2006	06/01/2014						
16590-0326-20		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	20	EA	BO	PO	EA	5	MG	4	06/01/2006	06/01/2014						
16590-0326-21		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	21	EA	BO	PO	EA	5	MG	4	06/01/2006	06/01/2014						
16590-0326-30		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	06/01/2006	06/01/2014						
16590-0326-45		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	45	EA	BO	PO	EA	5	MG	4	06/01/2006	06/01/2014						
16590-0326-60		J7506		11/01/2007	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	60	EA	BO	PO	EA	5	MG	4	11/01/2007	06/01/2014						
16590-0327-10		Q0165		04/01/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	10	MG	1	04/01/2007	12/31/2013						
16590-0357-09		Q0177		05/01/2006	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	9	EA	BO	PO	EA	25	MG	1	05/01/2006	06/01/2014						
16590-0357-12		Q0177		05/01/2006	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	12	EA	BO	PO	EA	25	MG	1	05/01/2006	06/01/2014						
16590-0357-20		Q0177		05/01/2006	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20	EA	BO	PO	EA	25	MG	1	05/01/2006	06/01/2014						
16590-0357-30		Q0177		05/01/2006	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	BO	PO	EA	25	MG	1	05/01/2006	06/01/2014						
16590-0362-06		Q0144		12/01/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6	EA	DP	PO	EA	1	GM	0.25	12/01/2006	06/01/2014						
16590-0370-20		J8499		06/01/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	20	EA	BO	PO	EA	1	EA	1	06/01/2006	06/01/2014						
16590-0370-30		J8499		06/01/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30	EA	BO	PO	EA	1	EA	1	06/01/2006	06/01/2014						
16590-0370-40		J8499		06/01/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	40	EA	BO	PO	EA	1	EA	1	06/01/2006	06/01/2014						
16590-0404-10		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	10	EA	BO	PO	EA	5	MG	2	06/01/2006	06/01/2014						
16590-0404-20		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	20	EA	BO	PO	EA	5	MG	2	06/01/2006	06/01/2014						
16590-0404-21		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21	EA	BO	PO	EA	5	MG	2	06/01/2006	06/01/2014						
16590-0404-30		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	30	EA	BO	PO	EA	5	MG	2	06/01/2006	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
16590-0404-45		J7506		06/01/2006	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	45	EA	BO	PO	EA	5	MG	2	06/01/2006	06/01/2014						
16714-0221-30		Q0166		05/15/2008	99/99/9999	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (FILM-COATED) 1 MG	2	EA	BX	PO	EA	1	MG	1	05/15/2008	99/99/9999						
16714-0221-32		Q0166		05/15/2008	99/99/9999	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (2X10 FILM-COATED) 1 MG	20	EA	BX	PO	EA	1	MG	1	05/15/2008	99/99/9999						
25021-0185-10		J1570		04/16/2018	99/99/9999	INJECTION, GANCICLOVIR SODIUM, 500 MG	GANCICLOVIR (PF) 50 MG/1 ML	10	ML	VL	IV	ML	500	MG	0.1	04/16/2018	99/99/9999						
17317-0022-01		J0280		01/01/2002	01/01/2014	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1	EA	VL	NA	GM	250	MG	4	01/01/2002	01/01/2014						
17317-0022-04		J0280		01/01/2002	01/01/2014	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1	EA	VL	NA	GM	250	MG	4	01/01/2002	01/01/2014						
17317-0022-05		J0280		01/01/2002	01/01/2014	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1	EA	VL	NA	GM	250	MG	4	01/01/2002	01/01/2014						
17317-0036-02		J7636		01/01/2002	01/01/2014	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	VL	NA	GM	1	MG	1000	01/01/2002	01/01/2014						
17317-0036-02	KO	J7636	KO	01/01/2002	01/01/2014	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	VL	NA	GM	1	MG	1000	01/01/2002	01/01/2014						
17317-0036-05		J7636		01/01/2002	01/01/2014	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	VL	NA	GM	1	MG	1000	01/01/2002	01/01/2014						
17317-0036-05	KO	J7636	KO	01/01/2002	01/01/2014	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	VL	NA	GM	1	MG	1000	01/01/2002	01/01/2014						
17317-0036-07		J7636		01/01/2002	01/01/2014	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	VL	NA	GM	1	MG	1000	01/01/2002	01/01/2014						
17317-0036-07	KO	J7636	KO	01/01/2002	01/01/2014	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	VL	NA	GM	1	MG	1000	01/01/2002	01/01/2014						
17317-0049-01		J3490		01/01/2002	01/01/2014	UNCLASSIFIED DRUGS	BENZOCANE (U.S.P.)	1	EA	NA	NA	GM	1	EA	1	01/01/2002	01/01/2014						
17317-0049-04		J3490		01/01/2002	01/01/2014	UNCLASSIFIED DRUGS	BENZOCANE (U.S.P.)	1	EA	NA	NA	GM	1	EA	1	01/01/2002	01/01/2014						
17317-0049-05		J3490		01/01/2002	01/01/2014	UNCLASSIFIED DRUGS	BENZOCANE (U.S.P.)	1	EA	NA	NA	GM	1	EA	1	01/01/2002	01/01/2014						
17317-0073-01		J0706		01/01/2002	01/01/2014	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATED (PURIFIED)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2014						
17317-0073-04		J0706		01/01/2002	01/01/2014	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATED (PURIFIED)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2014						
17317-0073-05		J0706		01/01/2002	01/01/2014	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATED (PURIFIED)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2014						
17317-0073-08		J0706		01/01/2002	01/01/2014	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATED (PURIFIED)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2014						
17317-0146-03		J1200		01/01/2002	01/01/2014	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (U.S.P.)	1	EA	NA	NA	GM	50	MG	20	01/01/2002	01/01/2014						
17317-0146-05		J1200		01/01/2002	01/01/2014	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (U.S.P.)	1	EA	NA	NA	GM	50	MG	20	01/01/2002	01/01/2014						
17317-0146-06		J1200		01/01/2002	01/01/2014	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (U.S.P.)	1	EA	NA	NA	GM	50	MG	20	01/01/2002	01/01/2014						
17317-0199-02		J1700		01/01/2002	01/01/2014	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P., MICRONIZED)	1	EA	BO	NA	GM	25	MG	40	01/01/2002	01/01/2014						
17317-0199-03		J1700		01/01/2002	01/01/2014	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P., MICRONIZED)	1	EA	BO	NA	GM	25	MG	40	01/01/2002	01/01/2014						
17317-0199-08		J1700		01/01/2002	01/01/2014	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P., MICRONIZED)	1	EA	BO	NA	GM	25	MG	40	01/01/2002	01/01/2014						
17317-0345-01		J3475		01/01/2002	01/01/2014	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (U.S.P.)	1	EA	BO	NA	GM	500	MG	2	01/01/2002	01/01/2014						
17317-0345-05		J3475		01/01/2002	01/01/2014	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE HEPTAHYDRATE (U.S.P.)	1	EA	BO	NA	GM	500	MG	2	01/01/2002	01/01/2014						
17317-0345-08		J3475		01/01/2002	01/01/2014	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (U.S.P.)	1	EA	BO	NA	GM	500	MG	2	01/01/2002	01/01/2014						
17317-0346-01		J3475		01/01/2002	01/01/2014	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (PURIFIED, U.S.P./F.C.C.)	1	EA	FC	NA	GM	500	MG	2	01/01/2002	01/01/2014						
17317-0346-05		J3475		01/01/2002	01/01/2014	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (PURIFIED, U.S.P./F.C.C.)	1	EA	FC	NA	GM	500	MG	2	01/01/2002	01/01/2014						
17317-0346-08		J3475		01/01/2002	01/01/2014	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (PURIFIED, U.S.P./F.C.C.)	1	EA	FC	NA	GM	500	MG	2	01/01/2002	01/01/2014						
17317-0398-01		J2440		01/01/2002	01/01/2014	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1	EA	BO	NA	GM	60	MG	16.66666	01/01/2002	01/01/2014						
17317-0398-04		J2440		01/01/2002	01/01/2014	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1	EA	BO	NA	GM	60	MG	16.66666	01/01/2002	01/01/2014						
17317-0413-01		J2560		01/01/2002	01/01/2014	INJECTION, PHENOBARBITAL SODIUM, UP TO 120 MG	SODIUM PHENOBARBITAL (U.S.P.)	1	EA	BO	NA	GM	120	MG	8.33333	01/01/2002	01/01/2014						
17317-0417-02		J7799		01/01/2002	01/01/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	PHENYLEPHRINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	EA	1	01/01/2002	01/01/2014						
17317-0417-03		J7799		01/01/2002	01/01/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	PHENYLEPHRINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	EA	1	01/01/2002	01/01/2014						
17317-0417-05		J7799		01/01/2002	01/01/2014	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	PHENYLEPHRINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	EA	1	01/01/2002	01/01/2014						
17317-0438-01		J3480		01/01/2002	01/01/2014	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P., F.C.C.)	1	EA	BO	NA	GM	2	MEQ	6.71141	01/01/2002	01/01/2014						
17317-0438-05		J3480		01/01/2002	01/01/2014	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P./F.C.C.)	1	EA	FC	NA	GM	2	MEQ	6.71141	01/01/2002	01/01/2014						
17317-0438-08		J3480		01/01/2002	01/01/2014	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P./F.C.C.)	1	EA	FC	NA	GM	2	MEQ	6.71141	01/01/2002	01/01/2014						
17317-0447-02		J7510		01/01/2002	01/01/2014	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2014						
17317-0447-03		J7510		01/01/2002	01/01/2014	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2014						
17317-0455-02		J3415		01/01/2004	01/01/2014	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	01/01/2004	01/01/2014						
17317-0455-03		J3415		01/01/2004	01/01/2014	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	01/01/2004	01/01/2014						
17317-0455-05		J3415		01/01/2004	01/01/2014	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	01/01/2004	01/01/2014						
17317-0455-06		J3415		01/01/2004	01/01/2014	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	01/01/2004	01/01/2014						
17317-0477-08		J7510		01/01/2002	01/01/2014	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2014						
17317-0567-02		J3140		01/01/2002	01/01/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P., MICRONIZED)	1	EA	BO	NA	GM	50	MG	20	01/01/2002	01/01/2014						
17317-0567-03		J3140		01/01/2002	01/01/2014																		

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
17714-0020-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
17714-0020-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	1000	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
17714-0021-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
17714-0021-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	1000	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
17714-0042-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 25 MG	100	EA	NA	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
17714-0042-24		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPLETE ALLERGY MEDICATION (CAPLET) 25 MG	24	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
18111-0002-02		J9206		02/28/2008	11/30/2012	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML) 20 MG/ML	2	ML	VL	IV	ML	20	MG	1	02/28/2008	11/30/2012						
18111-0002-03		J9206		02/28/2008	11/30/2012	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML) 20 MG/ML	5	ML	VL	IV	ML	20	MG	1	02/28/2008	11/30/2012						
18864-0211-03		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SERABRINA LA FRANCE 50 MG/15 ML	480	ML	NA	PO	ML	50	MG	0.06666	01/01/2002	99/99/9999						
20254-0018-01		Q0173		01/01/2002	09/11/2014	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 250 MG	100	EA	NA	PO	EA	250	MG	1	01/01/2002	09/11/2014						
20254-0018-03		Q0173		01/01/2002	09/11/2014	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 250 MG	500	EA	NA	PO	EA	250	MG	1	01/01/2002	09/11/2014						
20254-0207-06		Q0163		01/01/2002	09/11/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 25 MG	60	EA	NA	PO	EA	50	MG	0.5	01/01/2002	09/11/2014						
20254-0207-10		Q0163		01/01/2002	09/11/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 25 MG	10	EA	DP	PO	EA	50	MG	0.5	01/01/2002	09/11/2014						
20254-0208-06		Q0163		01/01/2002	09/11/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 50 MG	60	EA	NA	PO	EA	50	MG	1	01/01/2002	09/11/2014						
20254-0208-10		Q0163		01/01/2002	09/11/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (CAPLET) 50 MG	10	EA	NA	PO	EA	50	MG	1	01/01/2002	09/11/2014						
21695-0010-20		J8499		11/30/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	20	EA	BO	PO	EA	1	EA	1	11/30/2006	06/01/2014						
21695-0010-25		J8499		05/19/2008	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25	EA	BO	PO	EA	1	EA	1	05/19/2008	06/01/2014						
21695-0010-30		J8499		02/01/2007	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30	EA	BO	PO	EA	1	EA	1	02/01/2007	06/01/2014						
21695-0010-60		J8499		11/30/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	60	EA	BO	PO	EA	1	EA	1	11/30/2006	06/01/2014						
21695-0011-30		J8499		05/19/2008	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	05/19/2008	06/01/2014						
21695-0012-06		Q0144		07/19/2007	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6	EA	DP	PO	EA	1	GM	0.25	07/19/2007	06/01/2014						
21695-0080-21		J7509		01/01/2007	06/01/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISONE 4 MG	21	EA	BO	PO	EA	4	MG	1	01/01/2007	06/01/2014						
21695-0170-00		J7507		12/15/2006	06/01/2014	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	PROGRAF 1 MG	100	EA	BO	PO	EA	1	MG	1	12/15/2006	06/01/2014						
21695-0171-00		J7517		12/15/2006	06/01/2014	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT 250 MG	100	EA	BO	PO	EA	250	MG	1	12/15/2006	06/01/2014						
21695-0202-10		J0696		02/01/2007	06/01/2014	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (SDV) 500 MG	1	EA	VL	IJ	EA	250	MG	2	02/01/2007	06/01/2014						
21695-0241-01		J3070		01/01/2007	06/01/2014	INJECTION, PENTAZOCINE, 30 MG	TALWIN 30 MG/ML	1	ML	AM	IJ	ML	30	MG	1	01/01/2007	06/01/2014						
21695-0245-20		J7611		04/01/2008	06/01/2014	ALBUTEROL INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE 0.5%	20	ML	BO	IH	ML	1	MG	5	04/01/2008	06/01/2014						
21695-0304-30		Q0163		02/01/2007	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	30	EA	BO	PO	EA	50	MG	0.5	02/01/2007	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
21695-0304-90		Q0163		09/17/2007	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	90 EA	BO	PO	EA		50 MG		0.5	09/17/2007	06/01/2014						
21695-0306-20		J7506		04/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	04/01/2007	06/01/2014						
21695-0306-21		J7506		04/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	21 EA	BO	PO	EA		5 MG		2	04/01/2007	06/01/2014						
21695-0306-28		J7506		04/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	28 EA	BO	PO	EA		5 MG		2	04/01/2007	06/01/2014						
21695-0306-30		J7506		04/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30 EA	BO	PO	EA		5 MG		2	04/01/2007	06/01/2014						
21695-0306-42		J7506		04/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	42 EA	BO	PO	EA		5 MG		2	04/01/2007	06/01/2014						
21695-0307-10		J7506		02/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	10 EA	BO	PO	EA		5 MG		4	02/01/2007	06/01/2014						
21695-0307-18		J7506		04/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	18 EA	BO	PO	EA		5 MG		4	04/01/2007	06/01/2014						
21695-0307-20		J7506		07/27/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	20 EA	BO	PO	EA		5 MG		4	07/27/2007	06/01/2014						
21695-0307-30		J7506		02/01/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	30 EA	BO	PO	EA		5 MG		4	02/01/2007	06/01/2014						
21695-0332-25		J7613		04/01/2008	06/01/2014	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (3MLX25) 0.083%	3 ML	PC	IH	ML		1 MG		0.83	04/01/2008	06/01/2014						
21695-0332-25	KO	J7613	KO	04/01/2008	06/01/2014	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (3MLX25) 0.083%	3 ML	PC	IH	ML		1 MG		0.83	04/01/2008	06/01/2014						
21695-0365-08		J7510		10/15/2007	06/01/2014	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240 ML	BO	PO	ML		5 MG		0.6	10/15/2007	06/01/2014						
21695-0365-18		J7510		10/15/2007	06/01/2014	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	480 ML	BO	PO	ML		5 MG		0.6	10/15/2007	06/01/2014						
21695-0382-04		J8540		02/01/2007	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	4 EA	BO	PO	EA		0.25 MG		16	02/01/2007	06/01/2014						
21695-0414-60		Q0175		04/01/2007	06/01/2014	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	60 EA	BO	PO	EA		4 MG		1	04/01/2007	06/01/2014						
21695-0415-60		Q0176		06/27/2007	12/31/2013	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE (FILM-COATED) 8 MG	60 EA	BO	PO	EA		8 MG		1	06/27/2007	12/31/2013						
21695-0453-10		Q0170		04/01/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10 EA	BO	PO	EA		25 MG		1	04/01/2007	12/31/2013						
21695-0453-15		Q0170		01/15/2008	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	15 EA	BO	PO	EA		25 MG		1	01/15/2008	12/31/2013						
21695-0453-20		Q0170		04/01/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	20 EA	BO	PO	EA		25 MG		1	04/01/2007	12/31/2013						
21695-0453-25		Q0170		04/01/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	25 EA	BO	PO	EA		25 MG		1	04/01/2007	12/31/2013						
21695-0500-30		Q0163		04/15/2008	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30 EA	BO	PO	EA		50 MG		1	04/15/2008	06/01/2014						
21695-0572-30		Q0165		07/24/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (FILM-COATED) 10 MG	30 EA	BO	PO	EA		10 MG		1	07/24/2007	12/31/2013						
21695-0580-05		J7506		07/25/2007	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 50 MG	5 EA	BO	PO	EA		5 MG		10	07/25/2007	06/01/2014						
21695-0587-10		J2930		08/09/2007	06/01/2014	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	METHYLPREDNISOLONE 125 MG	1 EA	VL	IJ	EA		125 MG		1	08/09/2007	06/01/2014						
21695-0588-25		J1885		08/09/2007	06/01/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC (1MLX25) 30 MG/ML	1 ML	VL	IJ	ML		15 MG		2	08/09/2007	06/01/2014						
21695-0649-12		J8498		11/12/2007	06/01/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	12 EA	BX	RC	EA		1 EA		1	11/12/2007	06/01/2014						
21695-0703-04		Q0170		03/14/2008	12/31/2013	PROMETHAZINE HCL (1X120ML,FRUIT,TROPICAL) 6.25 MG/5 ML	PROMETHAZINE HCL (1X120ML,FRUIT,TROPICAL) 6.25 MG/5 ML	120 ML	BO	PO	ML		25 MG		0.05	03/14/2008	12/31/2013						
21695-0721-25		J1940		03/20/2008	06/01/2014	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (25X2ML) 10 MG/ML	2 ML	VL	IJ	ML		20 MG		0.5	03/20/2008	06/01/2014						
23490-1113-02		J7506		10/03/2006	01/01/2013	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	21 EA	NA	PO	EA		5 MG		2	10/03/2006	01/01/2013						
23490-1113-03		J7506		09/21/2006	01/01/2013	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30 EA	NA	PO	EA		5 MG		2	09/21/2006	01/01/2013						
23490-1911-02		J7509		10/03/2006	01/01/2013	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21 EA	NA	PO	EA		4 MG		1	10/03/2006	01/01/2013						
23490-5011-01		J8499		10/11/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG/5 ML	120 ML	BO	PO	ML		1 EA		1	10/11/2007	01/01/2013						
23490-5012-01		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25 EA	BO	PO	EA		1 EA		1	02/07/2007	01/01/2013						
23490-5012-02		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30 EA	BO	PO	EA		1 EA		1	02/07/2007	01/01/2013						
23490-5012-03		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40 EA	BO	PO	EA		1 EA		1	02/07/2007	01/01/2013						
23490-5012-04		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	100 EA	BO	PO	EA		1 EA		1	02/07/2007	01/01/2013						
23490-5013-01		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15 EA	BO	PO	EA		1 EA		1	02/07/2007	01/01/2013						
23490-5013-02		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	BO	PO	EA		1 EA		1	02/07/2007	01/01/2013						
23490-5013-03		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	40 EA	BO	PO	EA		1 EA		1	02/07/2007	01/01/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
23490-5013-04		J8499		10/11/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30 EA	BO PO EA	EA			1 EA		1	10/11/2007	01/01/2013						
23490-5015-01		J8499		02/07/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	25 EA	BO PO EA	EA			1 EA		1	02/07/2007	01/01/2013						
23490-5015-02		J8499		10/11/2007	01/01/2013	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30 EA	BO PO EA	EA			1 EA		1	10/11/2007	01/01/2013						
23490-5020-01		J7613		04/01/2008	01/01/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (24X3ML) 0.083%	3 ML	VL IH ML	ML			1 MG		0.83	04/01/2008	01/01/2013						
23490-5020-01	KO	J7613	KO	04/01/2008	01/01/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (24X3ML) 0.083%	3 ML	VL IH ML	ML			1 MG		0.83	04/01/2008	01/01/2013						
23490-5020-02		J7613		04/01/2008	01/01/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML) 0.083%	3 ML	PC IH ML	ML			1 MG		0.83	04/01/2008	01/01/2013						
23490-5020-02	KO	J7613	KO	04/01/2008	01/01/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML) 0.083%	3 ML	PC IH ML	ML			1 MG		0.83	04/01/2008	01/01/2013						
23490-5020-03		J7613		04/01/2008	01/01/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML) 0.083%	3 ML	PC IH ML	ML			1 MG		0.83	04/01/2008	01/01/2013						
23490-5020-03	KO	J7613	KO	04/01/2008	01/01/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML) 0.083%	3 ML	PC IH ML	ML			1 MG		0.83	04/01/2008	01/01/2013						
23490-5021-02		J7611		04/01/2008	01/01/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE (1X20ML) 0.5%	20 ML	BO IH ML	ML			1 MG		5	04/01/2008	01/01/2013						
23490-5110-09		J7500		04/30/2007	01/01/2013	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE 50 MG	90 EA	BO PO EA	EA			50 MG		1	04/30/2007	01/01/2013						
23490-5186-02		J0595		04/09/2007	01/01/2013	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE 2 MG/ML	10 ML	VL IJ ML	ML			1 MG		2	04/09/2007	01/01/2013						
23490-5404-01		J8540		02/07/2007	01/01/2013	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	12 EA	BO PO EA	EA			0.25 MG		3	02/07/2007	01/01/2013						
23490-5407-01		J8540		02/07/2007	01/01/2013	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	6 EA	BO PO EA	EA			0.25 MG		16	02/07/2007	01/01/2013						
23490-5407-02		J8540		11/30/2007	01/01/2013	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	12 EA	BO PO EA	EA			0.25 MG		16	11/30/2007	01/01/2013						
23490-5413-00		J1100		04/09/2007	01/01/2013	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE 4 MG/ML	5 ML	VL IJ ML	ML			1 MG		4	04/09/2007	01/01/2013						
23490-5455-01		Q0163		11/30/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE (1X120ML) 12.5 MG/5 ML	120 ML	BO PO ML	ML			50 MG		0.05	11/30/2007	01/01/2013						
23490-5457-00		Q0163		11/30/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	24 EA	BO PO EA	EA			50 MG		0.5	11/30/2007	01/01/2013						
23490-5457-01		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	6 EA	BO PO EA	EA			50 MG		0.5	02/07/2007	01/01/2013						
23490-5457-02		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	15 EA	BO PO EA	EA			50 MG		0.5	02/07/2007	01/01/2013						
23490-5457-03		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	20 EA	BO PO EA	EA			50 MG		0.5	02/07/2007	01/01/2013						
23490-5457-04		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	30 EA	BO PO EA	EA			50 MG		0.5	02/07/2007	01/01/2013						
23490-5457-05		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	60 EA	BO PO EA	EA			50 MG		0.5	02/07/2007	01/01/2013						
23490-5459-01		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 50 MG	6 EA	BO PO EA	EA			50 MG		1	02/07/2007	01/01/2013						
23490-5459-02		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 50 MG	15 EA	BO PO EA	EA			50 MG		1	02/07/2007	01/01/2013						
23490-5459-03		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 50 MG	30 EA	BO PO EA	EA			50 MG		1	02/07/2007	01/01/2013						
23490-5459-04		Q0163		02/07/2007	01/01/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 50 MG	60 EA	BO PO EA	EA			50 MG		1	02/07/2007	01/01/2013						
23490-5621-02		J1940		04/30/2007	01/01/2013	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE 10 MG/ML	2 ML	VL IJ ML	ML			20 MG		0.5	04/30/2007	01/01/2013						
23490-5733-01		Q0177		02/07/2007	01/01/2013	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20 EA	BO PO EA	EA			25 MG		1	02/07/2007	01/01/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
23490-5733-02		Q0177		02/07/2007	01/01/2013	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30 EA	BO	PO	EA		25 MG		1	02/07/2007	01/01/2013						
23490-5761-01		J7644		04/09/2007	01/01/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML) 0.02%	2.5 ML	PC	IH	ML		1 MG		0.2	04/09/2007	01/01/2013						
23490-5761-01	KO	J7644	KO	04/09/2007	01/01/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML) 0.02%	2.5 ML	PC	IH	ML		1 MG		0.2	04/09/2007	01/01/2013						
23490-5792-04		J1885		04/09/2007	01/01/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/ML	1 ML	NA	IJ	ML		15 MG		2	04/09/2007	01/01/2013						
23490-5854-01		J1055		02/07/2007	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	MEDROXYPROGESTERONE ACETATE 150 MG/ML	1 ML	VL	IM	ML		150 MG		1	02/07/2007	12/31/2012						
23490-5889-00		None		11/30/2007	01/01/2013	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	24 EA	BO	PO	EA		2.5 MG		1	11/30/2007	01/01/2013						
23490-5902-01		J7509		02/07/2007	01/01/2013	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21 EA	BO	PO	EA		4 MG		1	02/07/2007	01/01/2013						
23490-5914-01		J2765		04/09/2007	01/01/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HYDROCHLORIDE 5 MG/ML	2 ML	VL	IV	ML		10 MG		0.5	04/09/2007	01/01/2013						
23490-5932-01		J2250		04/30/2007	01/01/2013	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HYDROCHLORIDE 1 MG/ML	2 ML	VL	IJ	ML		1 MG		1	04/30/2007	01/01/2013						
23490-5933-01		J2250		04/30/2007	01/01/2013	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HYDROCHLORIDE 5 MG/ML	2 ML	VL	IJ	ML		1 MG		5	04/30/2007	01/01/2013						
23490-5933-02		J2250		04/30/2007	01/01/2013	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HYDROCHLORIDE (10X10ML) 5 MG/ML	10 ML	VL	IJ	ML		1 MG		5	04/30/2007	01/01/2013						
23490-5955-01		J2300		04/09/2007	01/01/2013	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HYDROCHLORIDE 10 MG/ML	10 ML	VL	IJ	ML		10 MG		1	04/09/2007	01/01/2013						
23490-6144-01		J7510		04/09/2007	01/01/2013	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240 ML	BO	PO	ML		5 MG		0.6	04/09/2007	01/01/2013						
23490-6144-02		J7510		10/11/2007	01/01/2013	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	180 ML	BO	PO	ML		5 MG		0.6	10/11/2007	01/01/2013						
23490-6144-03		J7510		10/11/2007	01/01/2013	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	120 ML	BO	PO	ML		5 MG		0.6	10/11/2007	01/01/2013						
23490-6145-01		J7510		10/11/2007	01/01/2013	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240 ML	BO	PO	ML		5 MG		0.6	10/11/2007	01/01/2013						
23490-6145-02		J7510		10/11/2007	01/01/2013	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	180 ML	BO	PO	ML		5 MG		0.6	10/11/2007	01/01/2013						
23490-6145-03		J7510		10/11/2007	01/01/2013	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	120 ML	BO	PO	ML		5 MG		0.6	10/11/2007	01/01/2013						
23490-6157-01		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	10 EA	BO	PO	EA		5 MG		2	02/07/2007	01/01/2013						
23490-6157-02		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	02/07/2007	01/01/2013						
23490-6157-03		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	40 EA	BO	PO	EA		5 MG		2	02/07/2007	01/01/2013						
23490-6157-04		J7506		04/09/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	37 EA	BO	PO	EA		5 MG		2	04/09/2007	01/01/2013						
23490-6157-05		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21 EA	BO	PO	EA		5 MG		2	02/07/2007	01/01/2013						
23490-6157-06		J7506		11/30/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	30 EA	BO	PO	EA		5 MG		2	11/30/2007	01/01/2013						
23490-6157-07		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	60 EA	BO	PO	EA		5 MG		2	02/07/2007	01/01/2013						
23490-6157-08		J7506		04/09/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100 EA	BO	PO	EA		5 MG		2	04/09/2007	01/01/2013						
23490-6158-00		J7506		04/09/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	6 EA	BO	PO	EA		5 MG		4	04/09/2007	01/01/2013						
23490-6158-01		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	10 EA	BO	PO	EA		5 MG		4	02/07/2007	01/01/2013						
23490-6158-02		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	18 EA	BO	PO	EA		5 MG		4	02/07/2007	01/01/2013						
23490-6158-03		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	20 EA	BO	PO	EA		5 MG		4	02/07/2007	01/01/2013						
23490-6158-04		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30 EA	BO	PO	EA		5 MG		4	02/07/2007	01/01/2013						
23490-6158-05		J7506		10/11/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	21 EA	BO	PO	EA		5 MG		4	10/11/2007	01/01/2013						
23490-6158-07		J7506		04/09/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	25 EA	BO	PO	EA		5 MG		4	04/09/2007	01/01/2013						
23490-6158-08		J7506		04/09/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	15 EA	BO	PO	EA		5 MG		4	04/09/2007	01/01/2013						
23490-6158-09		J7506		10/11/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	90 EA	BO	PO	EA		5 MG		4	10/11/2007	01/01/2013						
23490-6159-01		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	10 EA	BO	PO	EA		5 MG		1	02/07/2007	01/01/2013						
23490-6159-02		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	20 EA	BO	PO	EA		5 MG		1	02/07/2007	01/01/2013						
23490-6159-03		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	21 EA	BO	PO	EA		5 MG		1	02/07/2007	01/01/2013						
23490-6159-04		J7506		02/07/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	40 EA	BO	PO	EA		5 MG		1	02/07/2007	01/01/2013						
23490-6159-05		J7506		11/30/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	28 EA	BO	PO	EA		5 MG		1	11/30/2007	01/01/2013						
23490-6159-06		J7506		11/30/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	30 EA	BO	PO	EA		5 MG		1	11/30/2007	01/01/2013						
23490-6174-01		J8498		02/07/2007	01/01/2013	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	3 EA	BX	RC	EA		1 EA		1	02/07/2007	01/01/2013						
23490-6180-01		J8498		02/07/2007	01/01/2013	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 12.5 MG	12 EA	BX	RC	EA		1 EA		1	02/07/2007	01/01/2013						
23490-6182-01		J8498		02/07/2007	01/01/2013	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 25 MG	6 EA	BX	RC	EA		1 EA		1	02/07/2007	01/01/2013						
23490-6182-02		J8498		02/07/2007	01/01/2013	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 25 MG	12 EA	BX	RC	EA		1 EA		1	02/07/2007	01/01/2013						
23490-6182-03		J8498		11/30/2007	01/01/2013	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 25 MG	10 EA	BX	RC	EA		1 EA		1	11/30/2007	01/01/2013						
23490-6183-01		Q0170		02/07/2007	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	10 EA	BO	PO	EA		25 MG		1	02/07/2007	01/01/2013						
23490-6183-02		Q0170		02/07/2007	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	12 EA	BO	PO	EA		25 MG		1	02/07/2007	01/01/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
23490-6183-03		Q0170		02/07/2007	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	30	EA	BO	PO	EA	25	MG	1	02/07/2007	01/01/2013						
23490-6183-04		Q0170		04/09/2007	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	20	EA	BO	PO	EA	25	MG	1	04/09/2007	01/01/2013						
23490-6183-06		Q0170		11/30/2007	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60	EA	BO	PO	EA	25	MG	1	11/30/2007	01/01/2013						
23490-6183-07		Q0170		03/12/2008	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	90	EA	BO	PO	EA	25	MG	1	03/12/2008	01/01/2013						
23490-6183-08		Q0170		03/12/2008	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	100	EA	BO	PO	EA	25	MG	1	03/12/2008	01/01/2013						
23490-6187-01		Q0170		11/30/2007	01/01/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (1X120ML) 6.25 MG/5 ML	120	ML	BO	PO	ML	25	MG	0.05	11/30/2007	01/01/2013						
23490-6343-01		J1080		02/07/2007	01/01/2013	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TESTOSTERONE CYPIONATE 200 MG/ML	10	ML	NA	IM	ML	200	MG	1	02/07/2007	01/01/2013						
23490-6509-03		Q0165		11/30/2007	01/01/2013	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	10	MG	1	11/30/2007	01/01/2013						
23490-6512-01		Q0164		02/07/2007	01/01/2013	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	6	EA	BO	PO	EA	5	MG	1	02/07/2007	01/01/2013						
23490-6512-02		Q0164		02/07/2007	01/01/2013	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	10	EA	BO	PO	EA	5	MG	1	02/07/2007	01/01/2013						
23490-6687-00		J1815		04/30/2007	01/01/2013	INJECTION, INSULIN, PER 5 UNITS	INSULIN HUMAN REGULAR 100 U/ML	10	ML	NA	IJ	ML	5	U	20	04/30/2007	01/01/2013						
23490-6904-01		Q0144		11/12/2007	01/01/2013	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (1X15ML) 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	11/12/2007	01/01/2013						
23490-6905-00		Q0144		04/09/2007	01/01/2013	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN DIHYDRATE 200 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.04	04/09/2007	01/01/2013						
23490-6905-01		Q0144		10/11/2007	01/01/2013	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN DIHYDRATE 200 MG/5 ML	22.5	ML	BO	PO	ML	1	GM	0.04	10/11/2007	01/01/2013						
23490-6905-02		Q0144		10/11/2007	01/01/2013	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN DIHYDRATE 200 MG/5 ML	30	ML	BO	PO	ML	1	GM	0.04	10/11/2007	01/01/2013						
23490-7545-02		J3360		04/09/2007	01/01/2013	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM 5 MG/ML	10	ML	NA	IJ	ML	5	MG	1	04/09/2007	01/01/2013						
23490-7758-01		Q0144		02/07/2007	01/01/2013	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	3	EA	DP	PO	EA	1	GM	0.5	02/07/2007	01/01/2013						
23490-7760-01		Q0144		02/07/2007	01/01/2013	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	4	EA	BO	PO	EA	1	GM	0.25	02/07/2007	01/01/2013						
23490-7760-02		Q0144		04/09/2007	01/01/2013	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6	EA	BO	PO	EA	1	GM	0.25	04/09/2007	01/01/2013						
23490-7854-00		J7506		11/30/2007	01/01/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE (1X120ML) 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	11/30/2007	01/01/2013						
23535-0608-61		J3475		01/01/2002	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE	1	EA	NA	NA	GM	500	MG	2	01/01/2002	99/99/9999						
23535-0608-68		J3475		01/01/2002	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE	1	EA	NA	NA	GM	500	MG	2	01/01/2002	99/99/9999						
24208-0347-20		J7611		04/01/2008	06/05/2017	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE (STERILE) 0.5%	20	ML	BO	IH	ML	1	MG	5	04/01/2008	06/05/2017						
24385-0379-26		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHEDRYL (CHERRY) 12.5 MG/5 ML	120	ML	BO	PO	ML	50	MG	0.05	01/01/2002	99/99/9999						
24385-0406-73		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SLEEP TABLETS 25 MG	16	EA	NA	PO	EA	50	MG	0.5	01/01/2002	02/03/2016						
24385-0462-62		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHEDRYL 25 MG	24	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
24385-0462-78		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHEDRYL 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
24385-0479-62		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHEDRYL 25 MG	24	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
24385-0479-78		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHEDRYL 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
25208-0002-01		J3246		04/01/2008	12/31/2017	INJECTION, TIROFIBAN HCL, 0.25MG	AGGRASTAT (1X100ML) 0.05 MG/ML	100 ML	PC	IV	ML		0.25 MG		0.2	04/01/2008	12/31/2017						
25208-0002-02		J3246		04/01/2008	99/99/9999	INJECTION, TIROFIBAN HCL, 0.25MG	AGGRASTAT (1X250ML) 0.05 MG/ML	250 ML	PC	IV	ML		0.25 MG		0.2	04/01/2008	99/99/9999						
25332-0004-30		J3420		01/01/2002	01/06/2017	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	COBOLIN-M (VIAL) 1000 MCG/ML	30 ML	VL	IM	ML		1000 MCG		1	01/01/2002	01/06/2017						
25332-0073-30		J3415		01/01/2004	02/03/2016	INJECTION, PYRIDOXINE HCL, 100 MG	RODEX (VIAL) 100 MG/ML	30 ML	VL	IJ	ML		100 MG		1	01/01/2004	02/03/2016						
25332-0078-10		J3420		01/01/2002	01/06/2017	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	DEPO-COBOLIN (VIAL) 1000 MCG/ML	30 ML	VL	IM	ML		1000 MCG		1	01/01/2002	01/06/2017						
25332-0088-05		J3490		01/01/2002	08/06/2013	UNCLASSIFIED DRUGS	PRODRUX (VIAL) 250 MG/ML	5 ML	VL	IM	ML		1 EA		1	01/01/2002	08/06/2013						
25682-0001-01		J1300		01/01/2008	99/99/9999	INJECTION, EUCULIZUMAB, 10 MG	SOLIRIS (PF) 10 MG/ML	30 ML	VL	IV	ML		10 MG		1	01/01/2008	99/99/9999						
30103-0322-54		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DORMIN SLEEP AID 25 MG	32 EA	NA	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
30103-0722-54		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DORMIN SLEEP AID 25 MG	72 EA	NA	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
33261-0335-21		J7509		01/15/2008	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (DOSE PACK) 4 MG	21 EA	NA	PO	EA		4 MG		1	01/15/2008	99/99/9999						
33358-0009-25		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0010-15		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0010-28		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	28 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0010-30		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0010-60		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	60 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0011-25		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	25 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0011-30		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0011-35		J8499		07/10/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	35 EA	BO	PO	EA		1 EA		1	07/10/2007	99/99/9999						
33358-0040-06		Q0144		07/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6 EA	BO	PO	EA		1 GM		0.25	07/10/2007	99/99/9999						
33358-0041-10		Q0144		07/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	10 EA	BO	PO	EA		1 GM		0.5	07/10/2007	99/99/9999						
33358-0110-30		Q0163		07/10/2007	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	30 EA	BO	PO	EA		50 MG		0.5	07/10/2007	99/99/9999						
33358-0111-20		Q0163		07/10/2007	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 50 MG	20 EA	BO	PO	EA		50 MG		1	07/10/2007	99/99/9999						
33358-0111-30		Q0163		07/10/2007	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 50 MG	30 EA	BO	PO	EA		50 MG		1	07/10/2007	99/99/9999						
33358-0182-20		Q0177		07/10/2007	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAM 25 MG	20 EA	BO	PO	EA		25 MG		1	07/10/2007	99/99/9999						
33358-0182-30		Q0177		07/10/2007	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAM 25 MG	30 EA	BO	PO	EA		25 MG		1	07/10/2007	99/99/9999						
33358-0241-21		J7509		07/10/2007	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21 EA	BO	PO	EA		4 MG		1	07/10/2007	99/99/9999						
33358-0291-08		J7510		07/10/2007	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240 ML	BO	PO	ML		5 MG		0.6	07/10/2007	99/99/9999						
33358-0292-12		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	12 EA	BO	PO	EA		5 MG		1	07/10/2007	12/31/2015						
33358-0292-15		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	15 EA	BO	PO	EA		5 MG		1	07/10/2007	12/31/2015						
33358-0292-21		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	21 EA	BO	PO	EA		5 MG		1	07/10/2007	12/31/2015						
33358-0292-30		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	30 EA	BO	PO	EA		5 MG		1	07/10/2007	12/31/2015						
33358-0292-78		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	78 EA	BO	PO	EA		5 MG		1	07/10/2007	12/31/2015						
33358-0293-20		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	07/10/2007	12/31/2015						
33358-0293-30		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30 EA	BO	PO	EA		5 MG		2	07/10/2007	12/31/2015						
33358-0293-40		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	40 EA	BO	PO	EA		5 MG		2	07/10/2007	12/31/2015						
33358-0294-15		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	15 EA	BO	PO	EA		5 MG		4	07/10/2007	12/31/2015						
33358-0294-20		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	20 EA	BO	PO	EA		5 MG		4	07/10/2007	12/31/2015						
33358-0294-30		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	30 EA	BO	PO	EA		5 MG		4	07/10/2007	12/31/2015						
33358-0294-40		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	40 EA	BO	PO	EA		5 MG		4	07/10/2007	12/31/2015						
33358-0294-60		J7506		07/10/2007	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	60 EA	BO	PO	EA		5 MG		4	07/10/2007	12/31/2015						
33358-0299-20		Q0164		07/10/2007	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 5 MG	20 EA	BO	PO	EA		5 MG		1	07/10/2007	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
33358-0299-30		Q0164		07/10/2007	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 5 MG	30	EA	BO	PO	EA	5	MG	1	07/10/2007	99/99/9999						
33358-0300-10		Q0165		07/10/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	10	EA	BO	PO	EA	10	MG	1	07/10/2007	12/31/2013						
33358-0300-20		Q0165		07/10/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	20	EA	BO	PO	EA	10	MG	1	07/10/2007	12/31/2013						
33358-0300-30		Q0165		07/10/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	30	EA	BO	PO	EA	10	MG	1	07/10/2007	12/31/2013						
33358-0300-60		Q0165		07/10/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	60	EA	BO	PO	EA	10	MG	1	07/10/2007	12/31/2013						
33358-0301-02		J8498		07/10/2007	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	2	EA	BX	RC	EA	1	EA	1	07/10/2007	99/99/9999						
33358-0301-12		J8498		07/10/2007	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12	EA	BX	RC	EA	1	EA	1	07/10/2007	99/99/9999						
33358-0302-08		Q0170		07/10/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	8	EA	BO	PO	EA	25	MG	1	07/10/2007	12/31/2013						
33358-0302-10		Q0170		07/10/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10	EA	BO	PO	EA	25	MG	1	07/10/2007	12/31/2013						
33358-0302-30		Q0170		07/10/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	30	EA	BO	PO	EA	25	MG	1	07/10/2007	12/31/2013						
33358-0302-60		Q0170		07/10/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	60	EA	BO	PO	EA	25	MG	1	07/10/2007	12/31/2013						
33358-0313-01		J3415		07/10/2007	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE (SINGLE-DOSE) 100 MG/ML	1	ML	VL	U	ML	100	MG	1	07/10/2007	99/99/9999						
33358-0352-10		Q0173		07/10/2007	02/03/2016	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE 250 MG	10	EA	NA	PO	EA	250	MG	1	07/10/2007	02/03/2016						
33358-0352-20		Q0173		07/10/2007	02/03/2016	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE 250 MG	20	EA	NA	PO	EA	250	MG	1	07/10/2007	02/03/2016						
33358-0367-01		Q0144		07/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 1 GM/Packet	1	EA	BX	PO	EA	1	GM	1	07/10/2007	99/99/9999						
33358-0367-03		Q0144		07/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 1 GM/Packet	1	EA	BX	PO	EA	1	GM	1	07/10/2007	99/99/9999						
33358-0368-04		Q0144		07/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	4	EA	BO	PO	EA	1	GM	0.25	07/10/2007	99/99/9999						
33358-0368-30		Q0144		07/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	30	EA	BO	PO	EA	1	GM	0.25	07/10/2007	99/99/9999						
33358-0368-50		Q0144		07/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	50	EA	BO	PO	EA	1	GM	0.25	07/10/2007	99/99/9999						
33358-0418-30		Q0169		07/24/2007	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 12.5 MG	30	EA	BO	PO	EA	12.5	MG	1	07/24/2007	99/99/9999						
35356-0017-03		Q0144		09/14/2007	01/01/2015	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	3	EA	BO	PO	EA	1	GM	0.5	09/14/2007	01/01/2015						
35356-0019-10		J1650		09/14/2007	02/03/2016	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (100X0.6ML) 60 MG/0.6 ML	0.6	ML	SR	SC	ML	10	MG	10	09/14/2007	02/03/2016						
35356-0020-10		J1650		09/14/2007	02/03/2016	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (100X0.8ML) 80 MG/0.8 ML	0.8	ML	SR	SC	ML	10	MG	10	09/14/2007	02/03/2016						
35356-0038-12		J8498		10/19/2007	01/01/2015	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENADOZ 25 MG	12	EA	BX	RC	EA	1	EA	1	10/19/2007	01/01/2015						
35356-0044-15		Q0144		10/26/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	10/26/2007	99/99/9999						
35356-0058-10		J1070		11/09/2007	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	DEPO-TESTOSTERONE 100 MG/ML	10	ML	VL	IM	ML	100	MG	1	11/09/2007	12/31/2014						
35356-0082-01		J3301		02/08/2008	01/01/2015	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG 10 MG/ML	5	ML	VL	IJ	ML	10	MG	1	02/08/2008	01/01/2015						
35356-0083-01		J1030		02/08/2008	01/01/2015	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE 40 MG/ML	5	ML	VL	IJ	ML	40	MG	1	02/08/2008	01/01/2015						
35356-0084-01		J0702		02/08/2008	01/01/2015	INJECTION, BETAMETHASONE ACETATE 3MG AND BETAMETHASONE SODIUM PHOSPHATE 3MG	CELESTONE SOLUSPAN 3 MG/ML-3	5	ML	VL	IJ	ML	3	MG	1	02/08/2008	01/01/2015						
35356-0096-60		Q0176		02/29/2008	12/31/2013	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	60	EA	BO	PO	EA	8	MG	1	02/29/2008	12/31/2013						
35356-0098-90		Q0172		02/29/2008	12/31/2013	CHLORPROMAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE 100 MG	90	EA	BO	PO	EA	25	MG	4	02/29/2008	12/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
35356-0102-00		J1817		03/07/2008	01/01/2015	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	HUMALOG (100X10ML) 100 U/ML	10	ML	VL	SC	ML	50 U		2	03/07/2008	01/01/2015							
35356-0124-30		J7644		03/13/2008	01/01/2015	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1 MG		0.2	03/13/2008	01/01/2015							
35356-0124-30	KO	J7644	KO	03/13/2008	01/01/2015	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1 MG		0.2	03/13/2008	01/01/2015							
35356-0128-15		Q0144		03/13/2008	01/01/2015	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 100 MG/5 ML	15	ML	BO	PO	ML	1 GM		0.02	03/13/2008	01/01/2015							
35356-0177-15		J0696		05/16/2008	01/01/2015	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (1X15ML) 1 GM	15	ML	NA	IJ	ML	250 MG		4	05/16/2008	01/01/2015							
35356-0178-05		J1040		05/16/2008	01/01/2015	INJECTION, METHYLPREDNISOLONE ACETATE, 80 MG	METHYLPREDNISOLONE ACETATE (1X5ML) 80 MG/ML	5	ML	NA	IJ	ML	80 MG		1	05/16/2008	01/01/2015							
35356-0180-50		J2001		05/16/2008	01/01/2015	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (1X50ML,LATEX-FREE) 2%	50	ML	NA	IJ	ML	10 MG		2	05/16/2008	01/01/2015							
35356-0181-30		A4216		05/16/2008	01/01/2015	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE BACTERIOSTATIC (1X30ML,LATEX-FREE) 0.9%	30	ML	NA	IV	ML	10 ML		0.1	05/16/2008	01/01/2015							
35356-0194-21		J7509		05/16/2008	01/01/2015	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL (DOSE PACK) 4 MG	21	EA	NA	PO	EA	4 MG		1	05/16/2008	01/01/2015							
37205-0270-62		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPLETE ALLERGY MEDICINE 25 MG	24	EA	BX	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
37205-0270-78		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPLETE ALLERGY MEDICINE 25 MG	100	EA	BX	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
37205-0277-62		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPLETE ALLERGY MEDICINE 25 MG	24	EA	BX	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
37205-0277-78		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPLETE ALLERGY MEDICINE 25 MG	100	EA	BX	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
37205-0565-26		Q0163		01/01/2002	09/19/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPLETE ALLERGY (AF,CHERRY) 12.5 MG/5 ML	118	ML	BO	PO	ML	50 MG		0.05	01/01/2002	09/19/2017							
37205-0565-34		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPLETE ALLERGY (AF,CHERRY) 12.5 MG/5 ML	240	ML	BO	PO	ML	50 MG		0.05	01/01/2002	99/99/9999							
38423-0110-01		J1190		09/06/2007	04/21/2016	INJECTION, DEXRAXOXANE HYDROCHLORIDE, PER 250 MG	TOTECT (W/10 VIALS OF DILUENT) 500 MG	1	EA	VL	IV	EA	250 MG		2	09/06/2007	04/21/2016							
38779-0006-03		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1 EA		1	01/01/2002	99/99/9999							
38779-0006-04		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1 EA		1	01/01/2002	99/99/9999							
38779-0006-05		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1 EA		1	01/01/2002	99/99/9999							
38779-0008-01		J1700		01/01/2002	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	25 MG		40	01/01/2002	99/99/9999							
38779-0008-04		J1700		01/01/2002	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	25 MG		40	01/01/2002	99/99/9999							
38779-0008-05		J1700		01/01/2002	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	25 MG		40	01/01/2002	99/99/9999							
38779-0008-08		J1700		01/01/2002	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	25 MG		40	01/01/2002	99/99/9999							
38779-0008-09		J1700		01/01/2002	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	25 MG		40	01/01/2002	99/99/9999							
38779-0011-01		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0011-01	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0011-03		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0011-03	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0011-04		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0011-04	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0011-05		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0011-05	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 MG		1000	01/01/2002	99/99/9999							
38779-0015-01		J3490		04/26/2002	99/99/9999	UNCLASSIFIED DRUGS	BACITRACIN (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 EA		1	04/26/2002	99/99/9999							
38779-0015-04		J3490		04/26/2002	99/99/9999	UNCLASSIFIED DRUGS	BACITRACIN (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 EA		1	04/26/2002	99/99/9999							
38779-0015-05		J3490		04/26/2002	99/99/9999	UNCLASSIFIED DRUGS	BACITRACIN (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1 EA		1	04/26/2002	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
38779-0126-01		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
38779-0126-03		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
38779-0126-04		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
38779-0126-06		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
38779-0142-04		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		4 MG		250	01/01/2002	99/99/9999						
38779-0142-06		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		4 MG		250	01/01/2002	99/99/9999						
38779-0144-03		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		40 MG		25	01/01/2002	99/99/9999						
38779-0144-04		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		40 MG		25	01/01/2002	99/99/9999						
38779-0144-05		J1030		09/03/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		40 MG		25	09/03/2002	99/99/9999						
38779-0144-06		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		40 MG		25	01/01/2002	99/99/9999						
38779-0146-04		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
38779-0146-05		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
38779-0146-08		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
38779-0146-09		J3490		09/03/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	09/03/2002	99/99/9999						
38779-0150-03		J7510		01/01/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	99/99/9999						
38779-0150-04		J7510		01/01/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	99/99/9999						
38779-0150-05		J7510		01/01/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	99/99/9999						
38779-0150-08		J7510		04/25/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (ANHYDROUS,MICRONIZED)	1 EA	NA	NA	GM		5 MG		200	04/25/2002	99/99/9999						
38779-0150-09		J7510		09/03/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	09/03/2002	99/99/9999						
38779-0154-03		J7506		03/07/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	03/07/2002	12/31/2015						
38779-0154-04		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	12/31/2015						
38779-0154-05		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	12/31/2015						
38779-0154-08		J7506		08/26/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	08/26/2002	12/31/2015						
38779-0154-09		J7506		08/26/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		5 MG		200	08/26/2002	12/31/2015						
38779-0164-03		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0164-04		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0164-05		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0164-08		J1070		04/30/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	04/30/2002	12/31/2014						
38779-0164-09		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	JR	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0165-03		J3150		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (USP,MICRONIZED)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0165-04		J3150		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (USP,MICRONIZED)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0165-05		J3150		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0165-08		J3150		04/30/2002	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		100 MG		10	04/30/2002	12/31/2014						
38779-0166-03		J3302		01/01/2002	99/99/9999	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (USP)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	99/99/9999						
38779-0166-04		J3302		01/01/2002	99/99/9999	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (USP)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	99/99/9999						
38779-0166-05		J3302		01/01/2002	99/99/9999	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (USP)	1 EA	BO	NA	GM		5 MG		200	01/01/2002	99/99/9999						
38779-0173-01		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM		5 MG		200	01/01/2006	99/99/9999						
38779-0173-04		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM		5 MG		200	01/01/2006	99/99/9999						
38779-0173-05		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM		5 MG		200	01/01/2006	99/99/9999						
38779-0173-08		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM		5 MG		200	01/01/2006	99/99/9999						
38779-0180-04		Q0165		03/08/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	1 EA	BO	NA	GM		10 MG		100	03/08/2002	12/31/2013						
38779-0180-05		Q0165		01/01/2002	12/31/2013	DOSAGE REGIMEN PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR	PROCHLORPERAZINE MALEATE (U.S.P.)	1 EA	BO	NA	GM		10 MG		100	01/01/2002	12/31/2013						
38779-0180-08		Q0165		01/01/2002	12/31/2013	DOSAGE REGIMEN PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR	PROCHLORPERAZINE MALEATE (U.S.P.)	1 EA	BO	NA	GM		10 MG		100	01/01/2002	12/31/2013						
38779-0183-03		J1800		01/01/2002	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
38779-0183-04		J1800		01/01/2002	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0183-05		J1800		01/01/2002	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0183-08		J1800		01/01/2002	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0185-04		J7609		01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2007	99/99/9999						
38779-0185-04	KO	J7609	KO	01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2007	99/99/9999						
38779-0185-05		J7609		01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2007	99/99/9999						
38779-0185-05	KO	J7609	KO	01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2007	99/99/9999						
63323-0517-74		J1644		06/15/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM-SODIUM CHLORIDE (FREEFLEX BAG,LATEX-FREE) 25000 U/250 ML-0.45%	250 ML	BG IV ML		ML		1000 U		0.1	06/15/2018	99/99/9999						
63323-0518-77		J1644		06/15/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM-SODIUM CHLORIDE (FREEFLEX BAG,LATEX-FREE) 25000 U/500 ML-0.45%	500 ML	BG IV ML		ML		1000 U		0.05	06/15/2018	99/99/9999						
63323-0522-77		J1644		06/15/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM-DEXTROSE (FREEFLEX BAG,LATEX-FREE) 5%-25000 U/500 ML	500 ML	BG IV ML		ML		1000 U		0.05	06/15/2018	99/99/9999						
38779-0191-03		J0285		01/01/2002	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO NA GM	50 MG			20			01/01/2002	99/99/9999						
38779-0191-04		J0285		01/01/2002	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO NA GM	50 MG			20			01/01/2002	99/99/9999						
38779-0191-05		J0285		01/01/2002	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO NA GM	50 MG			20			01/01/2002	99/99/9999						
38779-0191-06		J0285		11/27/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO NA GM	50 MG			20			11/27/2003	99/99/9999						
38779-0191-08		J0285		01/01/2002	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	JR NA GM	50 MG			20			01/01/2002	99/99/9999						
63323-0523-74		J1644		06/15/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM-DEXTROSE (FREEFLEX BAG,LATEX-FREE) 5%-25000 U/250 ML	250 ML	BG IV ML		ML		1000 U		0.1	06/15/2018	99/99/9999						
38779-0194-03		J0515		01/01/2002	10/17/2016	INJECTION, BENZTROPINE MESYLATE, PER 1 MG	BENZTROPINE MESYLATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	10/17/2016						
38779-0195-01		J7624		01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0195-01	KO	J7624	KO	01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0195-03		J7624		01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0195-03	KO	J7624	KO	01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0195-06		J7624		01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0195-06	KO	J7624	KO	01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO NA GM	1 MG			1000			01/01/2002	99/99/9999						
38779-0198-00		J7627		01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0198-00	KO	J7627	KO	01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0198-03		J7627		01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0198-03	KO	J7627	KO	01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0198-04		J7626		04/19/2002	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			09/26/2008	99/99/9999	04/19/2002	04/25/2002	2000			
38779-0198-04	KO	J7626	KO	04/19/2002	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			09/26/2008	99/99/9999	04/19/2002	04/25/2002	2000			
38779-0198-05		J7627		01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED,MICRONIZED)	1 EA	NA NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0198-05	KO	J7627	KO	01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED,MICRONIZED)	1 EA	NA NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0198-06		J7627		01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0198-06	KO	J7627	KO	01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	BO NA GM	0.5 MG			2000			01/01/2006	99/99/9999						
38779-0215-00		J1160		02/05/2002	10/17/2016	INJECTION, DIGOXIN, UP TO 0.5 MG	DIGOXIN (U.S.P.)	1 EA	BO NA GM	0.5 MG			2000			02/05/2002	10/17/2016						
38779-0215-06		J1160		02/05/2002	10/17/2016	INJECTION, DIGOXIN, UP TO 0.5 MG	DIGOXIN (U.S.P.)	1 EA	BO NA GM	0.5 MG			2000			02/05/2002	10/17/2016						
38779-0215-09		J1160		02/05/2002	99/99/9999	INJECTION, DIGOXIN, UP TO 0.5 MG	DIGOXIN (U.S.P.)	1 EA	BO NA GM	0.5 MG			2000			02/05/2002	99/99/9999						
38779-0216-04		J1165		01/01/2002	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	BO NA GM	50 MG			20			01/01/2002	99/99/9999						
38779-0216-05		J1165		01/01/2002	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	BO NA GM	50 MG			20			01/01/2002	99/99/9999						
38779-0216-08		J1165		01/01/2002	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	BO NA GM	50 MG			20			01/01/2002	99/99/9999						
38779-0230-03		J7645		01/01/2007	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1 EA	BO NA GM	1 MG			1000			01/01/2007	99/99/9999						
38779-0230-03	KO	J7645	KO	01/01/2007	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1 EA	BO NA GM	1 MG			1000			01/01/2007	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
38779-0319-06		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (U.S.P.)	1 EA	BO	NA	GM	300 MG	3.33333		01/01/2007	99/99/9999							
38779-0319-06	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (U.S.P.)	1 EA	BO	NA	GM	300 MG	3.33333		01/01/2007	99/99/9999							
38779-0324-03		J1730		01/01/2002	99/99/9999	INJECTION, DIAZOXIDE, UP TO 300 MG	DIAZOXIDE (U.S.P.)	1 EA	BO	NA	GM	300 MG	3.33333		01/01/2002	99/99/9999							
38779-0324-04		J1730		01/01/2002	99/99/9999	INJECTION, DIAZOXIDE, UP TO 300 MG	DIAZOXIDE (U.S.P.)	1 EA	BO	NA	GM	300 MG	3.33333		01/01/2002	99/99/9999							
38779-0324-06		J1730		01/01/2002	99/99/9999	INJECTION, DIAZOXIDE, UP TO 300 MG	DIAZOXIDE (U.S.P.)	1 EA	BO	NA	GM	300 MG	3.33333		01/01/2002	99/99/9999							
38779-0330-01		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	99/99/9999							
38779-0330-03		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	99/99/9999							
38779-0330-04		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	99/99/9999							
38779-0330-05		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	99/99/9999							
38779-0330-06		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	99/99/9999							
38779-0364-01		J7622		02/07/2002	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		02/07/2002	99/99/9999							
38779-0364-01	KO	J7622	KO	02/07/2002	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		02/07/2002	99/99/9999							
38779-0364-03		J7622		02/07/2002	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		02/07/2002	99/99/9999							
38779-0364-03	KO	J7622	KO	02/07/2002	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		02/07/2002	99/99/9999							
38779-0364-06		J7622		02/07/2002	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		02/07/2002	99/99/9999							
38779-0364-06	KO	J7622	KO	02/07/2002	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		02/07/2002	99/99/9999							
38779-0388-03		J0475		01/01/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2002	99/99/9999							
38779-0388-04		J0475		01/01/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2002	99/99/9999							
38779-0388-05		J0475		01/01/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2002	99/99/9999							
38779-0388-09		J0475		04/22/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	JR	NA	GM	10 MG	100		04/22/2002	99/99/9999							
38779-0393-03		J0520		01/01/2002	10/17/2016	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	10/17/2016							
38779-0393-04		J0520		01/01/2002	10/17/2016	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	10/17/2016							
38779-0393-05		J0520		04/19/2002	10/17/2016	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		04/19/2002	10/17/2016							
38779-0393-06		J0520		01/01/2002	10/17/2016	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	10/17/2016							
38779-0403-01		J2765		04/25/2003	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P.)	1 EA	JR	NA	GM	10 MG	100		04/25/2003	99/99/9999							
38779-0403-04		J2765		01/01/2002	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2002	99/99/9999							
38779-0403-05		J2765		01/01/2002	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2002	99/99/9999							
38779-0405-01		J7638		01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-01	KO	J7638	KO	01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-03		J7638		01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-03	KO	J7638	KO	01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-04		J7638		01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-04	KO	J7638	KO	01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-05		J7638		01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-05	KO	J7638	KO	01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-06		J7638		01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0405-06	KO	J7638	KO	01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P., MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
38779-0423-04		J3230		01/01/2002	99/99/9999	INJECTION, CHLORPROMAZINE HCL, UP TO 50 MG	CHLORPROMAZINE HCL (U.S.P.)	1 EA	BO	NA	GM	50 MG	20		01/01/2002	99/99/9999							
38779-0423-05		J3230		01/01/2002	99/99/9999	INJECTION, CHLORPROMAZINE HCL, UP TO 50 MG	CHLORPROMAZINE HCL (U.S.P.)	1 EA	BO	NA	GM	50 MG	20		01/01/2002	99/99/9999							
38779-0454-03		J2440		01/01/2002	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		01/01/2002	99/99/9999							
38779-0454-04		J2440		01/01/2002	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		01/01/2002	99/99/9999							
38779-0454-05		J2440		01/01/2002	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		01/01/2002	99/99/9999							
38779-0468-03		J3420		04/25/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	1 EA	BO	NA	GM	1000 MCG	1000		04/25/2003	99/99/9999							
38779-0468-04		J3420		04/25/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	1 EA	BO	NA	GM	1000 MCG	1000		04/25/2003	99/99/9999							
38779-0468-05		J3420		04/25/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	1 EA	BO	NA	GM	1000 MCG	1000		04/25/2003	99/99/9999							
38779-0468-06		J3420		04/25/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	1 EA	BO	NA	GM	1000 MCG	1000		04/25/2003	99/99/9999							
38779-0495-04		J7604		01/01/2008	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1 EA	BO	NA	GM	1 GM	1		01/01/2008	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
38779-0495-04	KO	J7604	KO	01/01/2008	99/99/9999	ACETYLCSYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCSYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	99/99/9999						
38779-0495-05		J7604		01/01/2008	99/99/9999	ACETYLCSYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCSYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	99/99/9999						
38779-0495-05	KO	J7604	KO	01/01/2008	99/99/9999	ACETYLCSYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCSYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	99/99/9999						
38779-0495-08		J7604		01/01/2008	99/99/9999	ACETYLCSYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCSYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	99/99/9999						
38779-0495-08	KO	J7604	KO	01/01/2008	99/99/9999	ACETYLCSYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCSYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	99/99/9999						
38779-0495-09		J7604		01/01/2008	99/99/9999	ACETYLCSYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCSYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	99/99/9999						
38779-0495-09	KO	J7604	KO	01/01/2008	99/99/9999	ACETYLCSYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCSYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	99/99/9999						
38779-0534-05		J3490		04/25/2002	99/99/9999	UNCLASSIFIED DRUGS	CIPROFLOXACIN HCL (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	04/25/2002	99/99/9999						
38779-0534-08		J3490		04/25/2002	99/99/9999	UNCLASSIFIED DRUGS	CIPROFLOXACIN HCL (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	04/25/2002	99/99/9999						
38779-0534-09		J3490		04/25/2002	99/99/9999	UNCLASSIFIED DRUGS	CIPROFLOXACIN HCL (U.S.P.)	1 EA	JR	NA	GM		1 EA		1	04/25/2002	99/99/9999						
38779-0536-04		J2780		05/20/2002	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	JR	NA	GM		25 MG		40	05/20/2002	99/99/9999						
38779-0536-05		J2780		05/20/2002	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	JR	NA	GM		25 MG		40	05/20/2002	99/99/9999						
38779-0536-08		J2780		05/20/2002	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	JR	NA	GM		25 MG		40	05/20/2002	99/99/9999						
38779-0536-09		J2780		05/20/2002	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	JR	NA	GM		25 MG		40	05/20/2002	99/99/9999						
38779-0561-01		J0735		01/01/2002	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
38779-0561-03		J0735		01/01/2002	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
38779-0561-04		J0735		09/03/2002	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 EA	JR	NA	GM		1 MG		1000	09/03/2002	99/99/9999						
38779-0561-06		J0735		01/01/2002	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
38779-0571-05		J0280		01/01/2002	10/17/2016	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE DIHYDRATE (U.S.P.)	1 EA	BO	NA	GM		250 MG		4	09/26/2008	10/17/2016	01/01/2002	11/27/2003		4		
38779-0571-08		J0280		01/01/2002	10/17/2016	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE DIHYDRATE (U.S.P.)	1 EA	BO	NA	GM		250 MG		4	09/26/2008	10/17/2016	01/01/2002	11/27/2003		4		
38779-0599-01		J2150		01/01/2002	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	1 EA	BO	NA	GM		50 ML		0.08	01/01/2002	99/99/9999						
38779-0599-08		J2150		01/01/2002	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	1 EA	BO	NA	GM		50 ML		0.08	01/01/2002	99/99/9999						
38779-0599-09		J2150		01/01/2002	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	1 EA	BO	NA	GM		50 ML		0.08	01/01/2002	99/99/9999						
38779-0655-04		J3490		08/21/2002	99/99/9999	UNCLASSIFIED DRUGS	FAMOTIDINE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	08/21/2002	99/99/9999						
38779-0655-05		J3490		08/21/2002	99/99/9999	UNCLASSIFIED DRUGS	FAMOTIDINE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	08/21/2002	99/99/9999						
38779-0660-03		J7516		02/06/2002	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORIN A	1 EA	BO	NA	GM		250 MG		4	02/06/2002	99/99/9999						
38779-0660-04		J7516		02/06/2002	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORIN A	1 EA	BO	NA	GM		250 MG		4	02/06/2002	99/99/9999						
38779-0660-05		J7516		02/06/2002	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORIN A	1 EA	BO	NA	GM		250 MG		4	02/06/2002	99/99/9999						
38779-0660-06		J7516		02/06/2002	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORIN A	1 EA	BO	NA	GM		250 MG		4	02/06/2002	99/99/9999						
38779-0673-03		J2271		01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0673-04		J2271		01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0673-05		J2271		01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0673-07		J2271		01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2002	12/31/2014						
38779-0679-03		J0745		01/01/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		30 MG		33.33333	01/01/2002	99/99/9999						
38779-0679-04		J0745		01/01/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		30 MG		33.33333	01/01/2002	99/99/9999						
38779-0679-05		J0745		01/01/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		30 MG		33.33333	01/01/2002	99/99/9999						
38779-0731-01		J1170		04/23/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	04/23/2002	99/99/9999						
38779-0731-03		J1170		01/01/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	01/01/2002	99/99/9999						
38779-0731-04		J1170		01/01/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	01/01/2002	99/99/9999						
38779-0731-05		J1170		09/27/2007	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (1X100GM)	1 EA	JR	NA	GM		4 MG		250	09/27/2007	99/99/9999						
38779-0731-06		J1170		01/01/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	01/01/2002	99/99/9999						
38779-0767-03		J2310		01/01/2002	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL DIHYDRATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
38779-0767-06		J2310		01/01/2002	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL DIHYDRATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
38779-0855-03		J3130		04/25/2002	12/31/2014	INJECTION, TESTOSTERONE ENANTHATE, UP TO 200 MG	TESTOSTERONE ENANTHATE	1 EA	NA	NA	GM		200 MG		5	04/25/2002	12/31/2014						
38779-0855-04		J3130		04/25/2002	12/31/2014	INJECTION, TESTOSTERONE ENANTHATE, UP TO 200 MG	TESTOSTERONE ENANTHATE	1 EA	NA	NA	GM		200 MG		5	04/25/2002	12/31/2014						
38779-0873-04		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2004	99/99/9999						
38779-0873-05		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2004	99/99/9999						
38779-0873-08		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2004	99/99/9999						
38779-0873-09		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	01/01/2004	99/99/9999						
38779-0885-03		J1960		11/22/2002	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	1 EA	BO	NA	GM		2 MG		500	11/22/2002	99/99/9999						
38779-0885-06		J1960		11/22/2002	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	1 EA															

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
38779-0927-01		J2060		01/01/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO NA GM	2 MG			500		500	01/01/2002	99/99/9999						
38779-0927-03		J2060		01/01/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO NA GM	2 MG			500		500	01/01/2002	99/99/9999						
38779-0927-04		J2060		01/01/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO NA GM	2 MG			500		500	01/01/2002	99/99/9999						
38779-0927-05		J2060		01/01/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO NA GM	2 MG			500		500	01/01/2002	99/99/9999						
38779-0927-06		J2060		01/01/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO NA GM	2 MG			500		500	01/01/2002	99/99/9999						
38779-0927-08		J2060		01/01/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO NA GM	2 MG			500		500	01/01/2002	99/99/9999						
38779-0944-07		J0270		01/01/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO NA GM	1.25 MCG			800000		800000	01/01/2002	99/99/9999						
38779-0944-09		J0270		01/01/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO NA GM	1.25 MCG			800000		800000	01/01/2002	99/99/9999						
38779-0989-04		J3490		01/28/2002	99/99/9999	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	1 EA	BO NA GM	1 EA			1		1	01/28/2002	99/99/9999						
38779-0989-05		J3490		01/28/2002	99/99/9999	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	1 EA	BO NA GM	1 EA			1		1	01/28/2002	99/99/9999						
38779-0989-08		J3490		01/28/2002	99/99/9999	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	1 EA	BO NA GM	1 EA			1		1	01/28/2002	99/99/9999						
38779-0989-09		J3490		01/28/2002	99/99/9999	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	1 EA	BO NA GM	1 EA			1		1	01/28/2002	99/99/9999						
38779-1502-00		J2760		01/01/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM	5 MG			200		200	01/01/2002	99/99/9999						
38779-1502-03		J2760		05/22/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM	5 MG			200		200	05/22/2002	99/99/9999						
38779-1502-06		J2760		01/01/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM	5 MG			200		200	01/01/2002	99/99/9999						
38779-1502-09		J2760		05/22/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM	5 MG			200		200	05/22/2002	99/99/9999						
38779-1756-00		J3010		01/01/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 EA	BO NA GM	0.1 MG			10000		10000	01/01/2002	99/99/9999						
38779-1756-03		J3010		04/23/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 EA	JR NA GM	0.1 MG			10000		10000	04/23/2002	99/99/9999						
38779-1756-06		J3010		01/01/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 EA	JR NA GM	0.1 MG			10000		10000	01/01/2002	99/99/9999						
38779-1756-09		J3010		01/01/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 EA	BO NA GM	0.1 MG			10000		10000	01/01/2002	99/99/9999						
38779-1764-00		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1 EA	BO NA GM	1 MG			1000		1000	01/01/2007	99/99/9999						
38779-1764-03		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1 EA	BO NA GM	1 MG			1000		1000	01/01/2007	99/99/9999						
38779-1764-06		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1 EA	BO NA GM	1 MG			1000		1000	01/01/2007	99/99/9999						
38779-1766-03		J2175		01/01/2002	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	1 EA	BO NA GM	100 MG			10		10	01/01/2002	99/99/9999						
38779-1766-04		J2175		01/01/2002	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	1 EA	BO NA GM	100 MG			10		10	01/01/2002	99/99/9999						
38779-1766-05		J2175		01/01/2002	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	1 EA	BO NA GM	100 MG			10		10	01/01/2002	99/99/9999						
38779-1901-03		J1000		01/01/2002	99/99/9999	INJECTION, DEPO-ESTRADIOL CYPIONATE, UP TO 5 MG	ESTRADIOL CYPIONATE (U.S.P.)	1 EA	BO NA GM	5 MG			200		200	01/01/2002	99/99/9999						
38779-1901-04		J1000		01/01/2002	99/99/9999	INJECTION, DEPO-ESTRADIOL CYPIONATE, UP TO 5 MG	ESTRADIOL CYPIONATE (U.S.P.)	1 EA	BO NA GM	5 MG			200		200	01/01/2002	99/99/9999						
38779-1901-05		J1000		01/01/2002	99/99/9999	INJECTION, DEPO-ESTRADIOL CYPIONATE, UP TO 5 MG	ESTRADIOL CYPIONATE (U.S.P.)	1 EA	BO NA GM	5 MG			200		200	01/01/2002	99/99/9999						
38779-1905-01		J1094		01/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE MICRONIZED (ANHYDROUS)	1 EA	NA NA GM	1 MG			1000		1000	01/01/2003	99/99/9999						
38779-1905-03		J1094		01/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE MICRONIZED (ANHYDROUS)	1 EA	NA NA GM	1 MG			1000		1000	01/01/2003	99/99/9999						
38779-1905-04		J1094		01/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE MICRONIZED (ANHYDROUS)	1 EA	NA NA GM	1 MG			1000		1000	01/01/2003	99/99/9999						
38779-1905-05		J1094		01/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO NA GM	1 MG			1000		1000	01/01/2003	99/99/9999						
38779-1931-01		J1835		04/25/2002	99/99/9999	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE	1 EA	BO NA GM	50 MG			20		20	04/25/2002	99/99/9999						
38779-1943-05		J2800		04/25/2002	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (U.S.P.)	1 EA	BO NA GM	10 ML			1		1	04/25/2002	99/99/9999						
38779-1943-08		J2800		04/25/2002	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (U.S.P.)	1 EA	BO NA GM	10 ML			1		1	04/25/2002	99/99/9999						
38779-1943-09		J2800		04/25/2002	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (U.S.P.)	1 EA	BO NA GM	10 ML			1		1	04/25/2002	99/99/9999						
38779-1968-07		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (USP)	1 EA	BO NA GM	1 EA			1		1	01/01/2002	99/99/9999						
38779-1968-09		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (USP)	1 EA	BO NA GM	1 EA			1		1	01/01/2002	99/99/9999						
38779-2087-03		J7643		05/02/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1 EA	JR NA GM	1 MG			1000		1000	05/02/2002	99/99/9999						
38779-2087-03	KO	J7643	KO	05/02/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1 EA	JR NA GM	1 MG			1000		1000	05/02/2002	99/99/9999						
38779-2087-06		J7643		05/02/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1 EA	JR NA GM	1 MG			1000		1000	05/02/2002	99/99/9999						
38779-2087-06	KO	J7643	KO	05/02/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1 EA	JR NA GM	1 MG			1000		1000	05/02/2002	99/99/9999						
38779-2363-05		J1956		10/25/2007	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN HEMIHYDRATE (1X100MG)	1 EA	BO NA GM	250 MG			4		4	10/25/2007	99/99/9999						
39822-0277-02		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BACIIM (STERILE) 50000 U	1 EA	VL IM EA	1 EA			1		1	01/01/2002	99/99/9999						
39822-0412-01		J3260		01/01/2007	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (BULK VIAL,PF) 1.2 GM	1 EA	VL IV EA	80 MG			15		15	01/01/2007	99/99/9999						
39822-0412-06		J3260		01/01/2007	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (BULK VIAL,PF) 1.2 GM	6 EA	VL IV EA	80 MG			15		15	01/01/2007	99/99/9999						
39822-0615-01		J0770		01/01/2002	99/99/9999	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE SODIUM (VIAL,STERILE) 150 MG	1 EA	VL IJ EA	150 MG			1		1	01/01/2002	99/99/9999						
39822-0706-02		J3000		01/01/2002	99/99/9999	INJECTION, STREPTOMYCIN, UP TO 1 GM	STREPTOMYCIN SULFATE (STERILE) 1 GM	1 EA	VL IM EA	1 GM			1		1	01/01/2002	99/99/9999						
39822-0710-01		J1451		12/14/2007	06/06/2018	INJECTION, FOMEPIZOLE, 15 MG	FOMEPIZOLE (1X1.5ML,PF) 1 GM/ML	1.5 ML	VL IV ML	15 MG			66.66666		66.66666	12/14/2007	06/06/2018						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
42023-0116-01	J2590			02/29/2008	09/06/2018	INJECTION, OXYTOCIN, UP TO 10 UNITS	PITOCIN (1X10ML,MDV) 10 U/ML	10	ML	VL	IJ	ML	10 U		1	02/29/2008	09/06/2018							
42023-0116-25	J2590			02/01/2008	99/99/9999	INJECTION, OXYTOCIN, UP TO 10 UNITS	PITOCIN (25X1ML) 10 U/ML	1	ML	VL	IJ	ML	10 U		1	02/01/2008	99/99/9999							
43292-0556-31	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALERTAB 25 MG	100	EA	BX	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
43292-0557-05	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALERCAP 25 MG	100	EA	NA	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
43292-0557-19	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SLEEP-TABS 25 MG	36	EA	NA	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
43292-0557-65	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (MAX. STR.) 50 MG	50	EA	NA	PO	EA	50 MG		1	01/01/2002	99/99/9999							
43292-0557-78	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SLEEP-TABS 25 MG	100	EA	NA	PO	EA	50 MG		0.5	01/01/2002	99/99/9999							
44087-0004-07	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	SEROSTIM 4 MG	1	EA	VL	SC	EA	1 MG		4	01/01/2002	99/99/9999							
44087-0005-07	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	SEROSTIM (S.D.V., W/DILUENT) 5 MG	1	EA	VL	SC	EA	1 MG		5	01/01/2002	99/99/9999							
44087-0006-07	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	SEROSTIM (S.D.V., W/DILUENT) 6 MG	1	EA	VL	SC	EA	1 MG		6	01/01/2002	99/99/9999							
44087-0022-03	Q3026			01/01/2003	12/31/2013	INJECTION, INTERFERON BETA-1A, 11 MCG FOR SUBCUTANEOUS USE	REBIF (SRN,PREFILLED,27G,PF) 22 MCG/0.5 ML	0.5	ML	SR	SC	ML	11 MCG		4	01/01/2003	12/31/2013							
44087-0044-03	Q3026			01/01/2003	12/31/2013	INJECTION, INTERFERON BETA-1A, 11 MCG FOR SUBCUTANEOUS USE	REBIF (SRN,PREFILLED,27G,PF) 44 MCG/0.5 ML	0.5	ML	SR	SC	ML	11 MCG		8	01/01/2003	12/31/2013							
44087-1005-02	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	SAIZEN (VIAL, W/DILUENT) 5 MG	1	EA	VL	SC	EA	1 MG		5	01/01/2002	99/99/9999							
44087-1080-01	J2941			10/22/2004	06/01/2018	INJECTION, SOMATROPIN, 1 MG	SAIZEN (VIAL, W/DILUENT) 8.8 MG	1	CT	VL	IJ	EA	1 MG		8.8	10/22/2004	06/01/2018							
44087-1088-01	J2941			01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	SAIZEN (VIAL, W/DILUENT) 8.8 MG	1	EA	VL	IJ	EA	1 MG		8.8	01/01/2002	99/99/9999							
44087-1112-01	J3490			06/15/2004	99/99/9999	UNCLASSIFIED DRUGS	GONAL-F RFF (29GX1/2 NEEDLE,PEN) 450 IU/0.75 ML	0.75	ML	CR	SC	ML	1 EA		1	06/15/2004	99/99/9999							
44087-1113-01	J3490			06/15/2004	99/99/9999	UNCLASSIFIED DRUGS	GONAL-F RFF (29GX1/2,PEN) 300 IU/0.5 ML	0.5	ML	CR	SC	ML	1 EA		1	06/15/2004	99/99/9999							
44087-1114-01	J3490			06/15/2004	99/99/9999	UNCLASSIFIED DRUGS	GONAL-F RFF (29GX1/2,PEN) 900 IU/1.5 ML	1.5	ML	CR	SC	ML	1 EA		1	06/15/2004	99/99/9999							
44087-1150-01	J3490			11/10/2003	99/99/9999	UNCLASSIFIED DRUGS	OVIDREL (SRN,PREFILLED SYRINGE) 0.25 MG/0.5 ML	0.5	ML	SR	SC	ML	1 EA		1	11/10/2003	99/99/9999							
63323-0580-20	J0461			05/22/2018	99/99/9999	INJECTION, ATROPINE SULFATE, 0.01 MG	ATROPINE SULFATE 0.4 MG/1 ML	20	ML	VL	IJ	ML	0.01 MG		40	05/22/2018	99/99/9999							
44087-3388-07	J2941			04/07/2003	99/99/9999	INJECTION, SOMATROPIN, 1 MG	ZORBTIVE (MDV, VIALS W/ DILUENT) 8.8 MG	1	EA	VL	SC	EA	1 MG		8.8	04/07/2003	99/99/9999							
44087-6075-01	J3355			01/01/2006	99/99/9999	INJECTION, UROFOLLITROPIN, 75 IU	METRODIN 75 IU	1	EA	NA	IM	EA	75 IU		1	01/01/2006	99/99/9999							
44087-6075-03	J3355			01/01/2006	99/99/9999	INJECTION, UROFOLLITROPIN, 75 IU	METRODIN 75 IU	1	EA	NA	IM	EA	75 IU		1	01/01/2006	99/99/9999							
44087-6075-04	J3355			01/01/2006	99/99/9999	INJECTION, UROFOLLITROPIN, 75 IU	METRODIN 75 IU	1	EA	NA	IM	EA	75 IU		1	01/01/2006	99/99/9999							
44087-6150-01	J3355			01/01/2006	99/99/9999	INJECTION, UROFOLLITROPIN, 75 IU	METRODIN 150 IU	1	EA	NA	IM	EA	75 IU		2	01/01/2006	99/99/9999							
44087-8822-01	Q3026			02/14/2005	12/31/2013	INJECTION, INTERFERON BETA-1A, 11 MCG FOR SUBCUTANEOUS USE	REBIF (TITRATION PACK,PF) 44 MCG/ML	4.2	ML	BX	SC	ML	11 MCG		4	02/14/2005	12/31/2013							
44087-9005-01	J3490			06/07/2004	99/99/9999	UNCLASSIFIED DRUGS	GONAL-F RFF 75 IU	1	EA	VL	SC	EA	1 EA		1	06/07/2004	99/99/9999							
44087-9005-06	J3490			06/07/2004	99/99/9999	UNCLASSIFIED DRUGS	GONAL-F RFF 75 IU	1	EA	VL	SC	EA	1 EA		1	06/07/2004	99/99/9999							
44087-9030-01	J3490			05/10/2004	99/99/9999	UNCLASSIFIED DRUGS	GONAL-F (M.D.V.) 450 IU	1	EA	VL	SC	EA	1 EA		1	05/10/2004	99/99/9999							
44087-9070-01	J3490			05/07/2007	99/99/9999	UNCLASSIFIED DRUGS	GONAL-F (MDV) 1200 IU	1	EA	VL	SC	EA	1 EA		1	05/07/2007	99/99/9999							
44206-0300-01	J2791			01/01/2008	99/99/9999	INJECTION, RHO(D) IMMUNE GLOBULIN (HUMAN), (RHOPHYLAC), INTRAMUSCULAR OR INTRAVENOUS, 100 IU	RHOPHYLAC (W/SAFETY NEEDLE) 750 IU/ML	2	ML	SR	IJ	ML	100 IU		7.5	01/01/2008	99/99/9999							
44206-0300-10	J2791			01/01/2008	99/99/9999	INJECTION, RHO(D) IMMUNE GLOBULIN (HUMAN), (RHOPHYLAC), INTRAMUSCULAR OR INTRAVENOUS, 100 IU	RHOPHYLAC (W/SAFETY NEEDLE) 750 IU/ML	2	ML	SR	IJ	ML	100 IU		7.5	01/01/2008	99/99/9999							
44206-0416-03	J1566			01/01/2006	11/17/2016	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	CARMUNE NF (PF,NANOFILTERED) 3 GM	1	EA	VL	IV	EA	500 MG		6	01/01/2006	11/17/2016							
44206-0417-06	J1566			01/01/2006	99/99/9999	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	CARMUNE NF (PF,NANOFILTERED) 6 GM	1	EA	VL	IV	EA	500 MG		12	01/01/2006	99/99/9999							
44206-0418-12	J1566			01/01/2006	99/99/9999	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	CARMUNE NF (PF,NANOFILTERED) 12 GM	1	EA	VL	IV	EA	500 MG		24	01/01/2006	99/99/9999							
47781-0603-20	J9045			04/02/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	5	ML	VL	IV	ML	50 MG		0.2	04/02/2018	99/99/9999							
45802-0303-21	J7506			12/12/2007	04/16/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP,BLISTER PACK) 10 MG	21	EA	DP	PO	EA	5 MG		2	12/12/2007	04/16/2013							
45802-0303-67	J7506			12/12/2007	04/16/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP,BLISTER PACK) 10 MG	48	EA	DP	PO	EA	5 MG		2	12/12/2007	04/16/2013							
45802-0733-21	J7506			12/12/2007	04/16/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP,BLISTER PACK) 5 MG	21	EA	DP	PO	EA	5 MG		1	12/12/2007	04/16/2013							
45802-0733-67	J7506			12/12/2007	04/16/2013	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP,BLISTER PACK) 5 MG	48	EA	DP	PO	EA	5 MG		1	12/12/2007	04/16/2013							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
45802-0758-30		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 12.5 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
45802-0759-30		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 25 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
47682-0858-87		Q0163		01/01/2007	06/01/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MEDI-FIRST DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	100 EA	BX	PO	EA		50 MG		0.5	01/01/2007	06/01/2012						
00409-3459-07		J1170		06/27/2018	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (PF,LATEX-FREE) 2 MG/1 ML	1 ML	AM	IJ	ML		4 MG		0.5	06/27/2018	99/99/9999						
48879-0001-01		A4216		01/01/2006	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INHALATION (AL7023)	3 ML	EA	IH	ML		10 ML		0.1	01/01/2006	02/03/2016						
48879-0001-02		A4216		01/01/2006	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INHALATION (AL7025)	5 ML	EA	IH	ML		10 ML		0.1	01/01/2006	02/03/2016						
48879-0002-01		A4216		01/01/2006	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SALINE SOLUTION (AL7453) 0.45%	3 ML	EA	IH	ML		10 ML		0.1	01/01/2006	99/99/9999						
48879-0002-02		A4216		01/01/2006	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SALINE SOLUTION (AL7455) 0.45%	5 ML	EA	IH	ML		10 ML		0.1	01/01/2006	02/03/2016						
48879-0003-01		A4216		01/01/2006	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SALINE SOLUTION (AL7093) 0.9%	3 ML	EA	IH	ML		10 ML		0.1	01/01/2006	99/99/9999						
48879-0003-02		A4216		01/01/2006	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SALINE SOLUTION (AL7095) 0.9%	5 ML	EA	IH	ML		10 ML		0.1	01/01/2006	99/99/9999						
48879-0003-07		A4216		01/01/2006	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SALINE SOLUTION (AL4015) 0.9%	15 ML	PC	IH	ML		10 ML		0.1	01/01/2006	02/03/2016						
49281-0545-05		J3490		01/01/2002	12/14/2017	UNCLASSIFIED DRUGS	ACTHIB (SDV W/DIL,TAX INCL,PF) 10 MCG	1 EA	VL	IM	EA		1 EA		1	01/01/2002	12/14/2017						
49281-0880-01		J9031		01/01/2002	09/01/2013	BCG (INTRAVESICAL) PER INSTILLATION	THERACYS (S.D.V. W/DILUENT,PF) 81 MG	1 EA	VL	IL	EA		1 ATION		1	01/01/2002	09/01/2013						
49348-0044-04		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VALU-DRYL ALLERGY 25 MG	24 EA	BX	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
49348-0044-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VALU-DRYL ALLERGY 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
49348-0045-34		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VALU-DRYL ALLERGY CHILDREN'S 12.5 MG/5 ML	120 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
49348-0205-37		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VALU-DRYL ALLERGY CHILDREN'S (AF,CHERRY) 12.5 MG/5 ML	236 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
49348-0282-08		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VALU-DRYL ALLERGY 25 MG	48 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
49348-0564-04		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VALU-DRYL ALLERGY 25 MG	24 EA	BX	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
49483-0061-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ANTIHISTAMINE 25 MG CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
47781-0604-27		J9045		04/02/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	15 ML	VL	IV	ML		50 MG		0.2	04/02/2018	99/99/9999						
49483-0061-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ANTIHISTAMINE 25 MG SODIUM CHLORIDE (NEBU-SOL/MTR DOSE DSPNS) 0.9%	1000 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
49502-0501-20		A4218		01/01/2006	99/99/9999	STERILE SALINE OR WATER, METERED DOSE DISPENSER, 10 ML	DIJONER (VIAL,U.D.) 3 MG/3 ML-0.5 MG/3 ML	120 ML	EA	IH	ML		10 ML		0.1	01/01/2006	99/99/9999						
49502-0672-30		J7620		01/01/2006	04/30/2014	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	DIJONER (VIAL,U.D.) 3 MG/3 ML-0.5 MG/3 ML	30 ML	PC	IH	ML		3 MG		0.33333	01/01/2006	04/30/2014						
49502-0672-60		J7620		01/01/2006	06/30/2014	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	DIJONER (VIAL,U.D.) 3 MG/3 ML-0.5 MG/3 ML	60 ML	PC	IH	ML		3 MG		0.33333	01/01/2006	06/30/2014						
49502-0692-03		J7613		04/01/2008	06/17/2016	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ACCUNEB (PF) 0.021%	3 ML	PC	IH	ML		1 MG		0.21	04/01/2008	06/17/2016						
49502-0692-03	KO	J7613	KO	04/01/2008	06/17/2016	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ACCUNEB (PF) 0.021%	3 ML	PC	IH	ML		1 MG		0.21	04/01/2008	06/17/2016						
49502-0693-03		J7613		04/01/2008	08/31/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ACCUNEB (PF) 0.042%	3 ML	PC	IH	ML		1 MG		0.42	04/01/2008	08/31/2013						
49502-0693-03	KO	J7613	KO	04/01/2008	08/31/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ACCUNEB (PF) 0.042%	3 ML	PC	IH	ML		1 MG		0.42	04/01/2008	08/31/2013						
49614-0146-62		Q0163		10/13/2003	07/18/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MEDICINE SHOPPE NITE TIME SLEEP (MINI-CAPLET) 25 MG	24 EA	BO	PO	EA		50 MG		0.5	10/13/2003	07/18/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
49614-0379-26		Q0163		01/01/2004	07/18/2013	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	THE MEDICINE SHOPPE MEDI-PHEDRYL (MAY CAUSE DROWSINESS,AF) 12.5 MG/5 ML	118 ML	BO	PO	ML		50 MG		0.05	01/01/2004	07/18/2013						
49884-0289-01		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 20 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
49884-0290-01		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
49884-0290-04		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	250 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
49884-0290-05		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	500 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
49884-0673-14		J8515		01/01/2006	99/99/9999	CABERGOLINE, ORAL, 0.25 MG	CABERGOLINE 0.5 MG	8 EA	BO	PO	EA		0.25 MG		2	01/01/2006	99/99/9999						
49884-0724-01		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDROXYUREA 500 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
49884-0753-13		J8999		01/26/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	FLUTAMIDE 125 MG	180 EA	BO	PO	EA		1 EA		1	01/26/2006	99/99/9999						
49884-0907-38		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG/ML	240 ML	BO	PO	ML		1 EA		1	01/01/2002	99/99/9999						
49884-0907-61		J8999		05/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG/ML	480 ML	BO	PO	ML		1 EA		1	05/01/2004	99/99/9999						
49884-0922-02		J8999		02/09/2004	10/30/2014	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MERCAPTOPYRINE 50 MG	60 EA	BO	PO	EA		1 EA		1	02/09/2004	10/30/2014						
49884-0922-04		J8999		11/18/2004	10/30/2014	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MERCAPTOPYRINE 50 MG	250 EA	BO	PO	EA		1 EA		1	11/18/2004	10/30/2014						
49999-0003-15		Q0163		07/11/2002	06/01/2018	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15 EA	BO	PO	EA		50 MG		0.5	07/11/2002	06/01/2018						
49999-0003-20		Q0163		02/24/2005	06/01/2017	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	20 EA	BO	PO	EA		50 MG		0.5	02/24/2005	06/01/2017						
49999-0003-30		Q0163		07/11/2002	06/01/2018	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30 EA	BO	PO	EA		50 MG		0.5	07/11/2002	06/01/2018						
49999-0008-00		J7506		12/01/2003	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	100 EA	BO	PO	EA		5 MG		1	12/01/2003	06/01/2014						
49999-0008-05		J7506		05/16/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	5 EA	NA	PO	EA		5 MG		1	05/16/2008	12/31/2015						
49999-0008-20		J7506		07/16/2002	01/01/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	20 EA	BO	PO	EA		5 MG		1	07/16/2002	01/01/2015						
49999-0008-30		J7506		07/06/2004	01/01/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	30 EA	BO	PO	EA		5 MG		1	07/06/2004	01/01/2015						
49999-0008-40		J7506		01/27/2006	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	40 EA	BO	PO	EA		5 MG		1	01/27/2006	06/01/2014						
49999-0008-55		J7506		08/28/2002	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	55 EA	BO	PO	EA		5 MG		1	08/28/2002	06/01/2014						
49999-0028-05		J7506		03/13/2008	12/31/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	5 EA	BO	PO	EA		5 MG		2	03/13/2008	12/31/2014						
49999-0028-12		J7506		07/16/2002	12/31/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	12 EA	BO	PO	EA		5 MG		2	07/16/2002	12/31/2014						
49999-0028-14		J7506		01/27/2006	12/31/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	14 EA	BO	PO	EA		5 MG		2	01/27/2006	12/31/2014						
49999-0028-15		J7506		07/11/2002	01/01/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	15 EA	BO	PO	EA		5 MG		2	07/11/2002	01/01/2015						
49999-0028-20		J7506		07/16/2002	01/01/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	07/16/2002	01/01/2015						
49999-0028-28		J7506		07/01/2005	01/01/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	28 EA	BO	PO	EA		5 MG		2	07/01/2005	01/01/2015						
49999-0028-30		J7506		07/11/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30 EA	BO	PO	EA		5 MG		2	07/11/2002	12/31/2015						
49999-0028-40		J7506		07/16/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	40 EA	BO	PO	EA		5 MG		2	07/16/2002	12/31/2015						
49999-0028-48		J7506		07/06/2004	12/31/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	48 EA	BO	PO	EA		5 MG		2	07/06/2004	12/31/2014						
49999-0028-50		J7506		07/16/2002	12/31/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	50 EA	BO	PO	EA		5 MG		2	07/16/2002	12/31/2014						
49999-0028-60		J7506		03/30/2005	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	60 EA	BO	PO	EA		5 MG		2	03/30/2005	12/31/2015						
49999-0028-90		J7506		03/30/2005	12/31/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	90 EA	BO	PO	EA		5 MG		2	03/30/2005	12/31/2014						
49999-0036-12		Q0178		10/15/2004	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	12 EA	BO	PO	EA		50 MG		2	10/15/2004	12/31/2013						
49999-0036-60		Q0178		07/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	60 EA	BO	PO	EA		50 MG		2	07/01/2002	12/31/2013						
49999-0059-06		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	6 EA	BO	PO	EA		0.25 MG		16	01/01/2006	99/99/9999						
49999-0086-00		J8499		09/01/2006	01/01/2015	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100 EA	BO	PO	EA		1 EA		1	09/01/2006	01/01/2015						
49999-0086-25		J8499		07/29/2002	01/01/2015	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	BO	PO	EA		1 EA		1	07/29/2002	01/01/2015						
49999-0086-30		J8499		07/13/2005	06/01/2017	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30 EA	BO	PO	EA		1 EA		1	07/13/2005	06/01/2017						
49999-0086-90		J8499		07/13/2005	01/01/2015	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	90 EA	BO	PO	EA		1 EA		1	07/13/2005	01/01/2015						
49999-0090-05		Q0170		04/15/2005	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	5 EA	BO	PO	EA		25 MG		1	04/15/2005	12/31/2013						
49999-0090-10		Q0170		06/05/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10 EA	BO	PO	EA		25 MG		1	06/05/2002	12/31/2013						
49999-0090-12		Q0170		05/07/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12 EA	BO	PO	EA		25 MG		1	05/07/2003	12/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
49999-0090-15		Q0170		12/01/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	25	MG	1	12/01/2003	12/31/2013						
49999-0090-20		Q0170		10/15/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	25	MG	1	10/15/2003	12/31/2013						
49999-0090-30		Q0170		04/15/2005	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	25	MG	1	04/15/2005	12/31/2013						
49999-0090-60		Q0170		02/10/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	BO	PO	EA	25	MG	1	02/10/2004	12/31/2013						
49999-0091-04		Q0163		05/07/2003	05/10/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	4	EA	BO	PO	EA	50	MG	1	05/07/2003	05/10/2012						
49999-0091-15		Q0163		03/26/2003	12/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15	EA	BO	PO	EA	50	MG	1	03/26/2003	12/31/2014						
49999-0091-20		Q0163		09/03/2002	01/01/2015	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20	EA	BO	PO	EA	50	MG	1	09/03/2002	01/01/2015						
49999-0091-30		Q0163		05/07/2003	05/10/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30	EA	BO	PO	EA	50	MG	1	05/07/2003	05/10/2012						
49999-0091-60		Q0163		05/07/2003	01/01/2015	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	60	EA	BO	PO	EA	50	MG	1	05/07/2003	01/01/2015						
49999-0096-04		Q0144		01/27/2006	01/01/2015	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	4	EA	BO	PO	EA	1	GM	0.25	01/27/2006	01/01/2015						
49999-0096-06		Q0144		08/08/2002	01/01/2015	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6	EA	BO	PO	EA	1	GM	0.25	08/08/2002	01/01/2015						
49999-0110-00		J7506		07/06/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	100	EA	BO	PO	EA	5	MG	4	07/06/2004	12/31/2015						
49999-0110-06		J7506		08/27/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	6	EA	BO	PO	EA	5	MG	4	08/27/2002	12/31/2015						
49999-0110-07		J7506		04/06/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	7	EA	BO	PO	EA	5	MG	4	04/06/2005	12/31/2015						
49999-0110-10		J7506		07/06/2004	01/01/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	10	EA	BO	PO	EA	5	MG	4	07/06/2004	01/01/2015						
49999-0110-12		J7506		07/06/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	12	EA	BO	PO	EA	5	MG	4	07/06/2004	12/31/2015						
49999-0110-14		J7506		07/06/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	14	EA	BO	PO	EA	5	MG	4	07/06/2004	12/31/2015						
49999-0110-15		J7506		03/27/2006	01/01/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	15	EA	BO	PO	EA	5	MG	4	03/27/2006	01/01/2015						
49999-0110-18		J7506		10/15/2004	01/01/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	18	EA	BO	PO	EA	5	MG	4	10/15/2004	01/01/2015						
49999-0110-20		J7506		07/11/2002	01/01/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	20	EA	BO	PO	EA	5	MG	4	07/11/2002	01/01/2015						
49999-0110-21		J7506		02/24/2005	01/01/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	21	EA	BO	PO	EA	5	MG	4	02/24/2005	01/01/2015						
49999-0110-30		J7506		03/26/2003	01/01/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	03/26/2003	01/01/2015						
49999-0153-21		J7509		09/03/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21	EA	DP	PO	EA	4	MG	1	09/03/2002	99/99/9999						
49999-0231-35		J8499		06/02/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	35	EA	BO	PO	EA	1	EA	1	06/02/2005	99/99/9999						
49999-0260-15		Q0144		07/01/2003	01/01/2015	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.04	07/01/2003	01/01/2015						
49999-0262-04		Q0170		07/01/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120	ML	BO	PO	ML	25	MG	0.05	07/01/2003	12/31/2013						
49999-0335-08		J7510		02/10/2004	01/01/2015	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240	ML	BO	PO	ML	5	MG	0.6	02/10/2004	01/01/2015						
49999-0335-24		J7510		05/10/2004	01/01/2015	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (CHERRY) 15 MG/5 ML	240	ML	BO	PO	ML	5	MG	0.6	05/10/2004	01/01/2015						
49999-0339-12		J8498		09/01/2006	01/01/2015	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 12.5 MG	12	EA	BX	RC	EA	1	EA	1	09/01/2006	01/01/2015						
49999-0340-12		J8498		01/01/2006	01/01/2015	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	01/01/2015						
49999-0344-25		J7613		04/01/2008	01/01/2015	COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	01/01/2015						
49999-0344-25	KO	J7613	KO	04/01/2008	01/01/2015	COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3	ML	PC	IH	ML	1	MG	0.83	04/01/2008	01/01/2015						
49999-0380-24		None		06/09/2004	01/01/2015	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	24	EA	DP	PO	EA	2.5	MG	1	06/09/2004	01/01/2015						
49999-0385-10		J8499		06/09/2004	01/01/2015	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	10	EA	BO	PO	EA	1	EA	1	06/09/2004	01/01/2015						
49999-0385-15		J8499		06/09/2004	01/01/2015	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	15	EA	BO	PO	EA	1	EA	1	06/09/2004	01/01/2015						
49999-0385-25		J8499		06/09/2004	01/01/2015	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25	EA	BO	PO	EA	1	EA	1	06/09/2004	01/01/2015						
49999-0385-40		J8499		06/02/2005	01/01/2015	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40	EA	BO	PO	EA	1	EA	1	06/02/2005	01/01/2015						
49999-0437-03		J7506		08/12/2004	01/01/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 50 MG	3	EA	BO	PO	EA	5	MG	10	08/12/2004	01/01/2015						
49999-0525-10		J1200		01/25/2008	02/03/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE 50 MG/ML	1	ML	VL	U	ML	50	MG	1	01/25/2008	02/03/2016						
49999-0582-15		Q0144		01/27/2006	01/01/2015	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	01/27/2006	01/01/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
49999-0671-50		J2001		05/16/2008	01/01/2015	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (1X50ML) 1%	50	ML	NA	EP	ML	10	MG	1	05/16/2008	01/01/2015							
49999-0786-06		Q0144		01/11/2006	01/01/2015	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6	EA	BO	PO	EA	1	GM	0.25	01/11/2006	01/01/2015							
49999-0902-20		Q0169		01/11/2007	01/01/2015	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 12.5 MG	20	EA	BO	PO	EA	12.5	MG	1	01/11/2007	01/01/2015							
49999-0929-01		J7510		04/20/2007	01/01/2015	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	04/20/2007	01/01/2015							
49999-0936-00		J7517		12/21/2007	01/01/2015	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT 250 MG	100	EA	BO	PO	EA	250	MG	1	12/21/2007	01/01/2015							
49999-0936-30		J7517		04/30/2007	12/31/2014	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT 250 MG	30	EA	BO	PO	EA	250	MG	1	04/30/2007	12/31/2014							
49999-0937-30		J7517		04/30/2007	12/31/2014	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	CELLCEPT 500 MG	30	EA	BO	PO	EA	250	MG	2	04/30/2007	12/31/2014							
49999-0986-30		J8999		06/14/2007	01/01/2015	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	AROMASIN 25 MG	30	EA	BO	PO	EA	1	EA	1	06/14/2007	01/01/2015							
49999-0993-10		J1815		06/14/2007	01/01/2015	INJECTION, INSULIN, PER 5 UNITS	HUMULIN 70 U/ML-30 U/ML	10	ML	VL	SC	ML	5	U	20	06/14/2007	01/01/2015							
49999-0994-10		J1815		06/14/2007	01/01/2015	INJECTION, INSULIN, PER 5 UNITS	LANTUS 100 U/ML	10	ML	VL	SC	ML	5	U	20	06/14/2007	01/01/2015							
50111-0794-78		J0456		07/25/2007	06/30/2013	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN (USP) 500 MG	10	EA	VL	IV	EA	500	MG	1	07/25/2007	06/30/2013							
50242-0018-21		J2941		01/01/2002	07/31/2013	INJECTION, SOMATROPIN, 1 MG	NUTROPIN (VIAL W/DILUENT) 10 MG	1	EA	VL	SC	EA	1	MG	10	01/01/2002	07/31/2013							
50242-0020-20		J2941		01/01/2002	11/30/2013	INJECTION, SOMATROPIN, 1 MG	NUTROPIN (VIAL) 10 MG	1	EA	VL	SC	EA	1	MG	10	01/01/2002	11/30/2013							
70842-0160-10		J2265		08/24/2018	99/99/9999	INJECTION, MINOCYCLINE HYDROCHLORIDE, 1 MG	MINOCIN (LYOPHILIZED) 100 MG	10	EA	VL	IV	EA	1	MG	100	08/24/2018	99/99/9999							
72439-0500-10		J3480		08/29/2018	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (AMPULE) 2 MEQ/1 ML	10	ML	AM	IV	ML	2	MEQ	1	08/29/2018	99/99/9999							
52652-2001-01		None		04/25/2017	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	XATMEP 2.5 MG/1 ML	120	ML	BO	PO	ML	2.5	MG	1	04/25/2017	99/99/9999							
50242-0022-20		J2941		01/01/2002	03/31/2013	INJECTION, SOMATROPIN, 1 MG	NUTROPIN AQ (VIAL CARTON) 5 MG/ML	2	ML	VL	SC	ML	1	MG	5	01/01/2002	03/31/2013							
50242-0040-62		J2357		01/01/2005	99/99/9999	INJECTION, OMALIZUMAB, 5 MG	XOLAIR 150 MG	1	EA	VL	SC	EA	5	MG	30	01/01/2005	99/99/9999							
50242-0041-64		J2997		01/01/2002	99/99/9999	INJECTION, ALTEPLASE RECOMBINANT, 1 MG	CATHFLO ACTIVASE (VIAL) 2 MG	1	EA	VL	IV	EA	1	MG	2	01/01/2002	99/99/9999							
50242-0043-14		J2941		05/10/2002	12/31/2016	INJECTION, SOMATROPIN, 1 MG	NUTROPIN AQ PEN CARTRIDGE 5 MG/ML	2	ML	CT	SC	ML	1	MG	5	05/10/2002	12/31/2016							
50242-0044-13		J2997		01/01/2002	99/99/9999	INJECTION, ALTEPLASE RECOMBINANT, 1 MG	ACTIVASE (W/DILUENT) 50 MG	1	EA	VL	IV	EA	1	MG	50	01/01/2002	99/99/9999							
50242-0051-21		J9310		01/01/2002	12/31/2018	INJECTION, RITUXIMAB, 100 MG	RITUXAN (S.D.V.PF) 10 MG/ML	10	ML	VL	IV	ML	100	MG	0.1	01/01/2002	12/31/2018							
50242-0053-06		J9310		01/01/2002	12/31/2018	INJECTION, RITUXIMAB, 100 MG	RITUXAN (S.D.V.PF) 10 MG/ML	50	ML	VL	IV	ML	100	MG	0.1	01/01/2002	12/31/2018							
50242-0060-01		J9035		01/01/2005	99/99/9999	INJECTION, BEVACIZUMAB, 10 MG	AVASTIN (PF) 25 MG/ML	4	ML	VL	IV	ML	10	MG	2.5	01/01/2005	99/99/9999							
50242-0061-01		J9035		01/01/2005	99/99/9999	INJECTION, BEVACIZUMAB, 10 MG	AVASTIN (PF) 25 MG/ML	16	ML	VL	IV	ML	10	MG	2.5	01/01/2005	99/99/9999							
50242-0073-01		J2941		01/28/2008	07/31/2016	INJECTION, SOMATROPIN, 1 MG	NUTROPIN AQ PEN (1X2ML) 10 MG/ML	2	ML	CT	SC	ML	1	MG	10	01/28/2008	07/31/2016							
72205-0006-60		None		10/01/2018	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (FILM COATED) 150 MG	60	EA	BO	PO	EA	150	MG	1	10/01/2018	99/99/9999							
50242-0080-01		J2778		01/01/2008	99/99/9999	INJECTION, RANIBIZUMAB, 0.1 MG	LUCENTIS (INTRAVITREAL INJECTION) 0.5 MG/0.05 ML	0.05	ML	VL	IO	ML	0.1	MG	1	01/01/2008	99/99/9999							
50242-0085-27		J2997		01/01/2002	99/99/9999	INJECTION, ALTEPLASE RECOMBINANT, 1 MG	ACTIVASE (W/DILUENT) 100 MG	1	EA	VL	IV	EA	1	MG	100	01/01/2002	99/99/9999							
50242-0100-39		J7639		01/01/2002	99/99/9999	DORNASE ALPHA, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	PULMOZYME (AMP INNER NDC) 2.5 MG/2.5 ML	2.5	ML	PC	IH	ML	1	MG	1	01/01/2002	99/99/9999							
50242-0100-39	KO	J7639	KO	01/01/2002	99/99/9999	DORNASE ALPHA, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	PULMOZYME (AMP) 2.5 MG/2.5 ML	2.5	ML	PC	IH	ML	1	MG	1	01/01/2002	99/99/9999							
50242-0100-40		J7639		01/01/2002	99/99/9999	DORNASE ALPHA, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	PULMOZYME (AMP) 2.5 MG/2.5 ML	2.5	ML	PC	IH	ML	1	MG	1	01/01/2002	99/99/9999							
50242-0100-40	KO	J7639	KO	01/01/2002	99/99/9999	DORNASE ALPHA, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	PULMOZYME (AMP) 2.5 MG/2.5 ML	2.5	ML	PC	IH	ML	1	MG	1	01/01/2002	99/99/9999							
50242-0134-68		J9355		09/01/2003	99/99/9999	INJECTION, TRASTUZUMAB, 10 MG	HERCEPTIN (M.D.V.,W/DILUENT 20ML) 440 MG	1	EA	VL	IV	EA	10	MG	44	09/01/2003	99/99/9999							
50383-0040-04		J7510		01/22/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (AF,SF,DYE-FREE) 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	01/22/2003	99/99/9999							
50383-0042-24		J7510		03/24/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240	ML	BO	PO	ML	5	MG	0.6	03/24/2003	99/99/9999							
50383-0042-48		J7510		03/17/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	480	ML	BO	PO	ML	5	MG	0.6	03/17/2003	99/99/9999							
50383-0741-20		J7611		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE 0.5%	20	ML	BO	IH	ML	1	MG	5	04/01/2008	99/99/9999							
50383-0801-16		Q0170		03/01/2004	12/31/2019	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (CHERRY) 6.25 MG/5 ML	473	ML	BO	PO	ML	25	MG	0.05	03/01/2004	12/31/2013							
50383-0810-16		J8499		06/13/2005	07/31/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (BANANA) 200 MG/5 ML	473	ML	BO	PO	ML	1	EA	1	06/13/2005	07/31/2014							
50419-0150-57		J1945		01/01/2006	05/31/2012	INJECTION, LEPIDIRUDIN, 50 MG	REFLUDAN (VIAL) 50 MG	1	EA	VL	IV	EA	50	MG	1	01/01/2006	05/31/2012							
50419-0511-06		J9185		01/01/2002	06/30/2014	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARA 50 MG	1	EA	VL	IV	EA	50	MG	1	01/01/2002	06/30/2014							
50419-0523-25		J1830		01/02/2004	99/99/9999	FOR USE WHEN DRUG IS SELF ADMINISTERED)	BETASERON (15 BLISTER UNITS,PF) 0.3 MG-0.54%	15	EA	VL	MR	EA	0.25	MG	18	01/02/2004	99/99/9999							
50458-0306-11		J2794		01/01/2005	99/99/9999	INJECTION, RISPERIDONE, LONG ACTING, 0.5 MG	RISPERDAL CONSTA 25 MG	1	EA	VL	IM	EA	0.5	MG	50	01/01/2005	99/99/9999							
50458-0307-11		J2794		01/01/2005	99/99/9999	INJECTION, RISPERIDONE, LONG ACTING, 0.5 MG	RISPERDAL CONSTA 37.5 MG	1	EA	VL	IM	EA	0.5	MG	75	01/01/2005	99/99/9999							
50458-0308-11		J2794		01/01/2005	99/99/9999	INJECTION, RISPERIDONE, LONG ACTING, 0.5 MG	RISPERDAL CONSTA 50 MG	1	EA	VL	IM	EA	0.5	MG	100	01/01/2005	99/99/9999							
50458-0309-11		J2794		04/23/2007	99/99/9999	INJECTION, RISPERIDONE, LONG ACTING, 0.5 MG	RISPERDAL CONSTA 12.5 MG	1	EA	VL	IM	EA	0.5	MG	25	04/23/2007	99/99/9999							
50486-0078-22		A4216		01/01/2006	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	BRONCHO SALINE 0.9%	90	ML	BO	IH	ML	10	ML	0.1	01/01/2006	02/03/2016							
50486-0078-23		A4216		01/01/2006	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	BRONCHO SALINE 0.9																	

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
50486-0616-32		Q0163		12/04/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SLEEPINAL 50 MG	32	EA	NA	PO	EA	50	MG	1	12/04/2002	99/99/9999						
50962-0650-01		A4216		01/01/2006	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (INHALATION) 0.9%	1	ML	EA	IH	ML	10	ML	0.1	01/01/2006	99/99/9999						
51079-0066-01		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (USP) 50 MG	1	EA	BX	PO	EA	50	MG	1	01/01/2002	02/03/2016						
51079-0066-20		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (10X10) 50 MG	100	EA	BX	PO	EA	50	MG	1	01/01/2002	99/99/9999						
51079-0077-01		Q0177		11/26/2007	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (USP) 25 MG	1	EA	NA	PO	EA	25	MG	1	11/26/2007	99/99/9999						
51079-0077-20		Q0177		01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (10X10) 25 MG	100	EA	BX	PO	EA	25	MG	1	11/26/2007	99/99/9999	01/01/2002	04/01/2002	1			
51079-0078-01		Q0178		11/26/2007	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (USP) 50 MG	1	EA	NA	PO	EA	50	MG	1	11/26/2007	12/31/2013						
51079-0078-20		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (10X10) 50 MG	100	EA	BX	PO	EA	50	MG	1	11/26/2007	12/31/2013	01/01/2002	04/01/2002	1			
51079-0434-01		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (USP) 20 MG	1	EA	BX	PO	EA	1	EA	1	01/01/2002	99/99/9999						
51079-0434-20		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (10X10) 20 MG	100	EA	BX	PO	EA	1	EA	1	01/01/2002	99/99/9999						
51079-0435-01		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (USP) 40 MG	1	EA	BX	PO	EA	1	EA	1	01/01/2002	99/99/9999						
51079-0435-20		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (10X10) 40 MG	100	EA	BX	PO	EA	1	EA	1	01/01/2002	99/99/9999						
51079-0472-01		Q0166		03/03/2008	05/16/2012	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (FILM-COATED) 1 MG	1	EA	BX	PO	EA	1	MG	1	03/03/2008	05/16/2012						
51079-0472-05		Q0166		03/03/2008	05/16/2012	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (2CARDSX10,FILM-COATED) 1 MG	20	EA	BX	PO	EA	1	MG	1	03/03/2008	05/16/2012						
51079-0541-01		Q0164		01/01/2002	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (USP) 5 MG	1	EA	BX	PO	EA	5	MG	1	01/01/2002	99/99/9999						
51079-0541-20		Q0164		01/01/2002	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (10X10) 5 MG	100	EA	BX	PO	EA	5	MG	1	01/01/2002	99/99/9999						
51079-0542-01		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (USP) 10 MG	1	EA	BX	PO	EA	10	MG	1	01/01/2002	12/31/2013						
51079-0542-20		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (10X10) 10 MG	100	EA	BX	PO	EA	10	MG	1	01/01/2002	12/31/2013						
51079-0591-01		Q0144		06/25/2007	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	1	EA	BX	PO	EA	1	GM	0.25	06/25/2007	02/03/2016						
51079-0670-01		None		01/01/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM (USP) 2.5 MG	1	EA	BX	PO	EA	2.5	MG	1	01/01/1994	99/99/9999						
51079-0670-05		None		01/01/1994	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM (2X10) 2.5 MG	20	EA	BX	PO	EA	2.5	MG	1	01/01/1994	99/99/9999						
51079-0895-01		Q0170		02/01/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	1	EA	BX	PO	EA	25	MG	1	02/01/2007	12/31/2013						
51079-0895-20		Q0170		03/14/2005	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (10X10) 25 MG	100	EA	BX	PO	EA	25	MG	1	02/01/2007	12/31/2013	03/14/2005	05/24/2005	1			
51079-0967-01		Q0163		01/01/2002	05/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (MINITAB,MINITAB) 25 MG	1	EA	BX	PO	EA	50	MG	0.5	01/01/2002	05/16/2012						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51079-0967-17		Q0163		01/01/2002	05/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (MINITAB,MINITAB) 25 MG	1	EA	BX	PO	EA	50	MG	0.5	01/01/2002	05/16/2012						
51079-0967-19		Q0163		01/01/2002	05/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (MINITAB) 25 MG	25	EA	BX	PO	EA	50	MG	0.5	01/01/2002	05/16/2012						
51079-0967-20		Q0163		01/01/2002	05/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (10X10,MINITAB) 25 MG	100	EA	BX	PO	EA	50	MG	0.5	01/01/2002	05/16/2012						
51285-0366-01	None			03/09/2006	99/99/9999	METHOTREXATE, 5 MG	TREXALL (FILM-COATED) 5 MG	30	EA	BO	PO	EA	5	MG	1	03/09/2006	99/99/9999						
51285-0367-01	None			03/09/2006	99/99/9999	METHOTREXATE, 7.5 MG	TREXALL (FILM-COATED) 7.5 MG	30	EA	BO	PO	EA	7.5	MG	1	03/09/2006	99/99/9999						
51285-0368-01	None			12/01/2005	99/99/9999	METHOTREXATE, 10 MG	TREXALL (FILM-COATED) 10 MG	30	EA	BO	PO	EA	10	MG	1	12/01/2005	99/99/9999						
51285-0369-01	None			12/01/2005	99/99/9999	METHOTREXATE, 15 MG	TREXALL (FILM-COATED) 15 MG	30	EA	BO	PO	EA	15	MG	1	12/01/2005	99/99/9999						
51552-0005-01	J2675			09/01/2003	01/01/2015	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	01/01/2015						
51552-0005-04	J2675			09/01/2003	01/01/2015	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	01/01/2015						
51552-0005-07	J2675			09/01/2003	01/01/2015	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.)	1	EA	JR	NA	GM	50	MG	20	09/01/2003	01/01/2015						
51552-0006-01	J2675			09/01/2003	01/01/2015	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	01/01/2015						
51552-0006-03	J2675			09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE,U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0006-04	J2675			09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE,U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0006-05	J2675			09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE,U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0006-07	J2675			09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE,U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0021-01	J1700			01/01/2002	01/01/2015	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	25	MG	40	01/01/2002	01/01/2015						
51552-0021-02	J1700			09/01/2003	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	25	MG	40	09/01/2003	99/99/9999						
51552-0021-03	J1700			09/01/2003	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	25	MG	40	09/01/2003	99/99/9999						
51552-0021-04	J1700			09/01/2003	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	25	MG	40	09/01/2003	99/99/9999						
51552-0021-05	J1700			09/01/2003	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	25	MG	40	09/01/2003	99/99/9999						
51552-0024-01	J1094			01/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2003	99/99/9999						
51552-0024-02	J1094			09/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0024-03	J1094			09/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0024-04	J1094			09/01/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0025-01	J7638			01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
51552-0025-02	KO	J7638	KO	01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
51552-0025-03	J7638			09/01/2003	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0025-04	KO	J7638	KO	09/01/2003	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0025-05	KO	J7638	KO	09/01/2003	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0026-02	J7510			09/01/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	99/99/9999						
51552-0026-04	J7510			09/01/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	99/99/9999						
51552-0026-05	J7510			09/01/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	99/99/9999						
51552-0028-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE	1	EA	BO	NA	GM	5	MG	200	01/01/2002	12/31/2015						
51552-0028-02	J7506			09/01/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	12/31/2015						
51552-0028-04	J7506			09/01/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	12/31/2015						
51552-0028-05	J7506			09/01/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	12/31/2015						
51552-0029-01	J3140			01/01/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P.)	1	EA	BO	NA	GM	50	MG	20	01/01/2002	12/31/2014						
51552-0029-02	J3140			09/01/2003	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P.)	1	EA	JR	NA	GM	50	MG	20	09/01/2003	12/31/2014						
51552-0029-04	J3140			09/01/2003	07/30/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	07/30/2013						
51552-0029-07	J3140			09/01/2003	07/30/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	07/30/2013						
51552-0030-01	J3150			01/01/2002	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	01/01/2002	12/31/2014						
51552-0030-02	J3150			09/01/2003	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	09/01/2003	12/31/2014						
51552-0030-04	J3150			09/01/2003	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	09/01/2003	12/31/2014						
51552-0030-05	J3150			09/01/2003	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	100	MG	10	09/01/2003	12/31/2014						
51552-0030-08	J3150			09/01/2003	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	100	MG	10	09/01/2003	12/31/2014						
51552-0030-09	J3150			09/01/2003	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	1	EA	BO	NA	GM	100	MG	10	09/01/2003	12/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51552-0033-01		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
51552-0033-01	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
51552-0033-02		J7684		09/01/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0033-02	KO	J7684	KO	09/01/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0033-03		J7684		09/01/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0033-03	KO	J7684	KO	09/01/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0033-05		J7684		09/01/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0033-05	KO	J7684	KO	09/01/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0038-03		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	99/99/9999						
51552-0038-04		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	99/99/9999						
51552-0038-05		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	JR	NA	GM		1 EA		1	09/01/2003	99/99/9999						
51552-0038-06		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	99/99/9999						
51552-0042-01		J7643		01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
51552-0042-01	KO	J7643	KO	01/01/2002	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
51552-0044-02		J7609		01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-02	KO	J7609	KO	01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-04		J7609		01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-04	KO	J7609	KO	01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-05		J7609		01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-05	KO	J7609	KO	01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-06		J7609		01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-06	KO	J7609	KO	01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-07		J7609		01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0044-07	KO	J7609	KO	01/01/2007	01/01/2015	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2007	01/01/2015						
51552-0057-04		J3350		01/01/2002	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P.,N.F.)	1 EA	BO	NA	GM		40 GM		0.025	01/01/2002	99/99/9999						
51552-0057-04	KO	J3350	KO	09/01/2003	10/17/2016	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P.,N.F.)	1 EA	BO	NA	GM		40 GM		0.025	09/01/2003	10/17/2016						
51552-0057-08		J3350		09/01/2003	10/17/2016	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P.,N.F.)	1 EA	BO	NA	GM		40 GM		0.025	09/01/2003	10/17/2016						
51552-0061-06		J3480		09/01/2003	01/01/2015	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.,N.F.)	1 EA	BO	NA	GM		2 MEQ		6.71141	09/01/2003	01/01/2015						
51552-0064-01		J7624		01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
51552-0064-01	KO	J7624	KO	01/01/2002	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	01/01/2002	99/99/9999						
51552-0064-02		J7624		09/01/2003	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0064-02	KO	J7624	KO	09/01/2003	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0074-05		Q0165		09/01/2003	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	1 EA	BO	NA	GM		10 MG		100	09/01/2003	12/31/2013						
51552-0074-05	KO	Q0165	KO	09/01/2003	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	1 EA	BO	NA	GM		10 MG		100	09/01/2003	12/31/2013						
51552-0079-02		J7670		01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		10 MG		100	01/01/2007	01/01/2015						
51552-0079-02	KO	J7670	KO	01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		10 MG		100	01/01/2007	01/01/2015						
51552-0079-04		J7670		01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		10 MG		100	01/01/2007	01/01/2015						
51552-0079-04	KO	J7670	KO	01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		10 MG		100	01/01/2007	01/01/2015						
51552-0079-05		J7670		01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.,N.F.)	1 EA	BO	NA	GM		10 MG		100	01/01/2007	01/01/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51552-0079-05	KO	J7670	KO	01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	1	EA	BO	NA	GM	10	MG	100	01/01/2007	01/01/2015						
51552-0079-07		J7670		01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	1	EA	BO	NA	GM	10	MG	100	01/01/2007	01/01/2015						
51552-0079-07	KO	J7670	KO	01/01/2007	01/01/2015	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	1	EA	BO	NA	GM	10	MG	100	01/01/2007	01/01/2015						
51552-0106-04		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	1	EA	BO	NA	GM	10	MG	100	01/01/2004	99/99/9999						
51552-0106-05		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	1	EA	JR	NA	GM	10	MG	100	01/01/2004	99/99/9999						
51552-0106-06		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	1	EA	BO	NA	GM	10	MG	100	01/01/2004	99/99/9999						
51552-0124-02		J1200		09/01/2003	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0124-05		J1200		09/01/2003	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0124-06		J1200		09/01/2003	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0130-02		J3490		09/01/2003	07/30/2013	UNCLASSIFIED DRUGS	1	EA	BO	NA	GM	1	EA	1	09/01/2003	07/30/2013						
51552-0130-04		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	1	EA	BO	NA	GM	1	EA	1	01/01/2002	99/99/9999						
51552-0130-06		J3490		09/01/2003	07/30/2013	UNCLASSIFIED DRUGS	1	EA	BO	NA	GM	1	EA	1	09/01/2003	07/30/2013						
51552-0139-04		J3230		09/01/2003	99/99/9999	INJECTION, CHLORPROMAZINE HCL, UP TO 50 MG	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0139-05		J3230		09/01/2003	99/99/9999	INJECTION, CHLORPROMAZINE HCL, UP TO 50 MG	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0139-07		J3230		09/01/2003	01/01/2015	INJECTION, CHLORPROMAZINE HCL, UP TO 50 MG	1	EA	BO	NA	GM	50	MG	20	09/01/2003	01/01/2015						
51552-0141-02		J1980		09/01/2003	01/01/2015	INJECTION, HYOSCYAMINE SULFATE, UP TO 0.25 MG	1	EA	BO	NA	GM	0.25	MG	4000	09/01/2003	01/01/2015						
51552-0141-04		J1980		09/01/2003	01/01/2015	INJECTION, HYOSCYAMINE SULFATE, UP TO 0.25 MG	1	EA	BO	NA	GM	0.25	MG	4000	09/01/2003	01/01/2015						
51552-0147-01		J2550		01/01/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	1	EA	JR	NA	GM	50	MG	20	01/01/2002	99/99/9999						
51552-0147-02		J2550		09/01/2003	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0147-04		J2550		09/01/2003	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0147-05		J2550		09/01/2003	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0149-04		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	1	EA	JR	NA	GM	100	MG	10	01/01/2004	99/99/9999						
51552-0149-05		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	1	EA	BO	NA	GM	100	MG	10	01/01/2004	99/99/9999						
51552-0156-02		J7636		09/01/2003	99/99/9999	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0156-02	KO	J7636	KO	09/01/2003	99/99/9999	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0156-04		J7636		09/01/2003	99/99/9999	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0156-04	KO	J7636	KO	09/01/2003	99/99/9999	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0180-03		J2765		09/01/2003	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	1	EA	BO	NA	GM	10	MG	100	09/01/2003	99/99/9999						
51552-0180-04		J2765		09/01/2003	10/03/2017	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	1	EA	BO	NA	GM	10	MG	100	09/01/2003	10/03/2017						
51552-0180-05		J2765		09/01/2003	10/03/2017	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	1	EA	BO	NA	GM	10	MG	100	09/01/2003	10/03/2017						
51552-0188-01		J1330		01/01/2002	01/01/2015	INJECTION, ERGONOVINE MALEATE, UP TO 0.2 MG	1	EA	BO	NA	GM	0.2	MG	5000	01/01/2002	01/01/2015						
51552-0188-05		J1330		09/01/2003	01/01/2015	INJECTION, ERGONOVINE MALEATE, UP TO 0.2 MG	1	EA	VL	NA	GM	0.2	MG	5000	09/01/2003	01/01/2015						
51552-0188-07		J1330		09/01/2003	01/01/2015	INJECTION, ERGONOVINE MALEATE, UP TO 0.2 MG	1	EA	BO	NA	GM	0.2	MG	5000	09/01/2003	01/01/2015						
51552-0201-04		J7604		01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51552-0201-04	KO	J7604	KO	01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51552-0201-05		J7604		01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51552-0201-05	KO	J7604	KO	01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51552-0201-07		J7604		01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51552-0201-07	KO	J7604	KO	01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51552-0232-04		J7799		09/01/2003	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	1	EA	BO	NA	GM	1	EA	1	09/01/2003	99/99/9999						
51552-0232-04		J7799		09/01/2003	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	1	EA	BO	NA	GM	1	EA	1	09/01/2003	99/99/9999						
51552-0232-05		J7799		09/01/2003	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	1	EA	BO	NA	GM	1	EA	1	09/01/2003	99/99/9999						
51552-0233-01		J1110		01/01/2002	99/99/9999	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
51552-0233-02		J1110		09/01/2003	99/99/9999	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0278-01		J3302		01/01/2002	01/01/2015	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	1	EA	BO	NA	GM	5	MG	200	01/01/2002	01/01/2015						
51552-0278-02		J3302		09/01/2003	01/01/2015	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	1	EA	BO	NA	GM	5	MG	200	09/01/2003	01/01/2015						
51552-0278-03		J3302		09/01/2003	01/01/2015	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	1	EA	BO	NA	GM	5	MG	200	09/01/2003	01/01/2015						
51552-0304-01		J0285		09/01/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0304-02		J0285		09/01/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51552-0304-03		J0285		09/01/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
51552-0304-04		J0285		09/01/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (1X25GM)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999							
51552-0304-05		J0285		01/01/2002	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999	01/01/2002	08/31/2003	20				
51552-0304-06		J0285		09/01/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (1X500GM)	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999							
51552-0304-07		J0285		09/01/2003	01/01/2015	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1	EA	JR	NA	GM	50	MG	20	09/01/2003	01/01/2015							
51552-0304-09		J0285		09/01/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B	1	EA	JR	NA	GM	50	MG	20	09/01/2003	99/99/9999							
51552-0313-05		J0280		09/01/2003	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1	EA	JR	NA	GM	250	MG	4	09/01/2003	99/99/9999							
51552-0313-06		J0280		09/01/2003	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	250	MG	4	09/01/2003	99/99/9999							
51552-0324-06		J3480		09/01/2003	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	2	MEQ	6.71141	09/01/2003	10/17/2016							
51552-0324-08		J3480		09/01/2003	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	2	MEQ	6.71141	09/01/2003	10/17/2016							
51552-0324-09		J3480		09/01/2003	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	2	MEQ	6.71141	09/01/2003	99/99/9999							
51552-0380-01		J2150		09/01/2003	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P., N.F.)	1	EA	BO	NA	GM	50	ML	0.08	09/01/2003	99/99/9999							
51552-0380-05		J2150		09/01/2003	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P., N.F.)	1	EA	BO	NA	GM	50	ML	0.08	09/01/2003	99/99/9999							
51552-0380-06		J2150		09/01/2003	10/17/2016	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P., N.F.)	1	EA	BO	NA	GM	50	ML	0.08	09/01/2003	10/17/2016							
51552-0380-08		J2150		09/01/2003	10/17/2016	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P., N.F.)	1	EA	BO	NA	GM	50	ML	0.08	09/01/2003	10/17/2016							
51552-0380-09		J2150		09/01/2003	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	1	EA	BO	NA	GM	50	ML	0.08	09/01/2003	99/99/9999							
51552-0393-01		J7645		01/01/2007	01/01/2015	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0393-01	KO	J7645	KO	01/01/2007	01/01/2015	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0393-02		J7645		01/01/2007	01/01/2015	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0393-02	KO	J7645	KO	01/01/2007	01/01/2015	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0393-04		J7645		01/01/2007	01/01/2015	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0393-04	KO	J7645	KO	01/01/2007	01/01/2015	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0393-05		J7645		01/01/2007	01/01/2015	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0393-05	KO	J7645	KO	01/01/2007	01/01/2015	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (B.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	01/01/2015							
51552-0416-02		J2440		09/01/2003	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1	EA	BO	NA	GM	60	MG	16.66666	09/01/2003	99/99/9999							
51552-0416-04		J2440		09/01/2003	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1	EA	BO	NA	GM	60	MG	16.66666	09/01/2003	99/99/9999							
51552-0416-05		J2440		09/01/2003	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1	EA	BO	NA	GM	60	MG	16.66666	09/01/2003	99/99/9999							
51552-0416-07		J2440		09/01/2003	01/01/2015	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1	EA	BO	NA	GM	60	MG	16.66666	09/01/2003	01/01/2015							
51552-0423-02		J7632		01/01/2008	99/99/9999	CROMOLYN SODIUM, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
51552-0423-02	KO	J7632	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
51552-0423-04		J7632		01/01/2008	99/99/9999	CROMOLYN SODIUM, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
51552-0423-04	KO	J7632	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
51552-0423-05		J7632		01/01/2008	99/99/9999	CROMOLYN SODIUM, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
51552-0423-05	KO	J7632	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
51552-0423-07		J7632		01/01/2008	01/01/2015	CROMOLYN SODIUM, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	01/01/2015							
51552-0423-07	KO	J7632	KO	01/01/2008	01/01/2015	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	01/01/2015							
51552-0430-01		J7638		01/01/2002	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
51552-0430-01	KO	J7638	KO	01/01/2002	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
51552-0430-02		J7638		09/01/2003	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999							
51552-0430-02	KO	J7638	KO	09/01/2003	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999							
51552-0435-05		J0600		09/01/2003	01/01/2015	INJECTION, EDETATE CALCIUM DISODIUM, UP TO 1000 MG	EDETATE CALCIUM DISODIUM (U.S.P., F.C.C.)	1	EA	BO	NA	GM	1000	MG	1	09/01/2003	01/01/2015							
51552-0445-01		J1435		01/01/2002	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
51552-0445-02		J1435		09/01/2003	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999							
51552-0445-04		J1435		09/01/2003	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999							
51552-0446-03		J7681		09/01/2003	01/01/2015	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P., NF)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	01/01/2015							
51552-0446-03	KO	J7681	KO	09/01/2003	01/01/2015	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P., NF)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	01/01/2015							
51552-0446-04		J7681		09/01/2003	01/01/2015	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	01/01/2015							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51552-0446-04	KO	J7681	KO	09/01/2003	01/01/2015	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		09/01/2003	01/01/2015							
51552-0464-02		J1320		09/01/2003	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (1X5GM)	1 EA	BO	NA	GM	20 MG	50		09/01/2003	99/99/9999							
51552-0464-05		J1320		09/01/2003	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (1X100GM)	1 EA	BO	NA	GM	20 MG	50		09/01/2003	99/99/9999							
51552-0464-06		J1320		09/01/2003	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (1X500GM)	1 EA	JR	NA	GM	20 MG	50		09/01/2003	99/99/9999							
51552-0480-01		J0735		01/01/2002	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	99/99/9999							
51552-0480-02		J0735		09/01/2003	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 EA	JR	NA	GM	1 MG	1000		09/01/2003	99/99/9999							
51552-0487-05		J2810		09/01/2003	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE ANHYDROUS (U.S.P.)	1 EA	BO	NA	GM	40 MG	25		09/01/2003	99/99/9999							
51552-0496-01		J2760		01/01/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	99/99/9999							
51552-0496-02		J2760		09/01/2003	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		09/01/2003	99/99/9999							
51552-0496-04		J2760		09/01/2003	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		09/01/2003	99/99/9999							
51552-0496-05		J2760		09/01/2003	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		09/01/2003	99/99/9999							
51552-0496-09		J2760		09/01/2003	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		09/01/2003	99/99/9999							
51552-0498-01		J0270		09/01/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	PROSTAGLANDIN E1 (1X1MG,USP)	1 EA	BO	NA	GM	1.25 MCG	800000		09/01/2003	99/99/9999							
51552-0498-03		J0270		09/01/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	PROSTAGLANDIN E1 (U.S.P.)	1 EA	BO	NA	GM	1.25 MCG	800000		09/01/2003	99/99/9999							
51552-0498-05		J0270		09/01/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	PROSTAGLANDIN E1 (1X100MG,USP)	1 EA	BO	NA	GM	1.25 MCG	800000		09/01/2003	99/99/9999							
51552-0498-09		J0270		09/01/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	PROSTAGLANDIN E1 (1X5MG,USP)	1 EA	BO	NA	GM	1.25 MCG	800000		09/01/2003	99/99/9999							
51552-0519-01		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2002	99/99/9999							
51552-0519-02		J1630		09/01/2003	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		09/01/2003	99/99/9999							
51552-0526-05		J7799		09/01/2003	01/01/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	EPINEPHRINE (U.S.P., N.F.)	1 EA	BO	NA	GM	1 EA	1		09/01/2003	01/01/2015							
51552-0529-02		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (U.S.P., N.F.)	1 EA	BO	NA	GM	1 EA	1		09/01/2003	99/99/9999							
51552-0529-03		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (U.S.P., N.F.)	1 EA	BO	NA	GM	1 EA	1		09/01/2003	99/99/9999							
51552-0532-04		J1165		09/01/2003	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM	1 EA	JR	NA	GM	50 MG	20		09/01/2003	99/99/9999							
51552-0564-04		J3140		09/01/2003	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P.)	1 EA	JR	NA	GM	50 MG	20		09/01/2003	12/31/2014							
51552-0564-05		J3140		09/01/2003	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	50 MG	20		09/01/2003	12/31/2014							
51552-0564-07		J3140		09/01/2003	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE (U.S.P.)	1 EA	BO	NA	GM	50 MG	20		09/01/2003	12/31/2014							
51552-0588-06		J3520		09/01/2003	99/99/9999	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM (U.S.P.)	1 EA	BO	NA	GM	150 MG	6.66666		09/01/2003	99/99/9999							
51552-0603-02		J7509		09/01/2003	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	4 MG	250		09/01/2003	99/99/9999							
51552-0611-01		J7641		01/01/2002	01/01/2015	FLUNISOLIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, PER MILLIGRAM	FLUNISOLIDE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	01/01/2015							
51552-0611-01	KO	J7641	KO	01/01/2002	01/01/2015	FLUNISOLIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, PER MILLIGRAM	FLUNISOLIDE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		01/01/2002	01/01/2015							
51552-0611-02		J7641		09/01/2003	01/01/2015	FLUNISOLIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, PER MILLIGRAM	FLUNISOLIDE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		09/01/2003	01/01/2015							
51552-0611-02	KO	J7641	KO	09/01/2003	01/01/2015	FLUNISOLIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, PER MILLIGRAM	FLUNISOLIDE ANHYDROUS (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000		09/01/2003	01/01/2015							
51552-0613-02		J0475		09/01/2003	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (1X5GM)	1 EA	JR	NA	GM	10 MG	100		09/01/2003	99/99/9999							
51552-0613-04		J0475		09/01/2003	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (1X25GM)	1 EA	JR	NA	GM	10 MG	100		09/01/2003	99/99/9999							
51552-0613-05		J0475		09/01/2003	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (1X100GM)	1 EA	JR	NA	GM	10 MG	100		09/01/2003	99/99/9999							
51552-0620-02		J2780		09/01/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO	NA	GM	25 MG	40		09/01/2003	99/99/9999							
51552-0620-04		J2780		09/01/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO	NA	GM	25 MG	40		09/01/2003	99/99/9999							
51552-0620-05		J2780		09/01/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO	NA	GM	25 MG	40		09/01/2003	99/99/9999							
51552-0628-01		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	1 EA	1		01/01/2002	99/99/9999							
51552-0643-07		J2675		09/01/2003	01/01/2015	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (MILLED,U.S.P.)	1 EA	BO	NA	GM	50 MG	20		09/01/2003	01/01/2015							
51552-0652-01		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (1X1GM)	1 EA	BO	NA	GM	1 MG	1000		01/01/2007	99/99/9999							
51552-0652-02		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (1X5GM)	1 EA	BO	NA	GM	1 MG	1000		01/01/2007	99/99/9999							
51552-0652-04		J0364		01/01/2007	01/01/2015	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		01/01/2007	01/01/2015							
51552-0663-01		J7516		01/01/2002	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORIN A	1 EA	BO	NA	GM	250 MG	4		01/01/2002	99/99/9999							
51552-0663-02		J7516		09/01/2003	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORINE (1X5GM USP)	1 EA	BO	NA	GM	250 MG	4		09/01/2003	99/99/9999							
51552-0663-04		J7516		09/01/2003	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORINE (1X25GM,USP)	1 EA	BO	NA	GM	250 MG	4		09/01/2003	99/99/9999							
51552-0663-06		J7516		09/01/2003	01/01/2015	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	CYCLOSPORINE (1X500MG,USP)	1 EA	BO	NA	GM	250 MG	4		09/01/2003	01/01/2015							
51552-0668-01		J7627		01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	JR	NA	GM	0.5 MG	2000		01/01/2006	99/99/9999							
51552-0668-01	KO	J7627	KO	01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED)	1 EA	JR	NA	GM	0.5 MG	2000		01/01/2006	99/99/9999							
51552-0671-01		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		01/01/2006	99/99/9999							
51552-0671-02		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO														

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51552-0671-05		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO NA GM	5 MG	200		5		200	01/01/2006	99/99/9999						
51552-0671-06		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO NA GM	5 MG	200		5		200	01/01/2006	99/99/9999						
51552-0674-05		J2010		09/01/2003	01/01/2015	INJECTION, LINCOCYCIN HCL, UP TO 300 MG	LINCOCYCIN HYDROCHLORIDE (USP,1X100GM)	1 EA	BO NA GM	300 MG	3.33333		3.33333		3.33333	09/01/2003	01/01/2015						
51552-0674-07		J2010		09/01/2003	01/01/2015	INJECTION, LINCOCYCIN HCL, UP TO 300 MG	LINCOCYCIN HYDROCHLORIDE (USP,1X1000GM)	1 EA	BO NA GM	300 MG	3.33333		3.33333		3.33333	09/01/2003	01/01/2015						
51552-0676-04		J1240		09/01/2003	99/99/9999	INJECTION, DIMENHYDRINATE, UP TO 50 MG	DIMENHYDRINATE (1X25GM,USP)	1 EA	BO NA GM	50 MG	20		20		20	09/01/2003	99/99/9999						
51552-0676-05		J1240		09/01/2003	99/99/9999	INJECTION, DIMENHYDRINATE, UP TO 50 MG	DIMENHYDRINATE (1X100GM,USP)	1 EA	BO NA GM	50 MG	20		20		20	09/01/2003	99/99/9999						
51552-0678-02		J2271		09/01/2003	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (1X5GM,USP)	1 EA	NA NA GM	100 MG	10		10		10	09/01/2003	12/31/2014						
51552-0678-04		J2271		09/01/2003	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (1X25GM,USP)	1 EA	JR NA GM	100 MG	10		10		10	09/01/2003	12/31/2014						
51552-0678-06		J2271		09/01/2003	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (1X100GM,USP)	1 EA	JR NA GM	100 MG	10		10		10	09/01/2003	12/31/2014						
51552-0682-01		J1170		09/01/2003	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (1X1GM,USP)	1 EA	BO NA GM	4 MG	250		250		250	09/01/2003	99/99/9999						
51552-0682-02		J1170		09/01/2003	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (1X5GM,USP)	1 EA	BO NA GM	4 MG	250		250		250	09/01/2003	99/99/9999						
51552-0682-03		J1170		09/01/2003	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (1X10GM,USP)	1 EA	BO NA GM	4 MG	250		250		250	09/01/2003	99/99/9999						
51552-0682-04		J1170		09/01/2003	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (1X25GM,USP)	1 EA	BO NA GM	4 MG	250		250		250	09/01/2003	99/99/9999						
51552-0686-01		J2175		09/01/2003	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HYDROCHLORIDE (USP,1X1GM)	1 EA	BO NA GM	100 MG	10		10		10	09/01/2003	99/99/9999						
51552-0686-02		J2175		09/01/2003	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HYDROCHLORIDE (USP,1X5GM)	1 EA	BO NA GM	100 MG	10		10		10	09/01/2003	99/99/9999						
51552-0686-04		J2175		09/01/2003	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HYDROCHLORIDE (USP,1X25GM)	1 EA	BO NA GM	100 MG	10		10		10	09/01/2003	99/99/9999						
51552-0686-06		J2175		09/01/2003	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HYDROCHLORIDE (USP,1X100GM)	1 EA	BO NA GM	100 MG	10		10		10	09/01/2003	99/99/9999						
51552-0687-01		J3010		09/01/2003	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (1X1GM,USP)	1 EA	BO NA GM	0.1 MG	10000		10000		10000	09/01/2003	99/99/9999						
51552-0687-09		J3010		09/01/2003	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (1X500MG,USP)	500 ML	BO NA ML	0.1 MG	10000		10000		10000	09/01/2003	99/99/9999						
51552-0688-02		J0745		09/01/2003	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (1X5GM,USP)	1 EA	BO NA GM	30 MG	33.33333		33.33333		33.33333	09/01/2003	99/99/9999						
51552-0688-03		J0745		09/01/2003	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (1X10GM,USP)	1 EA	BO NA GM	30 MG	33.33333		33.33333		33.33333	09/01/2003	99/99/9999						
51552-0688-04		J0745		09/01/2003	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (1X25GM,USP)	1 EA	BO NA GM	30 MG	33.33333		33.33333		33.33333	09/01/2003	99/99/9999						
51552-0688-06		J0745		09/01/2003	01/01/2015	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (1X100GM,USP)	1 EA	BO NA GM	30 MG	33.33333		33.33333		33.33333	09/01/2003	01/01/2015						
51552-0701-02		J2710		09/01/2003	01/01/2015	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE	1 EA	BO NA GM	0.5 MG	2000		2000		2000	09/01/2003	01/01/2015						
51552-0715-04		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	RIFAMPIN (USP,1X25GM)	1 EA	BO NA GM	1 EA	1		1		1	09/01/2003	99/99/9999						
51552-0715-05		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	RIFAMPIN (USP,1X100GM)	1 EA	BO NA GM	1 EA	1		1		1	09/01/2003	99/99/9999						
51552-0715-06		J3490		09/01/2003	01/01/2015	UNCLASSIFIED DRUGS	RIFAMPIN (USP,1X500GM)	1 EA	BO NA GM	1 EA	1		1		1	09/01/2003	01/01/2015						
51552-0728-01		J1230		09/01/2003	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	BO NA GM	10 MG	100		100		100	09/01/2003	99/99/9999						
51552-0728-02		J1230		09/01/2003	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	BO NA GM	10 MG	100		100		100	09/01/2003	99/99/9999						
51552-0728-04		J1230		09/01/2004	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	JR NA GM	10 MG	100		100		100	09/01/2004	99/99/9999						
51552-0729-01		J2060		09/01/2003	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (1X1GM,USP)	1 EA	BO NA GM	2 MG	500		500		500	09/01/2003	99/99/9999						
51552-0729-02		J2060		09/01/2003	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (1X5GM,USP)	1 EA	BO NA GM	2 MG	500		500		500	09/01/2003	99/99/9999						
51552-0729-04		J2060		09/01/2003	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (1X25GM,USP)	1 EA	BO NA GM	2 MG	500		500		500	09/01/2003	99/99/9999						
51552-0729-05		J2060		09/01/2003	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (1X100GM,USP)	1 EA	NA NA GM	2 MG	500		500		500	09/01/2003	99/99/9999						
51552-0729-09		J2060		09/01/2003	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (1X500MG,USP)	1 EA	BO NA GM	2 MG	500		500		500	09/01/2003	99/99/9999						
51552-0733-01		J9190		09/01/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (1X1GM,USP)	1 EA	BO NA GM	500 MG	2		2		2	09/01/2003	99/99/9999						
51552-0733-02		J9190		09/01/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (1X5GM,USP)	1 EA	BO NA GM	500 MG	2		2		2	09/01/2003	99/99/9999						
51552-0733-04		J9190		09/01/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (1X25GM,USP)	1 EA	BO NA GM	500 MG	2		2		2	09/01/2003	99/99/9999						
51552-0733-05		J9190		09/01/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (1X100GM,USP)	1 EA	BO NA GM	500 MG	2		2		2	09/01/2003	99/99/9999						
51552-0737-01		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	NALTREXONE HYDROCHLORIDE (1X1GM,USP)	1 EA	JR NA GM	1 EA	1		1		1	09/01/2003	99/99/9999						
51552-0737-02		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	NALTREXONE HYDROCHLORIDE (1X5GM,USP)	1 EA	BO NA GM	1 EA	1		1		1	09/01/2003	99/99/9999						
51552-0738-04		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X25GM,USP, MICRONIZED)	1 EA	BO NA GM	50 MG	20		20		20	09/01/2003	99/99/9999						
51552-0738-05		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X100GM,USP, MICRONIZED)	1 EA	JR NA GM	50 MG	20		20		20	09/01/2003	99/99/9999						
51552-0738-06		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X500GM,USP, MICRONIZED)	1 EA	BO NA GM	50 MG	20		20		20	09/01/2003	99/99/9999						
51552-0738-07		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X1000GM,USP, MICRONIZED)	1 EA	BO NA GM	50 MG	20		20		20	09/01/2003	99/99/9999						
51552-0741-04		J0500		09/01/2003	99/99/9999	INJECTION, DICYCLIMINE HYDROCHLORIDE (USP)	DICYCLIMINE HYDROCHLORIDE (USP)	1 EA	BO NA GM	20 MG	50		50		50	09/01/2003	99/99/9999						
51552-0763-05		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	6-AMINOCAPROIC ACID (1X100GM)	1 EA	BO NA GM	1 EA	1		1		1	09/01/2003	99/99/9999						
51552-0763-07		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	6-AMINOCAPROIC ACID (1X1000GM)	1 EA	BO NA GM	1 EA	1		1		1	09/01/2003	99/99/9999						
51552-0768-01		J7684		09/01/2003	99/99/9999	INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE (1X1GM,USP)	1 EA	BO NA GM	1 MG	1000		1000		1000	09/01/2003	99/99/9999						
51552-0768-01	KO	J7684	KO	09/01/2003	99																		

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51552-0775-05		J7699		09/01/2003	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMYCIN SULFATE (1X100MG,USP)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	99/99/9999						
51552-0779-02		J7501		09/01/2003	99/99/9999	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (1X5GM)	1 EA	BO	NA	GM		100 MG		10	09/01/2003	99/99/9999						
51552-0779-04		J7501		09/01/2003	99/99/9999	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (1X25GM)	1 EA	BO	NA	GM		100 MG		10	09/01/2003	99/99/9999						
51552-0779-05		J7501		09/01/2003	01/01/2015	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (1X100GM)	1 EA	BO	NA	GM		100 MG		10	09/01/2003	01/01/2015						
51552-0789-01		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X1GM,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0789-01	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X1GM,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0789-02		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X5GM,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0789-02	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X5GM,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0789-04		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X25GM,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0789-04	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X25GM,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0789-05		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X100MG,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0789-05	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE (1X100MG,USP)	1 EA	BO	NA	GM		300 MG	3.33333		01/01/2007	99/99/9999						
51552-0802-02		J0360		09/01/2003	01/01/2015	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HCL (U.S.P.)	1 EA	BO	NA	GM		20 MG		50	09/01/2003	01/01/2015						
51552-0829-01		J2675		09/01/2003	01/01/2015	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X1GM,USP)	1 EA	NA	NA	GM		50 MG		20	09/01/2003	01/01/2015						
51552-0829-03		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X10GM,USP)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0829-04		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X25GM,USP)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0829-05		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X100GM,USP)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0829-06		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X500GM,USP)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0829-07		J2675		09/01/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X1000GM,USP)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0829-08		J2675		09/01/2003	01/01/2015	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (1X5000GM,USP)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	01/01/2015						
51552-0839-05		J2360		09/01/2003	01/01/2015	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORPHENADRINE CITRATE (U.S.P.)	1 EA	BO	NA	GM		60 MG	16.66666		09/01/2003	01/01/2015						
51552-0879-02		J0520		09/01/2003	99/99/9999	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (1X5GM,USP)	1 EA	JR	NA	GM		5 MG		200	09/01/2003	99/99/9999						
51552-0879-04		J0520		09/01/2003	99/99/9999	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (1X25GM,USP)	1 EA	JR	NA	GM		5 MG		200	09/01/2003	99/99/9999						
51552-0883-01		J7622		09/01/2003	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (1X1GM,USP)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0883-01	KO	J7622	KO	09/01/2003	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (1X1GM,USP)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0883-02		J7622		09/01/2003	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (1X5GM,USP)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0883-02	KO	J7622	KO	09/01/2003	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (1X5GM,USP)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0883-09		J7622		09/01/2003	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (1X250MG,USP)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0883-09	KO	J7622	KO	09/01/2003	99/99/9999	BECLOMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLOMETHASONE DIPROPIONATE (1X250MG,USP)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0889-02		J3490		09/01/2003	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (1X10MG,USP)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	99/99/9999						
51552-0889-03		J3490		09/01/2003	01/01/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (1X50MG,USP)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	01/01/2015						
51552-0889-04		J3490		09/01/2003	01/01/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (1X100MG,USP)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	01/01/2015						
51552-0889-09		J3490		09/01/2003	01/01/2015	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (1X500MG,USP)	1 EA	BO	NA	GM		1 EA		1	09/01/2003	01/01/2015						
51552-0894-02		J0945		09/01/2003	01/01/2015	INJECTION, BROMPHENIRAMINE MALEATE, PER 10 MG	BROMPHENIRAMINE MALEATE (1X5GM,USP)	1 EA	BO	NA	GM		10 MG		100	09/01/2003	01/01/2015						
51552-0894-04		J0945		09/01/2003	01/01/2015	INJECTION, BROMPHENIRAMINE MALEATE, PER 10 MG	BROMPHENIRAMINE MALEATE (1X25GM,USP)	1 EA	BO	NA	GM		10 MG		100	09/01/2003	01/01/2015						
51552-0894-05		J0945		09/01/2003	01/01/2015	INJECTION, BROMPHENIRAMINE MALEATE, PER 10 MG	BROMPHENIRAMINE MALEATE (1X100GM,USP)	1 EA	BO	NA	GM		10 MG		100	09/01/2003	01/01/2015						
51552-0910-04		J1800		09/01/2003	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HYDROCHLORIDE (USP, 1X25GM)	1 EA	JR	NA	GM		1 MG		1000	09/01/2003	99/99/9999						
51552-0910-05		J1800		09/01/2003	01/01/2015	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HYDROCHLORIDE (USP, 1X100GM)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	01/01/2015						
51552-0913-01		J1840		09/01/2003	01/01/2015	INJECTION, KANAMYCIN SULFATE, UP TO 500 MG	KANAMYCIN SULFATE (1X1GM,USP)	1 EA	BO	NA	GM		500 MG		2	09/01/2003	01/01/2015						
51552-0913-02		J1840		09/01/2003	01/01/2015	INJECTION, KANAMYCIN SULFATE, UP TO 500 MG	KANAMYCIN SULFATE (1X5GM,USP)	1 EA	BO	NA	GM		500 MG		2	09/01/2003	01/01/2015						
51552-0920-02		J1835		09/01/2003	99/99/9999	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE (1X5GM)	1 EA	JR	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0920-04		J1835		09/01/2003	99/99/9999	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE (1X25GM)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0920-05		J1835		09/01/2003	99/99/9999	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE (1X100GM)	1 EA	BO	NA	GM		50 MG		20	09/01/2003	99/99/9999						
51552-0920-06		J1835		09/01/2003	01/01/2015	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE (1X500GM)	1 EA	NA	NA	GM		50 MG		20	09/01/2003	01/01/2015						
51552-0940-02		J1940		09/01/2003	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (U.S.P.)	1 EA	BO	NA	GM		20 MG		50	09/01/2003	99/99/9999						
51552-0952-01		J0515		09/01/2003	01/01/2015	INJECTION, BENZTROPINE MESYLATE, PER 1 MG	BENZTROPINE MESYLATE (1X1GM,USP)	1 EA	BO	NA	GM		1 MG		1000	09/01/2003	01/01/2015						
51552-0958-02		J1030		09/01/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (USP, 1X5GM, MICRONIZED)	1 EA	BO	NA	GM		40 MG		25	09/01/2003	99/99/9999						
51552-0958-04		J1030		09/01/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (USP, 1X25GM, MICRONIZED)	1 EA	BO	NA	GM		40 MG		25	09/01/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51552-0958-05		J1030		09/01/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (USP, 1X100GM, MICRONIZED)	1	EA	BO	NA	GM	40	MG	25	09/01/2003	99/99/9999						
51552-0958-06		J1030		09/01/2003	01/01/2015	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (USP, 1X500GM, MICRONIZED)	1	EA	BO	NA	GM	40	MG	25	09/01/2003	01/01/2015						
51552-0978-05		J3000		09/01/2003	01/01/2015	INJECTION, STREPTOMYCIN, UP TO 1 GM	STREPTOMYCIN SULFATE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	09/01/2003	01/01/2015						
51552-0979-04		Q0178		09/01/2003	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	12/31/2013						
51552-0991-01		J0760		09/01/2003	99/99/9999	INJECTION, COLCHICINE, PER 1 MG	COLCHICINE (1X1GM, USP)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	99/99/9999						
51552-0999-02		J7636		09/01/2003	01/01/2015	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE (1X5GM)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	01/01/2015						
51552-0999-04		J7636		09/01/2003	01/01/2015	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE (1X25GM)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	01/01/2015						
51552-1018-05		J2800		09/01/2003	01/01/2015	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (USP, 1X100GM)	1	EA	BO	NA	GM	10	ML	1	09/01/2003	01/01/2015						
51552-1025-04		J3360		09/01/2003	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (1X5GM, USP)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	99/99/9999						
51552-1025-04		J3360		09/01/2003	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (1X25GM, USP)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	99/99/9999						
51552-1025-05		J3360		09/01/2003	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (1X100GM, USP)	1	EA	BO	NA	GM	5	MG	200	09/01/2003	99/99/9999						
51552-1031-01		J1450		09/01/2003	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (1X1GM)	1	EA	JR	NA	GM	200	MG	5	09/01/2003	99/99/9999						
51552-1031-02		J1450		09/01/2003	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (1X5GM)	1	EA	JR	NA	GM	200	MG	5	09/01/2003	99/99/9999						
51552-1031-04		J1450		09/01/2003	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (1X25GM)	1	EA	JR	NA	GM	200	MG	5	09/01/2003	99/99/9999						
51552-1036-01		J3370		09/01/2003	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HYDROCHLORIDE (1X1GM, USP)	1	EA	JR	NA	GM	500	MG	2	09/01/2003	99/99/9999						
51552-1036-09		J3370		09/01/2003	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HYDROCHLORIDE (1X250MG, USP)	1	EA	JR	NA	GM	500	MG	2	09/01/2003	99/99/9999						
51552-1045-01		J3420		09/01/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (1X1GM, USP)	1	EA	BO	NA	GM	1000	MCG	1000	09/01/2003	99/99/9999						
51552-1045-09		J3420		09/01/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (1X500MG, USP)	1	EA	BO	NA	GM	1000	MCG	1000	09/01/2003	99/99/9999						
51552-1053-06		J1212		09/01/2003	99/99/9999	INJECTION, DMSO, DIMETHYL SULFOXIDE, 50%, 50 ML	DIMETHYLSULFOXIDE	473	ML	BO	NA	ML	50	%	0.02	09/01/2003	99/99/9999						
51552-1054-01		J8610		09/01/2003	01/01/2015	METHOTREXATE; ORAL, 2.5 MG	METHOTREXATE (USP, 1X1GM)	1	EA	BO	NA	GM	2.5	MG	400	09/01/2003	01/01/2015						
51552-1054-09		J8610		09/01/2003	01/01/2015	METHOTREXATE; ORAL, 2.5 MG	METHOTREXATE (USP, 1X100MG)	1	EA	BO	NA	GM	2.5	MG	400	09/01/2003	01/01/2015						
51552-1063-02		J3430		09/01/2003	01/01/2015	INJECTION, PHYTONADIONE (VITAMIN K), PER 1 MG	PHYTONADIONE (USP, 1X5GM)	1	EA	BO	NA	GM	1	MG	1000	09/01/2003	01/01/2015						
51552-1069-02		J2460		09/01/2003	99/99/9999	INJECTION, OXYTETRACYCLINE HCL, UP TO 50 MG	OXYTETRACYCLINE HCL (U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/01/2003	99/99/9999						
51655-0020-24		J7506		01/01/2002	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	01/01/2002	11/16/2012						
51655-0020-52		J7506		01/01/2002	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	20	EA	BO	PO	EA	5	MG	4	01/01/2002	11/16/2012						
51655-0020-53		J7506		01/01/2002	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	10	EA	BO	PO	EA	5	MG	4	01/01/2002	11/16/2012						
51655-0020-80		J7506		06/22/2005	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	8	EA	NA	PO	EA	5	MG	4	06/22/2005	11/16/2012						
51655-0084-27		Q0170		01/01/2002	11/16/2012	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	12	EA	BO	PO	EA	25	MG	1	01/01/2002	11/16/2012						
51655-0084-53		Q0170		06/22/2005	11/16/2012	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10	EA	NA	PO	EA	25	MG	1	06/22/2005	11/16/2012						
51655-0086-24		J7506		01/01/2002	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	30	EA	BO	PO	EA	5	MG	1	01/01/2002	11/16/2012						
51655-0086-27		J7506		01/01/2002	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	12	EA	BO	PO	EA	5	MG	1	01/01/2002	11/16/2012						
51655-0086-51		J7506		06/22/2005	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	40	EA	NA	PO	EA	5	MG	1	06/22/2005	11/16/2012						
51655-0087-24		J7506		01/01/2005	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	30	EA	NA	PO	EA	5	MG	2	01/01/2005	11/16/2012						
51655-0087-28		J7506		06/22/2005	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21	EA	NA	PO	EA	5	MG	2	06/22/2005	11/16/2012						
51655-0087-49		J7506		06/22/2005	11/16/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	42	EA	NA	PO	EA	5	MG	2	06/22/2005	11/16/2012						
51655-0088-24		Q0163		01/01/2002	11/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30	EA	BX	PO	EA	50	MG	1	01/01/2002	11/16/2012						
51655-0088-52		Q0163		01/01/2002	11/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20	EA	BO	PO	EA	50	MG	1	01/01/2002	11/16/2012						
51655-0093-87		Q0164		06/22/2005	11/16/2012	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 5 MG	6	EA	NA	PO	EA	5	MG	1	06/22/2005	11/16/2012						
51655-0113-24		Q0163		01/01/2002	11/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30	EA	BX	PO	EA	50	MG	0.5	01/01/2002	11/16/2012						
51655-0113-25		Q0163		01/01/2002	11/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60	EA	BO	PO	EA	50	MG	0.5	01/01/2002	11/16/2012						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51655-0113-27		Q0163		01/01/2002	11/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	12	EA	BX	PO	EA	50	MG	0.5	01/01/2002	11/16/2012						
51655-0113-80		Q0163		06/22/2005	11/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	8	EA	NA	PO	EA	50	MG	0.5	06/22/2005	11/16/2012						
51655-0133-54		Q0163		06/22/2005	11/16/2012	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15	EA	NA	PO	EA	50	MG	0.5	06/22/2005	11/16/2012						
51655-0294-89		Q0165		06/22/2005	11/16/2012	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	4	EA	NA	PO	EA	10	MG	1	06/22/2005	11/16/2012						
51655-0296-51		J8499		06/22/2005	11/16/2012	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40	EA	NA	PO	EA	1	EA	1	06/22/2005	11/16/2012						
51655-0296-54		J8499		06/22/2005	11/16/2012	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	15	EA	NA	PO	EA	1	EA	1	06/22/2005	11/16/2012						
51655-0296-76		J8499		06/22/2005	11/16/2012	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25	EA	NA	PO	EA	1	EA	1	06/22/2005	11/16/2012						
51655-0300-51		J8499		06/22/2005	11/16/2012	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	40	EA	NA	PO	EA	1	EA	1	06/22/2005	11/16/2012						
51655-0300-54		J8499		06/22/2005	11/16/2012	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15	EA	NA	PO	EA	1	EA	1	06/22/2005	11/16/2012						
51655-0300-76		J8499		06/22/2005	11/16/2012	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25	EA	NA	PO	EA	1	EA	1	06/22/2005	11/16/2012						
51655-0523-53		Q0173		01/01/2002	11/16/2012	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 250 MG	10	EA	BO	PO	EA	250	MG	1	01/01/2002	11/16/2012						
51655-0533-52		Q0177		06/22/2005	11/16/2012	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20	EA	NA	PO	EA	25	MG	1	06/22/2005	11/16/2012						
51927-1000-00		J2271		09/08/2003	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.; CII)	1	EA	JR	NA	GM	100	MG	10	09/08/2003	12/31/2014						
51927-1001-00		J7636		09/08/2003	99/99/9999	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1001-00	KO	J7636	KO	09/08/2003	99/99/9999	ATROPINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ATROPINE SULFATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1003-00		J1170		09/08/2003	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.; CII)	1	EA	JR	NA	GM	4	MG	250	09/08/2003	99/99/9999						
51927-1005-00		J2060		09/08/2003	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.; CIV)	1	EA	JR	NA	GM	2	MG	500	09/08/2003	99/99/9999						
51927-1007-00		J1960		09/08/2003	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.; CII)	1	EA	BO	NA	GM	2	MG	500	09/08/2003	99/99/9999						
51927-1012-00		J0592		09/08/2003	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE (U.S.P.; CII)	1	EA	JR	NA	GM	0.1	MG	10000	09/08/2003	99/99/9999						
51927-1013-00		J0745		09/08/2003	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.; CII)	1	EA	BO	NA	GM	30	MG	33.33333	09/08/2003	99/99/9999						
51927-1014-00		J3360		09/08/2003	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (U.S.P.; CIV)	1	EA	JR	NA	GM	5	MG	200	09/08/2003	99/99/9999						
51927-1017-00		J1230		09/08/2003	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.; CII)	1	EA	BO	NA	GM	10	MG	100	09/08/2003	99/99/9999						
51927-1018-00		J2175		09/08/2003	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.; CII)	1	EA	BO	NA	GM	100	MG	10	09/08/2003	99/99/9999						
51927-1019-00		J3010		09/08/2003	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1	EA	JR	NA	GM	0.1	MG	10000	09/08/2003	99/99/9999						
51927-1026-00		J3140		09/08/2003	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE	1	EA	JR	NA	GM	50	MG	20	09/08/2003	12/31/2014						
51927-1027-00		J3140		09/08/2003	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.; SOY; CII)	1	EA	JR	NA	GM	50	MG	20	09/08/2003	12/31/2014						
51927-1029-00		J3150		09/08/2003	12/31/2014	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE MICRONIZED (U.S.P., MICRONIZED)	1	EA	JR	NA	GM	100	MG	10	09/08/2003	12/31/2014						
51927-1046-00		J2675		09/08/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (U.S.P.)	1	EA	JR	NA	GM	50	MG	20	09/08/2003	99/99/9999						
51927-1079-00		J1200		09/08/2003	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (U.S.P.)	1	EA	JR	NA	GM	50	MG	20	09/08/2003	99/99/9999						
51927-1080-00		J1240		09/08/2003	99/99/9999	INJECTION, DIMENHYDRINATE, UP TO 50 MG	DIMENHYDRINATE (U.S.P.)	1	EA	BO	NA	GM	50	MG	20	09/08/2003	99/99/9999						
51927-1082-00		J2765		09/08/2003	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	09/08/2003	99/99/9999						
51927-1085-00		J9190		09/08/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (U.S.P. -5 FU)	1	EA	JR	NA	GM	500	MG	2	09/08/2003	99/99/9999						
51927-1090-00		J3480		12/04/2003	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (USP; GRANULAR)	1	EA	BO	NA	GM	2	MEQ	6.71141	12/04/2003	99/99/9999						
51927-1093-00		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (USP)	1	EA	BO	NA	GM	100	MG	10	01/01/2004	99/99/9999						
51927-1110-00		J1700		09/08/2003	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE MICRONIZED (U.S.P.)	1	EA	JR	NA	GM	25	MG	40	09/08/2003	99/99/9999						
51927-1148-00		J7510		09/08/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE MICRONIZED (ANHYDROUS)	1	EA	JR	NA	GM	5	MG	200	09/08/2003	99/99/9999						
51927-1194-00		J3490		09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	BENZOCANE	1	EA	JR	NA	GM	1	EA	1	09/08/2003	99/99/9999						
51927-1202-00		J0706		12/04/2003	99/99/9999	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATE (PURIFIED)	1	EA	BO	NA	GM	5	MG	200	12/04/2003	99/99/9999						
51927-1213-00		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	1	EA	JR	NA	GM	10	MG	100	01/01/2004	99/99/9999						
51927-1225-00		J7799		09/08/2003	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	PHENYLEPHRINE HCL (U.S.P.)	1	EA	JR	NA	GM	1	EA	1	09/08/2003	99/99/9999						
51927-1242-00		J3411		01/01/2004	99/99/9999	INJECTION, THIAMINE HCL, 100 MG	THIAMINE HYDROCHLORIDE (USP)	1	EA	BO	NA	GM	100	MG	10	01/01/2004	99/99/9999						
51927-1269-00		J3350		12/04/2003	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (USP)	1	EA	BO	NA	GM	40	GM	0.025	12/04/2003	99/99/9999						
51927-1317-00		J3520		12/04/2003	99/99/9999	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM (USP; DIHYDRATE)	1	EA	BO	NA	GM	150	MG	6.66666	12/04/2003	99/99/9999						
51927-1325-00		J2650		09/08/2003	99/99/9999	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1	EA	JR	NA	GM	1	ML	20	09/08/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51927-1326-00		J7684		09/08/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., MICRONIZED)	1	EA	JR	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1326-00	KO	J7684	KO	09/08/2003	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., MICRONIZED)	1	EA	JR	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1332-00		J1030		09/08/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1	EA	BO	NA	GM	40	MG	25	09/08/2003	99/99/9999						
51927-1347-00		J0500		09/08/2003	99/99/9999	INJECTION, DICYCLIMINE HCL, UP TO 20 MG	DICYCLIMINE HYDROCHLORIDE (U.S.P.)	1	EA	BO	NA	GM	20	MG	50	09/08/2003	99/99/9999						
51927-1400-00		J3410		09/08/2003	99/99/9999	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (U.S.P.)	1	EA	JR	NA	GM	25	MG	40	09/08/2003	99/99/9999						
51927-1430-00		J7638		09/08/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	JR	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1430-00	KO	J7638	KO	09/08/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1	EA	JR	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1433-00		J1630		09/08/2003	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1	EA	JR	NA	GM	5	MG	200	09/08/2003	99/99/9999						
51927-1435-00		J7506		09/08/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE MICRONIZED (USP)	1	EA	BO	NA	GM	5	MG	200	09/08/2003	12/31/2015						
51927-1441-00		J9017		12/04/2003	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (TECHNICAL)	1	EA	BO	NA	GM	1	MG	1000	12/04/2003	99/99/9999						
51927-1444-00		J0280		09/08/2003	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE (U.S.P.; ANHYDROUS)	1	EA	JR	NA	GM	250	MG	4	09/08/2003	99/99/9999						
51927-1449-00		J3490		09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1	EA	JR	NA	GM	1	EA	1	09/08/2003	99/99/9999						
51927-1454-00		J7624		09/08/2003	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1	EA	JR	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1454-00	KO	J7624	KO	09/08/2003	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE DIPROPIONATE (U.S.P., MICRONIZED)	1	EA	JR	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1510-00		J2810		09/08/2003	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE (USP, ANHYDROUS)	1	EA	BO	NA	GM	40	MG	25	09/08/2003	99/99/9999						
51927-1565-00		J8610		09/08/2003	99/99/9999	METHOTREXATE, ORAL, 2.5 MG	METHOTREXATE (U.S.P.)	1	EA	BO	NA	GM	2.5	MG	400	09/08/2003	99/99/9999						
51927-1571-00		J1245		09/08/2003	99/99/9999	INJECTION, DIPYRIDAMOLE, PER 10 MG	DIPYRIDAMOLE (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	09/08/2003	99/99/9999						
51927-1573-00		J7609		01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1	EA	JR	NA	GM	1	MG	1000	01/01/2007	99/99/9999						
51927-1573-00	KO	J7609	KO	01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1	EA	JR	NA	GM	1	MG	1000	01/01/2007	99/99/9999						
51927-1575-00		J7643		09/08/2003	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1575-00	KO	J7643	KO	09/08/2003	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1597-00		J3490		12/04/2003	99/99/9999	UNCLASSIFIED DRUGS	ETHANOLAMINE (MONOETHANOLAMINE)	1	EA	BO	NA	GM	1	EA	1	12/04/2003	99/99/9999						
51927-1601-00		J7604		01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51927-1601-00	KO	J7604	KO	01/01/2008	99/99/9999	ACETYL CYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
51927-1603-00		J1320		09/08/2003	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	1	EA	JR	NA	GM	20	MG	50	09/08/2003	99/99/9999						
51927-1606-00		J1800		09/08/2003	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1610-00		J7699		09/08/2003	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1	EA	JR	NA	GM	1	EA	1	09/08/2003	99/99/9999						
51927-1612-00		J1212		12/04/2003	99/99/9999	INJECTION, DIMETHYL SULFOXIDE, 50%, 50 ML	DIMETHYL SULFOXIDE (USP)	1	ML	BO	NA	ML	50	%	0.02	12/04/2003	99/99/9999						
51927-1641-00		J7622		09/08/2003	99/99/9999	BECLMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLMETHASONE DIPROPIONATE (U.S.P. (ANHYDROUS))	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1641-00	KO	J7622	KO	09/08/2003	99/99/9999	BECLMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BECLMETHASONE DIPROPIONATE (U.S.P. (ANHYDROUS))	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1648-00		J7645		01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE	1	EA	JR	NA	GM	1	MG	1000	01/01/2007	99/99/9999						
51927-1648-00	KO	J7645	KO	01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE	1	EA	JR	NA	GM	1	MG	1000	01/01/2007	99/99/9999						
51927-1659-00		J1180		09/08/2003	99/99/9999	INJECTION, DYPHYLLINE, UP TO 500 MG	DYPHYLLINE	1	EA	BO	NA	GM	500	MG	2	09/08/2003	99/99/9999						
51927-1662-00		J3420		12/04/2003	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (USP)	1	EA	BO	NA	GM	1000	MCG	1000	12/04/2003	99/99/9999						
51927-1683-00		J3490		09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN PHOSPHATE (U.S.P.)	1	EA	JR	NA	GM	1	EA	1	09/08/2003	99/99/9999						
51927-1706-00		J1110		09/08/2003	99/99/9999	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG	DIHYDROERGOTAMINE MESYLATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1709-00		J1435		09/08/2003	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P. E-1)	1	EA	JR	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1715-00		J7799		09/08/2003	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	EPINEPHRINE HCL (USP)	1	EA	BO	NA	GM	1	EA	1	09/08/2003	99/99/9999						
51927-1722-00		J3430		12/04/2003	99/99/9999	INJECTION, PHYTONADIONE (VITAMIN K), PER 1 MG	MENADIOLONE (USP)	1	EA	BO	NA	GM	1	MG	1000	12/04/2003	99/99/9999						
51927-1726-00		J0285		09/08/2003	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.; ORAL GRADE)	1	EA	JR	NA	GM	50	MG	20	09/08/2003	99/99/9999						
51927-1742-00		J3370		09/08/2003	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (U.S.P.)	1	EA	JR	NA	GM	500	MG	2	09/08/2003	99/99/9999						
51927-1775-00		J2440		09/08/2003	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1	EA	JR	NA	GM	60	MG	16.66666	09/08/2003	99/99/9999						
51927-1776-00		J3490		09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (USP (6))	1	EA	BO	NA	GM	1	EA	1	09/08/2003	99/99/9999						
51927-1781-00		J2150		12/04/2003	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (USP)	1	EA	BO	NA	GM	50	ML	0.08	12/04/2003	99/99/9999						
51927-1784-00		J1940		09/08/2003	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (U.S.P.)	1	EA	BO	NA	GM	20	MG	50	09/08/2003	99/99/9999						
51927-1788-00		J3000		09/08/2003	99/99/9999	INJECTION, STREPTOMYCIN, UP TO 1 GM	STREPTOMYCIN SULFATE	1	EA	BO	NA	GM	1	GM	1	09/08/2003	99/99/9999						
51927-1794-00		J7641		09/08/2003	99/99/9999	FLUNISOLIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, PER MILLIGRAM	FLUNISOLIDE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						
51927-1794-00	KO	J7641	KO	09/08/2003	99/99/9999	FLUNISOLIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, PER MILLIGRAM	FLUNISOLIDE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/08/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51927-1829-00		J3490		09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	CORTISONE ACETATE MICRONIZED (USP)	1 EA	JR	NA	GM		1 EA		1	09/08/2003	99/99/9999						
51927-1831-00		J1980		09/08/2003	99/99/9999	INJECTION, HYOSCYAMINE SULFATE, UP TO 0.25 MG	HYOSCYAMINE SULFATE (U.S.P.)	1 EA	BO	NA	GM	0.25 MG	4000		4000	09/08/2003	99/99/9999						
51927-1838-00		J1165		09/08/2003	99/99/9999	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	JR	NA	GM	50 MG	20		20	09/08/2003	99/99/9999						
51927-1865-00		J1955		12/04/2003	99/99/9999	INJECTION, LEVOCARNITINE, PER 1 GM	LEVOCARNITINE (USP)	1 EA	BO	NA	GM	1 GM	1		1	12/04/2003	99/99/9999						
51927-1895-00		J0760		09/08/2003	99/99/9999	INJECTION, COLCHICINE, PER 1MG	COLCHICINE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-1925-00		J3430		09/08/2003	99/99/9999	INJECTION, PHYTONADIONE (VITAMIN K), PER 1 MG	PHYTONADIONE (USP; VITAMIN K1)	1 EA	BO	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-1950-00		J0945		09/08/2003	99/99/9999	INJECTION, BROMPHENIRAMINE MALEATE, PER 10 MG	BROMPHENIRAMINE MALEATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		100	09/08/2003	99/99/9999						
51927-1951-00		J7624		09/08/2003	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	JR	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-1951-00	KO	J7624	KO	09/08/2003	99/99/9999	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 EA	JR	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-1954-00		J3490		09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 EA	JR	NA	GM	1 EA	1		1	09/08/2003	99/99/9999						
51927-1956-00		J3490		09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	RIFAMPIN (U.S.P.)	1 EA	JR	NA	GM	1 EA	1		1	09/08/2003	99/99/9999						
51927-1981-00		J3250		09/12/2003	99/99/9999	INJECTION, TRIMETHOBENZAMIDE HCL, UP TO 200 MG	TRIMETHOBENZAMIDE HCL	1 EA	BO	NA	GM	200 MG	5		5	09/12/2003	99/99/9999						
51927-2007-00		J0475		09/08/2003	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	JR	NA	GM	10 MG	100		100	09/08/2003	99/99/9999						
51927-2097-00		J0520		09/08/2003	99/99/9999	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (U.S.P.)	1 EA	JR	NA	GM	5 MG	200		200	09/08/2003	99/99/9999						
51927-2101-00		J0770		09/08/2003	99/99/9999	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE SODIUM (USP)	1 EA	BO	NA	GM	150 MG	6.66666		6.66666	09/08/2003	99/99/9999						
51927-2116-00		J0152		01/01/2004	12/31/2013	INJECTION, ADENOSINE FOR DIAGNOSTIC USE, 30 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOSINE	1 EA	BO	NA	GM	30 MG	33.33333		33.33333	01/01/2004	12/31/2013						
51927-2118-00		J2360		09/08/2003	99/99/9999	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORPHENADRINE CITRATE (USP)	1 EA	BO	NA	GM	60 MG	16.66666		16.66666	09/08/2003	99/99/9999						
51927-2132-00		J0152		01/01/2004	12/31/2013	INJECTION, ADENOSINE FOR DIAGNOSTIC USE, 30 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOSINE (TRIHYDRATE)	1 EA	BO	NA	GM	30 MG	33.33333		33.33333	01/01/2004	12/31/2013						
51927-2132-00		J0152		01/01/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		100	09/08/2003	12/31/2013						
51927-2140-00		J2300		09/08/2003	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL	1 EA	BO	NA	GM	10 MG	100		100	09/08/2003	99/99/9999						
51927-2182-00		J1790		09/08/2003	99/99/9999	INJECTION, DROPERIDOL, UP TO 5 MG	DROPERIDOL (USP)	1 EA	BO	NA	GM	5 MG	200		200	09/08/2003	99/99/9999						
51927-2196-00		J0270		09/08/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	JR	NA	GM	1.25 MCG	800000		800000	09/08/2003	99/99/9999						
51927-2206-00		J0780		09/08/2003	99/99/9999	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE (USP)	1 EA	BO	NA	GM	10 MG	100		100	09/08/2003	99/99/9999						
51927-2231-00		J1094		09/08/2003	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 EA	JR	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-2234-00		J2680		09/08/2003	99/99/9999	INJECTION, FLUPHENAZINE DECANOATE, UP TO 25 MG	FLUPHENAZINE DECANOATE (U.S.P.)	1 EA	BO	NA	GM	25 MG	40		40	09/08/2003	99/99/9999						
51927-2258-00		J7501		09/08/2003	99/99/9999	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (USP)	1 EA	BO	NA	GM	100 MG	10		10	09/08/2003	99/99/9999						
51927-2303-00		J0364		01/01/2007	99/99/9999	INJECTION, APMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P., HEMIHYDRATE)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
51927-2316-00		J0178		09/08/2003	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (U.S.P.)	1 EA	JR	NA	GM	50 MG	20		20	09/08/2003	12/31/2013						
51927-2319-00		J1265		01/01/2006	99/99/9999	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HCL	1 EA	BO	NA	GM	40 MG	25		25	01/01/2006	99/99/9999						
51927-2375-00		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN (USP)	1 EA	BO	NA	GM	300 MG	3.33333		3.33333	01/01/2007	99/99/9999						
51927-2375-00	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN (USP)	1 EA	BO	NA	GM	300 MG	3.33333		3.33333	01/01/2007	99/99/9999						
51927-2379-00		J0735		09/08/2003	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 EA	JR	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-2319-00		J2800		09/08/2003	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (U.S.P.)	1 EA	BO	NA	GM	10 ML	1		1	09/08/2003	99/99/9999						
51927-2669-00		J2760		09/08/2003	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		200	09/08/2003	99/99/9999						
51927-2692-00		J0640		09/08/2003	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (USP; ANHYDROUS)	1 EA	JR	NA	GM	50 MG	20		20	09/08/2003	99/99/9999						
51927-2704-00		J0278		01/01/2006	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (U.S.P.)	1 EA	BO	NA	GM	100 MG	10		10	01/01/2006	99/99/9999						
51927-2706-00		J1070		09/08/2003	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.; CIII)	1 EA	JR	NA	GM	100 MG	10		10	09/08/2003	12/31/2014						
51927-2732-00		J3475		12/04/2003	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (USP; HEPTAHYDRATE)	1 EA	BO	NA	GM	500 MG	2		2	12/04/2003	99/99/9999						
51927-2742-00		J1730		09/08/2003	99/99/9999	INJECTION, DIAZOXIDE, UP TO 300 MG	DIAZOXIDE (U.S.P.)	1 EA	BO	NA	GM	300 MG	3.33333		3.33333	09/08/2003	99/99/9999						
51927-2762-00		J9340		09/08/2003	99/99/9999	INJECTION, THIOTEPA, 15 MG	TRIETHYLENETHIOPHOSPHORAMIDE/T	1 EA	BO	NA	GM	15 MG	66.66666		66.66666	09/08/2003	99/99/9999						
51927-2765-00		J7681		09/08/2003	99/99/9999	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-2765-00	KO	J7681	KO	09/08/2003	99/99/9999	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	09/08/2003	99/99/9999						
51927-2895-00		J1600		09/08/2003	99/99/9999	INJECTION, GOLD SODIUM THIOALATE, UP TO 50 MG	GOLD SODIUM THIOALATE	1 EA	BO	NA	GM	50 MG	20		20	09/08/2003	99/99/9999						
51927-2986-00		J0595		01/01/2004	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (U.S.P.; CIV)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2004	99/99/9999						
51927-2994-00		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		200	01/01/2006	99/99/9999						
51927-3023-00		J2780		09/08/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	JR	NA	GM	25 MG	40		40	09/08/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3		
51927-3115-00	J2690			09/08/2003	99/99/9999	INJECTION, PROCAINAMIDE HCL, UP TO 1 GM	PROCAINAMIDE HCL (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	09/08/2003	99/99/9999								
51927-3163-00	J1000			09/08/2003	99/99/9999	INJECTION, DEPO-ESTRADIOL CYPIONATE, UP TO 5 MG	ESTRADIOL CYPIONATE (U.S.P.)	1 EA	JR	NA	GM		5 MG		200	09/08/2003	99/99/9999								
51927-3177-00	J2010			09/08/2003	99/99/9999	INJECTION, LINCOMYCIN HCL, UP TO 300 MG	LINCOMYCIN HCL (U.S.P.)	1 EA	BO	NA	GM		300 MG		3.33333	09/08/2003	99/99/9999								
51927-3196-00	J7516			09/08/2003	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORIN A (USP)	1 EA	JR	NA	GM		250 MG		4	09/08/2003	99/99/9999								
51927-3258-00	J2460			09/08/2003	99/99/9999	INJECTION, OXYTETRACYCLINE HCL, UP TO 50 MG	OXYTETRACYCLINE HCL (U.S.P.)	1 EA	BO	NA	GM		50 MG		20	09/08/2003	99/99/9999								
51927-3286-00	J1644			09/08/2003	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (USP)	1 EA	BO	NA	GM		1000 U		160	09/08/2003	99/99/9999								
51927-3335-00	J2310			09/08/2003	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL DIHYDRATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	09/08/2003	99/99/9999								
51927-3370-00	J3302			09/08/2003	99/99/9999	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (USP)	1 EA	JR	NA	GM		5 MG		200	09/08/2003	99/99/9999								
51927-3408-00	J3490			09/08/2003	99/99/9999	UNCLASSIFIED DRUGS	FAMOTIDINE (U.S.P.)	1 EA	JR	NA	GM		1 EA		1	09/08/2003	99/99/9999								
51927-3422-00	J0636			09/08/2003	99/99/9999	INJECTION, CALCITRIOL, 0.1 MCG	CALCITRIOL IN ALMOND OIL (NF) 1 MCG/ML	1 ML	BO	NA	ML		0.1 MCG		10	09/08/2003	99/99/9999								
51927-3484-00	J2725			09/08/2003	99/99/9999	INJECTION, PROTIRELIN, PER 250 MCG	PROTIRELIN	1 EA	BO	NA	GM		250 MCG		4000	09/08/2003	99/99/9999								
51927-3530-00	J2675			09/08/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (U.S.P.)	1 EA	JR	NA	GM		50 MG		20	09/08/2003	99/99/9999								
51927-3557-00	J7507			01/01/2004	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS	0.001 GM	JR	NA	GM		1 MG		1000	01/01/2004	99/99/9999								
51927-3613-00	J2515			03/26/2004	99/99/9999	INJECTION, PENTOBARBITAL SODIUM, PER 50 MG	PENTOBARBITAL SODIUM (U.S.P.)	1 EA	BO	NA	GM		50 MG		20	03/26/2004	99/99/9999								
51927-3634-00	J3490			01/04/2008	99/99/9999	UNCLASSIFIED DRUGS	CIPROFLOXACIN HYDROCHLORIDE (USP)	1 EA	BO	NA	GM		1 EA		1	01/04/2008	99/99/9999								
51927-3643-00	J7640			01/01/2006	99/99/9999	FORMOTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 12 MICROGRAMS	FORMOTEROL FUMARATE (DIHYDRATE)	1 EA	BO	NA	GM		12 MCG		83333.33	01/01/2006	99/99/9999								
51927-3643-00	KO	J7640	KO	01/01/2006	99/99/9999	FORMOTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 12 MICROGRAMS	FORMOTEROL FUMARATE (DIHYDRATE)	1 EA	BO	NA	GM		12 MCG		83333.33	01/01/2006	99/99/9999								
51927-9017-00	J2675			09/08/2003	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.; WETTABLE POWDER)	1 EA	JR	NA	GM		50 MG		20	09/08/2003	99/99/9999								
51927-9018-00	J2550			09/08/2003	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	JR	NA	GM		50 MG		20	09/08/2003	99/99/9999								
51991-0188-31	J7509			11/05/2003	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (UNIT OF USE) 4 MG	21 EA	DP	PO	EA		4 MG		1	11/05/2003	99/99/9999								
51991-0458-01	J7506			01/16/2006	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.) 1 MG	100 EA	BO	PO	EA		5 MG		0.2	01/16/2006	12/31/2015								
52544-0153-02	J3315			12/30/2004	03/12/2017	INJECTION, TRIPTORELIN PAMOATE, 3.75 MG	TRELSTAR DEPOT (SDV) 3.75 MG	1 EA	VL	IM	EA		3.75 MG		1	12/30/2004	03/12/2017								
52544-0154-02	J3315			12/30/2004	03/12/2017	INJECTION, TRIPTORELIN PAMOATE, 3.75 MG	TRELSTAR LA (SDV) 11.25 MG	1 EA	VL	IM	EA		3.75 MG		3	12/30/2004	03/12/2017								
00517-0710-01	J1451			07/16/2018	99/99/9999	INJECTION, FOMEPIZOLE, 15 MG	FOMEPIZOLE (1X1.5ML,PF) 1 GM/1 ML	1.5 ML	VL	IV	ML		15 MG		66.66666	07/16/2018	99/99/9999								
13533-0335-04	J1460			08/24/2018	99/99/9999	INJECTION, GAMMA GLOBULIN, INTRAMUSCULAR, 1 CC	GAMASTAN (SDV,PF,LATEX-FREE) 15%-18%	2 ML	VL	IM	ML		1 CC		1	08/24/2018	99/99/9999								
13533-0335-12	J1460			08/24/2018	99/99/9999	INJECTION, GAMMA GLOBULIN, INTRAMUSCULAR, 1 CC	GAMASTAN (SDV,PF,LATEX-FREE) 15%-18%	10 ML	VL	IM	ML		1 CC		1	08/24/2018	99/99/9999								
52769-0470-72	J1566			01/01/2006	99/99/9999	OTHERWISE SPECIFIED, 500 MG	POLYGM (W/50 ML DILUENT) 2.5 MG	1 EA	NA	IV	EA		500 MG		0.005	01/01/2006	99/99/9999								
52959-0043-00	Q0163			06/17/2003	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100 EA	BO	PO	EA		50 MG		0.5	06/17/2003	99/99/9999								
42023-0221-10	J1335			07/26/2018	99/99/9999	INJECTION, ERTAPENEM SODIUM, 500 MG	ERTAPENEM 1 GM	10 EA	VL	IJ	EA		500 MG		2	07/26/2018	99/99/9999								
44567-0420-24	J3475			07/23/2018	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (NEXCEL BAG,LATEX-FREE) 40 MG/1 ML	50 ML	FC	IV	ML		500 MG		0.08	07/23/2018	99/99/9999								
44567-0421-24	J3475			07/23/2018	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (NEXCEL BAG,LATEX-FREE) 40 MG/1 ML	100 ML	FC	IV	ML		500 MG		0.08	07/23/2018	99/99/9999								
52959-0043-04	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	4 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999								
52959-0043-10	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	10 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999								
52959-0043-15	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999								
52959-0043-20	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999								
52959-0043-24	Q0163			05/12/2003	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	24 EA	BO	PO	EA		50 MG		0.5	05/12/2003	99/99/9999								
52959-0043-30	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999								
52959-0043-50	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	50 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999								

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
52959-0043-60		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60	EA	BO	PO	EA	50	MG	0.5	01/01/2002	02/03/2016						
52959-0053-06		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	6	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
52959-0053-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	10	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
52959-0053-12		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	12	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
52959-0053-15		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
52959-0053-20		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
52959-0053-30		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
52959-0053-52		Q0163		01/24/2005	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	52	EA	BO	PO	EA	50	MG	1	01/24/2005	99/99/9999						
52959-0079-00		J7500		01/01/2002	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	IMURAN 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
52959-0100-00		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (DOSE PACK) 4 MG	21	EA	DP	PO	EA	4	MG	1	01/01/2002	99/99/9999						
52959-0123-03		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	120	ML	BO	PO	ML	50	MG		0.05	01/01/2002	99/99/9999					
52959-0123-06		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	180	ML	BO	PO	ML	50	MG		0.05	01/01/2002	99/99/9999					
52959-0126-00		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-05		J7506		11/06/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	5	EA	BO	PO	EA	5	MG	2	11/06/2002	12/31/2015						
52959-0126-07		J7506		11/06/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	7	EA	BO	PO	EA	5	MG	2	11/06/2002	12/31/2015						
52959-0126-10		J7506		08/19/2003	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	10	EA	BO	PO	EA	5	MG	2	08/19/2003	12/31/2015						
52959-0126-12		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	12	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-15		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	15	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-18		J7506		01/15/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	18	EA	BO	PO	EA	5	MG	2	01/15/2002	12/31/2015						
52959-0126-20		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	20	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-21		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-25		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	25	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-37		J7506		07/18/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	37	EA	BO	PO	EA	5	MG	2	07/18/2007	12/31/2015						
52959-0126-40		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	40	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-42		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	42	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-44		J7506		03/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	44	EA	BO	PO	EA	5	MG	2	03/01/2004	12/31/2015						
52959-0126-45		J7506		09/19/2006	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	45	EA	NA	PO	EA	5	MG	2	09/19/2006	12/31/2015						
52959-0126-50		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	50	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0126-60		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	60	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015						
52959-0127-00		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	100	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-07		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	7	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-10		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	10	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-12		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	12	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-15		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	15	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-20		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	20	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-21		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	21	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-25		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	25	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-37		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	37	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
52959-0127-42		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	42	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
52959-0158-06		J7669		01/01/2002	02/03/2016	METAPROTERENOL SULFATE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	ALUPENT (VIAL) 0.6%	2.5	ML	AM	IH	ML	10	MG	0.6	01/01/2002	02/03/2016							
52959-0158-06	KO	J7669	KO	01/01/2002	02/03/2016	METAPROTERENOL SULFATE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	ALUPENT (VIAL) 0.6%	2.5	ML	AM	IH	ML	10	MG	0.6	01/01/2002	02/03/2016							
52959-0179-06		J2360		01/01/2002	01/27/2016	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	NORFLEX 30 MG/ML	2	ML	AM	IU	ML	60	MG	0.5	01/01/2002	01/27/2016							
52959-0220-00		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0220-10		J7506		08/19/2003	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	10	EA	BO	PO	EA	5	MG	1	08/19/2003	12/31/2015							
52959-0220-20		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	20	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0220-21		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	21	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0220-30		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	30	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0220-36		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	36	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0220-40		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	40	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0220-60		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	60	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0220-75		J7506		01/01/2002	12/31/2015	PREDNISON, ORAL, PER 5MG	PREDNISON 5 MG	75	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
52959-0237-12		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999							
52959-0244-00		None		10/02/2000	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	100	EA	BO	PO	EA	2.5	MG	1	10/02/2000	99/99/9999							
52959-0291-00		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	COMPazine 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	02/03/2016							
52959-0313-15		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	01/01/2002	99/99/9999							
52959-0330-00		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 200 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
52959-0330-25		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 200 MG	25	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
52959-0330-50		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 200 MG	50	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
52959-0355-06		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	6	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999							
52959-0355-12		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999							
52959-0391-15		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPazine 10 MG	15	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013							
52959-0392-12		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	12	EA	BO	PO	EA	0.25	MG	3	01/01/2006	99/99/9999							
52959-0392-21		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	21	EA	DP	PO	EA	0.25	MG	3	01/01/2006	99/99/9999							
52959-0392-28		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	28	EA	BO	PO	EA	0.25	MG	3	01/01/2006	99/99/9999							
52959-0433-10		Q0177		06/06/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	10	EA	BO	PO	EA	25	MG	1	06/06/2002	99/99/9999							
52959-0433-15		Q0177		02/28/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	15	EA	BO	PO	EA	25	MG	1	02/28/2002	99/99/9999							
52959-0433-20		Q0177		12/27/2004	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20	EA	BO	PO	EA	25	MG	1	12/27/2004	99/99/9999							
52959-0433-30		Q0177		10/17/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	BO	PO	EA	25	MG	1	10/17/2002	99/99/9999							
52959-0433-40		Q0177		01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	40	EA	BO	PO	EA	25	MG	1	01/01/2002	99/99/9999							
52959-0433-60		Q0177		12/27/2004	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	60	EA	BO	PO	EA	25	MG	1	12/27/2004	99/99/9999							
52959-0476-02		Q0165		08/09/2005	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	120	EA	BO	PO	EA	10	MG	1	08/09/2005	12/31/2013							
52959-0476-10		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013							
52959-0476-15		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013							
52959-0476-20		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
52959-0476-24		Q0165		10/27/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	24	EA	BO	PO	EA	10 MG		1	10/27/2004	12/31/2013						
52959-0476-30		Q0165		11/22/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	10 MG		1	11/22/2004	12/31/2013						
52959-0476-60		Q0165		11/22/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	10 MG		1	11/22/2004	12/31/2013						
52959-0479-10		Q0173		01/01/2002	02/03/2016	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 250 MG	10	EA	BO	PO	EA	250 MG		1	01/01/2002	02/03/2016						
52959-0479-12		Q0173		01/01/2002	02/03/2016	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 250 MG	12	EA	BO	PO	EA	250 MG		1	01/01/2002	02/03/2016						
52959-0479-20		Q0173		01/01/2002	02/03/2016	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 250 MG	20	EA	BO	PO	EA	250 MG		1	01/01/2002	02/03/2016						
52959-0479-30		Q0173		01/01/2002	10/17/2016	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 250 MG	30	EA	BO	PO	EA	250 MG		1	01/01/2002	10/17/2016						
52959-0505-06		Q0144		01/01/2002	99/99/9999	AZITHROMYICIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX Z-PAK 250 MG	6	EA	DP	PO	EA	1 GM		0.25	01/01/2002	99/99/9999						
52959-0517-25		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0517-30		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0517-35		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	35	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-01		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-10		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	10	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-12		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	12	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-15		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-21		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	21	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-25		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-30		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0544-40		J8499		08/24/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	40	EA	BO	PO	EA	1 EA		1	08/24/2007	99/99/9999						
52959-0544-50		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	50	EA	BO	PO	EA	1 EA		1	01/01/2002	99/99/9999						
52959-0547-04		J8540		05/16/2007	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	4	EA	BO	PO	EA	0.25 MG		16	05/16/2007	99/99/9999						
52959-0547-10		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	10	EA	BO	PO	EA	0.25 MG		16	01/01/2006	99/99/9999						
52959-0547-11		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	11	EA	BO	PO	EA	0.25 MG		16	01/01/2006	99/99/9999						
52959-0547-12		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	12	EA	BO	PO	EA	0.25 MG		16	01/01/2006	99/99/9999						
52959-0547-16		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	16	EA	BO	PO	EA	0.25 MG		16	01/01/2006	99/99/9999						
52959-0547-20		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	20	EA	BO	PO	EA	0.25 MG		16	01/01/2006	99/99/9999						
52959-0547-30		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	30	EA	BO	PO	EA	0.25 MG		16	01/01/2006	99/99/9999						
52959-0547-50		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	50	EA	BO	PO	EA	0.25 MG		16	01/01/2006	99/99/9999						
52959-0561-01		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 12.5 MG	12	EA	BX	RC	EA	1 EA		1	01/01/2006	02/03/2016						
52959-0561-04		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 12.5 MG	4	EA	BX	RC	EA	1 EA		1	01/01/2006	02/03/2016						
52959-0562-01		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 25 MG	12	EA	NA	RC	EA	1 EA		1	01/01/2006	02/03/2016						
52959-0562-06		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 25 MG	6	EA	NA	RC	EA	1 EA		1	01/01/2006	02/03/2016						
52959-0622-60		J7510		01/01/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (CHERRY) 15 MG/5 ML	480	ML	BO	PO	ML	5 MG		0.6	01/01/2002	99/99/9999						
52959-0657-03		Q0144		01/01/2002	99/99/9999	AZITHROMYICIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	15	ML	BO	PO	ML	1 GM		0.04	01/01/2002	99/99/9999						
52959-0657-06		Q0144		01/01/2006	99/99/9999	AZITHROMYICIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	22.5	ML	BO	PO	ML	1 GM		0.04	01/01/2006	99/99/9999						
52959-0678-30		J8499		10/07/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30	EA	BO	PO	EA	1 EA		1	10/07/2003	99/99/9999						
52959-0741-20		J7611		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DMC, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE 0.5%	20	ML	BO	IH	ML	1 MG		5	04/01/2008	99/99/9999						
52959-0748-01		J8501		08/22/2007	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND 40 MG	1	EA	BO	PO	EA	5 MG		8	08/22/2007	99/99/9999						
52959-0804-04		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	120	ML	BO	PO	ML	25 MG		0.05	01/01/2002	12/31/2013						
52959-0804-08		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	240	ML	BO	PO	ML	25 MG		0.05	01/01/2002	12/31/2013						
52959-0817-10		Q0173		10/04/2005	99/99/9999	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 300 MG	10	EA	BO	PO	EA	250 MG		1.2	10/04/2005	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
52959-0833-06		Q0178		10/14/2005	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	6 EA	BO	PO	EA		50 MG		1	10/14/2005	12/31/2013						
52959-0833-20		Q0178		10/14/2005	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20 EA	BO	PO	EA		50 MG		1	10/14/2005	12/31/2013						
52959-0838-06		Q0144		11/22/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6 EA	BO	PO	EA		1 GM		0.25	11/22/2005	99/99/9999						
52959-0914-30		Q0169		11/26/2007	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 12.5 MG	30 EA	BO	PO	EA		12.5 MG		1	11/26/2007	99/99/9999						
52959-0927-03		Q0144		04/24/2008	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	3 EA	BO	PO	EA		1 GM		0.5	04/24/2008	02/03/2016						
52959-0928-30		J8999		05/15/2008	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 20 MG	30 EA	NA	PO	EA		1 EA		1	05/15/2008	99/99/9999						
52959-0932-30		Q0144		05/23/2008	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (1X30ML CHERRY) 200 MG/5 ML	30 ML	BO	PO	ML		1 GM		0.04	05/23/2008	99/99/9999						
53014-0250-01		J7510		01/01/2002	05/21/2012	PREDNISOLONE ORAL, PER 5 MG	PEDIAPRED (SF,DYE-FREE,RASPBERRY) 5 MG/5 ML	120 ML	BO	PO	ML		5 MG		0.2	01/01/2002	05/21/2012						
53100-0128-22		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SOMINEX 25 MG	16 EA	NA	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
53100-0128-32		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SOMINEX 25 MG	32 EA	NA	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
53100-0128-51		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SOMINEX 25 MG	72 EA	NA	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
53100-0128-75		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SOMINEX 50 MG	16 EA	NA	PO	EA		50 MG		1	01/01/2002	99/99/9999						
54092-0376-01		Q0173		08/29/2003	99/99/9999	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 300 MG	100 EA	BO	PO	EA		250 MG		1.2	08/29/2003	99/99/9999						
54092-0700-01		J1743		01/01/2008	99/99/9999	INJECTION, IDURSULFASE, 1 MG	ELAPRASE (PF) 2 MG/ML	3 ML	VL	IV	ML		1 MG		2	01/01/2008	99/99/9999						
54482-0053-01		J8999		01/01/2002	03/29/2018	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MATULANE 50 MG	100 EA	BO	PO	EA		1 EA		1	01/01/2002	03/29/2018						
69097-0319-87		J7626		11/14/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.5 MG/2 ML	2 ML	AM	IH	ML		0.5 MG		0.5	11/14/2017	99/99/9999						
54482-0147-01		J1955		01/01/2002	99/99/9999	INJECTION, LEVOCARNITINE, PER 1 GM	CARNITOR (S.D.V.) 200 MG/ML	5 ML	VL	IV	ML		1 GM		0.2	01/01/2002	99/99/9999						
54569-0239-00		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
54569-0239-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	24 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
54569-0239-02		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
54569-0239-03		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
54569-0239-08		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	6 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
54569-0241-00		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30 EA	BO	PO	EA		50 MG		1	01/01/2002	99/99/9999						
54569-0241-02		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15 EA	BO	PO	EA		50 MG		1	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
54569-0241-03		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999							
54569-0241-05		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	10	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999							
54569-0322-00	J8540			01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	12	EA	BO	PO	EA	0.25	MG	3	01/01/2006	99/99/9999							
54569-0322-03	J8540			01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	20	EA	BO	PO	EA	0.25	MG	3	01/01/2006	99/99/9999							
54569-0324-04	J8540			01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	6	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999							
54569-0327-00	J7509			01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	MEDROL (UNIT OF USE) 4 MG	21	EA	DP	PO	EA	4	MG	1	01/01/2002	99/99/9999							
54569-0330-00	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	21	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
54569-0330-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	50	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
54569-0330-03	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
54569-0330-04	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	30	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
54569-0330-07	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	60	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015							
54569-0331-00	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015							
54569-0331-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	15	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015							
54569-0331-02	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015							
54569-0331-04	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	50	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015							
54569-0331-05	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015							
54569-0331-07	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015							
54569-0331-08	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	40	EA	BO	PO	EA	5	MG	2	01/01/2002	12/31/2015							
54569-0332-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	10	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015							
54569-0332-02	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	21	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015							
54569-0332-03	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015							
54569-0332-05	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	100	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015							
54569-0332-09	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	18	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015							
54569-0333-00	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 50 MG	8	EA	BO	PO	EA	5	MG	10	01/01/2002	12/31/2015							
54569-0336-01	J8540			01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 2 MG	6	EA	BO	PO	EA	0.25	MG	8	01/01/2006	99/99/9999							
54569-0350-05	Q0164			01/01/2002	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	6	EA	BO	PO	EA	5	MG	1	01/01/2002	99/99/9999							
54569-0355-00	Q0165			01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	30	EA	BO	PO	EA	10	MG	1	12/07/2005	12/31/2013	01/01/2002	01/31/2003		1			
54569-0355-02	Q0165			01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013							
54569-1036-00	J7509			01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21	EA	DP	PO	EA	4	MG	1	01/01/2002	99/99/9999							
54569-1046-00	Q0170			01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120	ML	BO	PO	ML	25	MG	0.05	01/01/2002	12/31/2013							
54569-1335-00	J7510			01/01/2002	11/08/2012	PREDNISOLONE ORAL, PER 5 MG	PEDIAPRED 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	01/01/2002	11/08/2012							
54569-1377-00	J0696			01/01/2002	01/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	ROCEPHIN (VIAL) 500 MG	1	EA	VL	IJ	EA	250	MG	2	01/01/2002	01/31/2014							
54569-1387-00	J2010			01/01/2002	99/99/9999	INJECTION, LINCOSAMYIN HCL, UP TO 300 MG	LINCOCIN (VIAL) 300 MG/ML	10	ML	VL	IJ	ML	300	MG	1	01/15/2004	99/99/9999	01/01/2002	01/31/2003		1			
54569-1411-00	J1080			01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML	10	ML	VL	IM	ML	200	MG	1	01/15/2004	12/31/2014	01/01/2002	01/31/2003		1			
54569-1522-00	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (AMP) 0.9%	10	ML	AM	IJ	ML	10	ML	0.1	01/01/2004	99/99/9999							
54569-1555-00	J2930			01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	SOLU-MEDROL (ACT-O-VIAL) 125 MG	1	EA	VL	IJ	EA	125	MG	1	05/23/2007	99/99/9999	01/01/2002	01/31/2003		1			
54569-1555-01	J2930			06/05/2002	02/03/2016	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	SOLU-MEDROL (ACT-O-VIAL) 125 MG	1	EA	VL	IJ	EA	125	MG	1	06/05/2002	02/03/2016							
54569-1754-00	Q0170			01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013							
54569-1754-01	Q0170			01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013							
54569-1754-05	Q0170			12/07/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	BO	PO	EA	25	MG	1	12/07/2007	12/31/2013							
54569-1754-06	Q0170			07/02/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	25	MG	1	07/02/2002	12/31/2013							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54569-1754-09		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30 EA	BO	PO	EA		25 MG		1	01/01/2002	12/31/2013						
54569-1818-08		None		10/20/2000	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	32 EA	NA	PO	EA		2.5 MG		1	10/20/2000	99/99/9999						
54569-1827-01		J3301		01/01/2002	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG-10 (VIAL) 10 MG/ML	5 ML	VL	IJ	ML		10 MG		1	01/15/2004	99/99/9999	01/01/2002	01/31/2003				1
54569-1901-01		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (M.D.V.) 40 MG/ML	5 ML	VL	IJ	ML		40 MG		1	01/15/2004	99/99/9999	01/01/2002	01/31/2003				1
54569-2318-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN N (VIAL) 100 U/ML	10 ML	VL	SC	ML		5 U		20	01/01/2003	99/99/9999						
54569-2319-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN R (VIAL) 100 U/ML	10 ML	VL	IJ	ML		5 U		20	01/01/2003	99/99/9999						
54569-2353-05		Q0177		01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30 EA	BO	PO	EA		25 MG		1	01/01/2002	99/99/9999						
54569-2571-01		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20 EA	BO	PO	EA		50 MG		1	09/01/2005	12/31/2013	01/01/2002	06/10/2003				1
54569-2580-00		J1000		01/01/2002	10/17/2016	INJECTION, DEPO-ESTRADIOL CYPIONATE, UP TO 5 MG	DEPO-ESTRADIOL 5 MG/ML	5 ML	VL	IM	ML		5 MG		1	01/01/2004	10/17/2016	01/01/2002	01/31/2003				1
54569-2646-00		J3355		01/01/2006	99/99/9999	INJECTION, UROFOLLITROPIN, 75 IU	METRODIN 75 IU	1 EA	NA	IM	EA		75 IU		1	01/01/2006	99/99/9999						
54569-2918-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN 70/30 (VIAL) 70 U/ML-30 U/ML	10 ML	VL	SC	ML		5 U		20	01/01/2003	99/99/9999						
54569-2918-02		J1815		09/22/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN 70/30 (10X10ML) 70 U/ML-30 U/ML	10 ML	VL	SC	ML		5 U		20	09/22/2003	99/99/9999						
54569-3043-00		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	20 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
54569-3043-01		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	12 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
54569-3043-02		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	6 EA	BO	PO	EA		5 MG		4	11/17/2003	12/31/2015	01/01/2002	06/10/2003				4
54569-3043-05		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	14 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
54569-3043-06		J7506		11/07/2006	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	25 EA	BO	PO	EA		5 MG		4	11/07/2006	12/31/2015						
54569-3078-00		A4216		01/18/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE/RESPIRATORY THERAPY 0.9%	5 ML	VL	IH	ML		10 ML		0.1	01/18/2007	99/99/9999						
54569-3260-00		J3490		01/01/2002	02/03/2016	UNCLASSIFIED DRUGS	MARCAINE HCL (M.D.V.) 0.25%	50 ML	VL	IJ	ML		1 EA		1	01/01/2002	02/03/2016						
54569-3302-00		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	60 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
54569-3302-01		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
54569-3413-00		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	21 EA	DP	PO	EA		5 MG		1	01/01/2002	12/31/2015						
54569-3467-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN 70/30 70 U/ML-30 U/ML	10 ML	VL	SC	ML		5 U		20	01/01/2003	99/99/9999						
54569-3504-00		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	8 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
54569-3504-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	10 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
54569-3701-00		J1055		01/15/2004	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	DEPO-PROVERA CONTRACEPTIVE (VIAL) 150 MG/ML	1 ML	VL	IM	ML		150 MG		1	01/15/2004	12/31/2012						
54569-3704-00		J3030		01/01/2002	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX (S.D.V.) 6 MG/0.5 ML	0.5 ML	VL	SC	ML		6 MG		2	01/01/2002	99/99/9999						
54569-3765-01		J8999		10/20/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 10 MG	60 EA	BO	PO	EA		1 EA		1	10/20/2005	99/99/9999						
54569-3833-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN R (VIAL) 100 U/ML	10 ML	VL	IJ	ML		5 U		20	01/26/2004	99/99/9999	01/01/2003	06/10/2003				20
54569-3835-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN N (VIAL) 100 U/ML	10 ML	VL	SC	ML		5 U		20	09/22/2003	99/99/9999	01/01/2003	06/10/2003				20
54569-3899-00		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3 ML	PC	IH	ML		1 MG		0.83	04/01/2008	99/99/9999						
54569-3899-00	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3 ML	PC	IH	ML		1 MG		0.83	04/01/2008	99/99/9999						
54569-3900-00		J7611		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE 0.5%	20 ML	BO	IH	ML		1 MG		5	04/01/2008	99/99/9999						
54569-3946-00		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (VIAL) 40 MG/ML	1 ML	VL	IJ	ML		40 MG		1	01/22/2004	99/99/9999	01/01/2002	01/31/2003				1
54569-4112-00		J2300		01/01/2002	02/03/2016	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HYDROCHLORIDE (10X1ML) 20 MG/ML	1 ML	NA	IJ	ML		10 MG		2	01/01/2002	02/03/2016						
54569-4168-00		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	5 EA	BO	PO	EA		25 MG		1	01/01/2002	12/31/2013						
54569-4197-00		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (AF) 12.5 MG/5 ML	120 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
54569-4230-00		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.04	01/01/2002	99/99/9999						
54569-4232-00		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 100 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.02	01/01/2002	99/99/9999						
54569-4265-00		J1030		01/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL (M.D.V.) 40 MG/ML	10 ML	VL	IJ	ML		40 MG		1	01/15/2004	99/99/9999	01/01/2002	01/31/2003				1
69097-0321-87		J7626		11/14/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML, SINGLE-DOSE) 1 MG/2 ML	2 ML	AM	IH	ML		0.5 MG		1	11/14/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54569-4482-00		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4482-01		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	50 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4482-04		J8499		09/11/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40 EA	BO PO EA	EA			1 EA		1	01/01/2005	99/99/9999	09/11/2002	06/10/2003	1			
54569-4482-06		J8499		04/26/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	21 EA	BO PO EA	EA			1 EA		1	04/26/2005	99/99/9999						
54569-4497-00		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX Z-PAK 250 MG	6 EA	DP PO EA	EA			1 GM		0.25	01/01/2002	99/99/9999						
54569-4522-00		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	4 EA	BO PO EA	EA			1 GM		0.25	01/01/2002	99/99/9999						
54569-4522-01		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	2 EA	BO PO EA	EA			1 GM		0.25	01/01/2002	99/99/9999						
54569-4522-02		Q0144		08/26/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	30 EA	BO PO EA	EA			1 GM		0.25	01/05/2004	99/99/9999	08/26/2002	06/10/2003	0.25			
54569-4567-00		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX (SINGLE DOSE PACKETS) 1 GM/PACKET	1 EA	BX PO EA	EA			1 GM		1	01/01/2002	99/99/9999						
54569-4648-00		J1100		01/01/2002	02/03/2016	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (25X5ML) 4 MG/ML	5 ML	NA IJ ML	ML			1 MG		4	01/01/2002	02/03/2016						
54569-4720-00		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12 EA	BX RC EA	EA			1 EA		1	01/01/2006	99/99/9999						
54569-4720-02		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	3 EA	BX RC EA	EA			1 EA		1	01/01/2006	99/99/9999						
54569-4724-00		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	35 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4734-00		J1610		01/01/2002	99/99/9999	INJECTION, GLUCAGON HYDROCHLORIDE, PER 1 MG	GLUCAAGON EMERGENCY KIT 1 MG	1 EA	VL IJ EA	EA			1 MG		1	01/01/2002	99/99/9999						
54569-4748-00		J7614		04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.021%	3 ML	PC IH ML	ML			0.5 MG		0.42	04/01/2008	99/99/9999						
54569-4748-00	KO	J7614	KO	04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.021%	3 ML	PC IH ML	ML			0.5 MG		0.42	04/01/2008	99/99/9999						
54569-4765-01		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	14 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4765-02		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4765-03		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	50 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4765-04		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4765-05		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	45 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4765-06		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	60 EA	BO PO EA	EA			1 EA		1	01/01/2002	99/99/9999						
54569-4765-09		J8499		06/01/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	45 EA	BO PO EA	EA			1 EA		1	06/01/2006	99/99/9999						
54569-4904-00		J1055		01/15/2004	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	DEPO-PROVERA (SRN, PREFILLED) 150 MG/ML	1 ML	SR IM ML	ML			150 MG		1	01/15/2004	12/31/2012						
54569-4910-00		J7644		01/01/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (VIAL) 0.02%	2.5 ML	PC IH ML	ML			1 MG		0.2	01/01/2002	99/99/9999						
54569-4910-00	KO	J7644	KO	01/01/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (VIAL) 0.02%	2.5 ML	PC IH ML	ML			1 MG		0.2	01/01/2002	99/99/9999						
45963-0637-49		J9263		08/03/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF, LATEX-FREE) 5 MG/1 ML	10 ML	VL IV ML	ML			0.5 MG		10	08/03/2018	99/99/9999						
45963-0638-58		J9263		08/03/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF, LATEX-FREE) 5 MG/1 ML	20 ML	VL IV ML	ML			0.5 MG		10	08/03/2018	99/99/9999						
54569-4930-00		J2941		01/01/2002	99/99/9999	INJECTION, SOMATROPIN, 1 MG	SAIZEN (VIAL, W/DILUENT) 5 MG	1 EA	VL SC EA	EA			1 MG		5	01/01/2002	99/99/9999						
54569-5247-00		J2310		01/01/2002	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (VIAL, FLIPTOP) 0.4 MG/ML	1 ML	VL IJ ML	ML			1 MG		0.4	01/01/2002	99/99/9999						
54569-5311-00		J3490		01/01/2002	02/03/2016	UNCLASSIFIED DRUGS	ENGERIX-B PEDIATRIG (S.D.V., TAX INCL, PF) 10 MCG/0.5 ML	0.5 ML	VL IM ML	ML			1 EA		1	01/01/2002	02/03/2016						
54569-5312-00		J2001		11/08/2007	02/03/2016	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL 2%	5 ML	SR IJ ML	ML			10 MG		2	11/08/2007	02/03/2016						
54569-5312-01		J2001		11/08/2007	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (5X5ML) 2%	5 ML	SR IJ ML	ML			10 MG		2	11/08/2007	99/99/9999						
54569-5408-00		J3490		07/18/2002	99/99/9999	UNCLASSIFIED DRUGS	ENGERIX-B (TIP-LOK W/O NDL, TAX, PF) 20 MCG/ML	1 ML	SR IM ML	ML			1 EA		1	07/18/2002	99/99/9999						
54569-5445-00		J7614		04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.042%	3 ML	VL IH ML	ML			0.5 MG		0.84	04/01/2008	99/99/9999						
54569-5445-00	KO	J7614	KO	04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.042%	3 ML	VL IH ML	ML			0.5 MG		0.84	04/01/2008	99/99/9999						
54569-5448-00		Q0144		09/09/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX TRI-PAK 500 MG	1 EA	DP PO EA	EA			1 GM		0.5	09/09/2002	99/99/9999						
54569-5527-00		J1055		08/15/2003	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	DEPO-PROVERA CONTRACEPTIVE 150 MG/ML	1 ML	VL IM ML	ML			150 MG		1	08/15/2003	12/31/2012						
54569-5533-00		J3420		09/19/2003	01/28/2013	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (M.D.V.) 1000 MCG/ML	30 ML	VL IM ML	ML			1000 MCG		1	09/19/2003	01/28/2013						
54569-5578-00		J3490		07/21/2004	02/03/2016	UNCLASSIFIED DRUGS	TWINRIX (TIP-LOK SYRINGE) 720 EL U/ML-20 MCG/ML	1 ML	SR IM ML	ML			1 EA		1	07/21/2004	02/03/2016						
54569-5589-00		Q0173		08/26/2004	99/99/9999	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 300 MG	12 EA	BO PO EA	EA			250 MG		1.2	08/26/2004	99/99/9999						
54569-5589-01		Q0173		09/02/2005	99/99/9999	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE HCL 300 MG	6 EA	BO PO EA	EA			250 MG		1.2	09/02/2005	99/99/9999						
54569-5605-00		J1815		02/16/2006	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	LANTUS 100 U/ML	10 ML	VL SC ML	ML			5 U		20	02/16/2006	99/99/9999						
54569-5610-00		J0150		09/30/2004	12/31/2014	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS, INSTEAD USE A9270)	ADENOSINE 3 MG/ML	2 ML	NA IV ML	ML			6 MG		0.5	09/30/2004	12/31/2014						
54569-5629-00		J3490		11/10/2004	02/03/2016	UNCLASSIFIED DRUGS	RECOMBIVAX HB PEDIATRIG/ADOLESCENT (S.D.V., TAX INCL, PF) 5 MCG/0.5 ML	0.5 ML	VL IM ML	ML			1 EA		1	11/10/2004	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54569-5630-00	J3490			11/10/2004	02/03/2016	UNCLASSIFIED DRUGS	RECOMBIVAX HB (S.D.V., TAX INCL) 10 MCG/ML	1 ML	VL	IM	ML		1 EA		1	11/10/2004	02/03/2016						
54569-5715-00	J8999			07/15/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDROXYUREA 500 MG	100 EA	BO	PO	EA		1 EA		1	07/15/2005	99/99/9999						
54569-5720-00	J0696			07/26/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 250 MG	1 EA	VL	IJ	EA		250 MG		1	07/26/2005	99/99/9999						
54569-5721-00	J0696			07/26/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 500 MG	1 EA	VL	IJ	EA		250 MG		2	07/26/2005	99/99/9999						
54569-5722-00	J0696			07/26/2005	10/01/2012	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/26/2005	10/01/2012						
54569-5723-00	J0696			07/27/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 250 MG	1 EA	VL	IJ	EA		250 MG		1	07/27/2005	99/99/9999						
54569-5724-00	J0696			07/27/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 500 MG	1 EA	VL	IJ	EA		250 MG		2	07/27/2005	99/99/9999						
54569-5725-00	J0696			07/27/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/27/2005	99/99/9999						
54569-5729-00	J8540			01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	28 EA	BO	PO	EA		0.25 MG		16	01/01/2006	99/99/9999						
54569-5741-00	J8501			10/24/2005	99/99/9999	APREPITANT, ORAL, 5 MG	EMEND TRI-FOLD PACK	3 EA	PG	PO	EA		5 MG		19	10/24/2005	99/99/9999						
54569-5744-00	J8498			01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 12.5 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
54569-5744-01	J8498			01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 12.5 MG	6 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
54569-5745-00	J8498			01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 25 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
54569-5745-01	J8498			01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 25 MG	4 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
54569-5745-02	J8498			01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 25 MG	6 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
54569-5754-00	Q0144			11/24/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	4 EA	BO	PO	EA		1 GM		0.25	11/24/2005	99/99/9999						
54569-5755-00	Q0144			11/24/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6 EA	DP	PO	EA		1 GM		0.25	11/24/2005	99/99/9999						
54569-5756-00	Q0144			11/24/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	3 EA	DP	PO	EA		1 GM		0.5	11/24/2005	99/99/9999						
54569-5764-00	J2792			01/12/2006	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT DETERGENT, 100 IU	HYPERRHO S/D (FULL DOSE)	1 ML	SR	IM	ML		100 IU		15	01/12/2006	99/99/9999						
54569-5781-00	J1324			01/01/2007	10/17/2016	INJECTION, ENFUVIRTIDE, 1 MG	FUZEON 90 MG	60 EA	PG	SC	EA		1 MG		90	01/01/2007	10/17/2016						
61314-0318-10	Q5101			07/20/2018	99/99/9999	INJECTION, FILGRASTIM-SDZ, BIOSIMILAR, (ZARXIO), 1 MICROGRAM	ZARXIO (PF) 300 MCG/0.5 ML	0.5 ML	SR	IJ	ML		1 MCG		600	07/20/2018	99/99/9999						
54569-5795-00	J2300			05/12/2006	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL (10X1ML) 10 MG/ML	1 ML	AM	IJ	ML		10 MG		1	05/12/2006	99/99/9999						
54569-5804-00	Q0144			06/30/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 600 MG	8 EA	BO	PO	EA		1 GM		0.6	06/30/2006	99/99/9999						
54569-5806-00	Q0144			07/24/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 1 GM/Package	1 EA	BX	PO	EA		1 GM		1	07/24/2006	99/99/9999						
54569-5807-00	Q0144			07/24/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 100 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.02	07/24/2006	99/99/9999						
54569-5808-00	Q0144			07/24/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.04	07/24/2006	99/99/9999						
54569-5809-00	Q0144			07/24/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	22.5 ML	BO	PO	ML		1 GM		0.04	07/24/2006	99/99/9999						
54569-5810-00	Q0144			07/25/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	30 ML	BO	PO	ML		1 GM		0.04	07/25/2006	99/99/9999						
54569-5815-00	J1200			08/03/2006	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HYDROCHLORIDE (25X1ML) 50 MG/ML	1 ML	VL	IJ	ML		50 MG		1	08/03/2006	99/99/9999						
54569-5828-00	J1460			09/26/2006	99/99/9999	INJECTION, GAMMA GLOBULIN, INTRAMUSCULAR, 1 CC	GAMASTAN S/D (SDV)	2 ML	VL	IM	ML		1 ML		1	09/26/2006	99/99/9999						
54569-5840-00	J7506			10/10/2006	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21 EA	BO	PO	EA		5 MG		2	10/10/2006	12/31/2015						
54569-5841-00	J7506			10/10/2006	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	48 EA	BO	PO	EA		5 MG		2	10/10/2006	12/31/2015						
54569-5857-00	J8999			11/06/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 20 MG	30 EA	BO	PO	EA		1 EA		1	11/06/2006	99/99/9999						
54569-5862-00	J3490			11/13/2006	09/07/2016	UNCLASSIFIED DRUGS	PROPOFOL (SDV, 5X20ML) 10 MG/ML	20 ML	VL	IJ	ML		1 EA		1	11/13/2006	09/07/2016						
54569-5874-00	J2405			01/12/2007	03/14/2016	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (5X2ML,SDV) 2 MG/ML	2 ML	VL	IJ	ML		1 MG		2	01/12/2007	03/14/2016						
54569-5911-00	J7506			05/10/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (PACK) 5 MG	48 EA	BO	PO	EA		5 MG		1	05/10/2007	12/31/2015						
54746-0001-01	J9215			01/01/2002	99/99/9999	INJECTION, INTERFERON, ALFA-N3, (HUMAN LEUKOCYTE DERIVED), 250,000 IU	ALFERON N (M.D.V.) 5 Million IU/ML	1 ML	VL	IJ	ML		250000 IU		20	01/01/2002	99/99/9999						
54838-0135-40	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SILADRYL ALLERGY (AF,SF) 12.5 MG/5 ML	118 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
54838-0135-70	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SILADRYL ALLERGY 12.5 MG/5 ML	237 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
54838-0135-80	Q0163			01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SILADRYL ALLERGY (AF,SF) 12.5 MG/5 ML	473 ML	BO	PO	ML		50 MG		0.05	01/01/2002	99/99/9999						
54838-0154-40	Q0163			01/01/2002	03/01/2018	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SILPHEN 12.5 MG/5 ML	118 ML	BO	PO	ML		50 MG		0.05	01/01/2002	03/01/2018						
54838-0154-70	Q0163			01/01/2002	03/01/2018	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SILPHEN 12.5 MG/5 ML	237 ML	BO	PO	ML		50 MG		0.05	01/01/2002	03/01/2018						
54838-0154-80	Q0163			01/01/2002	03/01/2018	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SILPHEN 12.5 MG/5 ML	473 ML	BO	PO	ML		50 MG		0.05	01/01/2002	03/01/2018						
54868-0007-00	J1200			01/01/2002	02/03/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	BENADRYL (VIAL) 50 MG/ML	10 ML	AM	IJ	ML		50 MG		1	01/01/2002	02/03/2016						
54868-0015-00	J1265			12/11/2006	02/03/2016	INJECTION, DOPAMINE HCL, 40 MG	DOPAMINE HYDROCHLORIDE 80 MG/ML	125 ML	NA	IV	ML		40 MG		2	12/11/2006	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-0026-00		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
54868-0026-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
54868-0026-04		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	1000	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
54868-0026-05		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	10	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
54868-0026-06		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
54868-0026-07		Q0163		06/29/2006	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60	EA	BO	PO	EA	50	MG	0.5	06/29/2006	99/99/9999						
54868-0102-00		J1720		12/11/2006	02/03/2016	RINGERS LACTATE INFUSION, UP TO 1000 CC	LACTATED RINGERS (12X1000ML)	1000	ML	PC	IV	ML	1000	ML	0.001	12/11/2006	02/03/2016						
54868-0163-02		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOVIRAX 200 MG	25	EA	BO	PO	EA	1	EA	1	01/01/2002	02/03/2016						
54868-0169-01		Q0177		01/01/2002	02/03/2016	REGIMEN	VISTARIL 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	02/03/2016						
54868-0173-00		J9250		03/26/2003	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (PF) 25 MG/ML WATER FOR INJECTION BACTERIOSTATIC (VIAL)	2	ML	EA	IJ	ML	5	MG	5	03/26/2003	99/99/9999						
54868-0183-00		A4216		01/01/2004	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	30	ML	VL	IV	ML	10	ML	0.1	01/01/2004	02/03/2016						
54868-0186-00		J0595		01/01/2004	02/03/2016	INJECTION, BUTORPHANOL TARTRATE, 1 MG	STADOL (M.D.V.) 2 MG/ML	10	ML	VL	IJ	ML	1	MG	2	01/01/2004	02/03/2016						
54868-0206-00		J0702		01/01/2002	02/03/2016	INJECTION, BETAMETHASONE ACETATE 3MG AND BETAMETHASONE SODIUM PHOSPHATE 3MG	CELESTONE SOLUSPAN (M.D.V.) 3 MG/ML 3 MG/ML	5	ML	VL	IJ	ML	3	MG	1	01/01/2002	02/03/2016						
54868-0216-00		J1080		09/20/2007	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML	10	ML	VL	IM	ML	200	MG	1	09/20/2007	12/31/2014						
54868-0218-00		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	20	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999						
54868-0218-01		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	10	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999						
54868-0218-03		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	3	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999						
54868-0218-04		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	30	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999						
54868-0218-05		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	16	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999						
54868-0218-06		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	100	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999						
54868-0218-07		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	40	EA	BO	PO	EA	0.25	MG	16	01/01/2006	99/99/9999						
54868-0218-08		J8540		09/11/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (USP) 4 MG	50	EA	BO	PO	EA	0.25	MG	16	09/11/2006	99/99/9999						
54868-0218-09		J8540		04/03/2008	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	5	EA	BO	PO	EA	0.25	MG	16	04/03/2008	99/99/9999						
54868-0231-00		J3410		01/01/2002	02/03/2016	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (M.D.V.) 50 MG/ML	10	ML	VL	IM	ML	25	MG	2	01/01/2002	02/03/2016						
54868-0234-00		J3301		01/01/2002	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG-10 (VIAL) 10 MG/ML	5	ML	VL	IJ	ML	10	MG	1	01/01/2002	99/99/9999						
54868-0258-01		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	30	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
54868-0258-02		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
54868-0258-04		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	20	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
54868-0258-05		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	36	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
54868-0258-06		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	55	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
54868-0258-08		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	60	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
54868-0258-09		J7506		03/14/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	15	EA	BO	PO	EA	5	MG	1	03/14/2002	12/31/2015						
54868-0261-00		J0780		01/01/2002	06/14/2016	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE (M.D.V.) 5 MG/ML	10	ML	VL	IJ	ML	10	MG	0.5	01/01/2002	06/14/2016						
54868-0262-00		J2550		01/01/2002	02/03/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (M.D.V.) 50 MG/ML PROMETHAZINE HCL (10X25ML,MDV) 50 MG/ML	10	ML	VL	IJ	ML	50	MG	1	01/01/2002	02/03/2016						
54868-0262-01		J2550		09/29/2005	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (M.D.V.) 50 MG/ML	10	ML	VL	IJ	ML	50	MG	1	09/29/2005	99/99/9999						
54868-0296-01		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE 5%	500	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999						
54868-0296-02		J7060		01/01/2002	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE 5%	250	ML	FC	IV	ML	500	ML	0.002	01/01/2002	99/99/9999						
54868-0296-04		J7060		12/12/2006	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (48X100ML) 5%	100	ML	FC	IV	ML	500	ML	0.002	12/12/2006	99/99/9999						
54868-0554-00		J1200		01/01/2002	02/03/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	BENADRYL (AMP) 50 MG/ML	1	ML	VL	IJ	ML	50	MG	1	01/01/2002	02/03/2016						
54868-0559-00		J0690		01/01/2002	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (VIAL) 1 GM	1	EA	VL	IJ	EA	500	MG	2	01/01/2002	99/99/9999						
54868-0597-00		J2550		01/01/2002	02/03/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PHENERGAN (AMP) 25 MG/ML	1	ML	AM	IJ	ML	50	MG	0.5	01/01/2002	02/03/2016						
54868-0601-01		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 25 MG	2	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999						
54868-0601-02		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999						
54868-0605-00		J1720		01/01/2002	02/03/2016	INJECTION, HYDROCORTISONE SODIUM SUCCINATE, UP TO 100 MG	SOLU-CORTEF (S.D.V.) 100 MG	1	EA	VL	IJ	EA	100	MG	1	01/01/2002	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-1050-04		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	40	EA	BO	PO	EA	50	MG	1	01/01/2002	02/03/2016						
54868-1050-05		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	99/99/9999						
54868-1050-06		Q0163		04/15/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15	EA	NA	PO	EA	50	MG	1	04/15/2002	99/99/9999						
54868-1082-00		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	10	MG	1	01/01/2002	12/31/2013						
54868-1082-01		Q0165		01/29/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	10	MG	1	01/29/2004	12/31/2013						
54868-1082-02		Q0165		06/03/2005	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20	EA	BO	PO	EA	10	MG	1	06/03/2005	12/31/2013						
54868-1082-03		Q0165		08/24/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	10	MG	1	08/24/2007	12/31/2013						
54868-1082-04		Q0165		02/10/2005	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	10	MG	1	02/10/2005	12/31/2013						
54868-1082-05		Q0165		06/09/2005	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	10	MG	1	06/09/2005	12/31/2013						
54868-1082-06		Q0165		04/16/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	90	EA	BO	PO	EA	10	MG	1	04/16/2007	12/31/2013						
54868-1119-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 1 MG	100	EA	BO	PO	EA	5	MG	0.2	01/01/2002	12/31/2015						
54868-1119-02	J7506			12/09/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 1 MG	90	EA	BO	PO	EA	5	MG	0.2	12/09/2002	12/31/2015						
54868-1119-03	J7506			12/09/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 1 MG	30	EA	BO	PO	EA	5	MG	0.2	12/09/2002	12/31/2015						
54868-1119-04	J7506			06/01/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 1 MG	15	EA	BO	PO	EA	5	MG	0.2	06/01/2004	12/31/2015						
54868-1119-05	J7506			10/05/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 1 MG	60	EA	BO	PO	EA	5	MG	0.2	10/05/2004	12/31/2015						
54868-1126-00	J8999			08/11/2003	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN 2 MG	50	EA	BO	PO	EA	1	EA	1	08/11/2003	02/03/2016						
54868-1126-01	J8999			11/22/2005	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN 2 MG	30	EA	BO	PO	EA	1	EA	1	11/22/2005	02/03/2016						
54868-1126-02	J8999			11/22/2005	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN 2 MG	10	EA	BO	PO	EA	1	EA	1	11/22/2005	02/03/2016						
54868-1126-03	J8999			11/22/2005	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN 2 MG	25	EA	BO	PO	EA	1	EA	1	11/22/2005	02/03/2016						
54868-1126-04	J8999			05/23/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN 2 MG	5	EA	BO	PO	EA	1	EA	1	05/23/2006	02/03/2016						
54868-1126-05	J8999			10/17/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN 2 MG	100	EA	BO	PO	EA	1	EA	1	10/17/2006	02/03/2016						
54868-1183-00	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	100	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
54868-1183-01	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	15	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
54868-1183-02	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	60	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
54868-1183-03	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
54868-1183-04	J7506			01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	20	EA	BO	PO	EA	5	MG	4	01/01/2002	12/31/2015						
54868-1183-05	J7506			08/19/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	10	EA	BO	PO	EA	5	MG	4	08/19/2003	12/31/2015						
54868-1183-06	J7506			08/15/2005	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	25	EA	BO	PO	EA	5	MG	4	08/15/2005	12/31/2015						
54868-1227-00		Q0163		02/23/2006	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE (AF) 12.5 MG/5 ML	473	ML	BO	PO	ML	50	MG	0.05	02/23/2006	99/99/9999						
54868-1227-02		Q0163		10/22/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GENAHIST (AF,SF,CHERRY) 12.5 MG/5 ML	120	ML	BO	PO	ML	50	MG	0.05	10/22/2002	99/99/9999						
54868-1323-00		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
54868-1323-01		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						

NDC	NDC Mod	HPCCS	HPCCS Mod	Relationship Start Date	Relationship End Date	HPCCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HPCCS Amount #1	HPCCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-1323-02		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	BO	PO	EA	25	MG	1	07/02/2003	12/31/2013	01/01/2002	04/15/2002	1			
54868-1323-04		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
54868-1323-05		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
54868-1323-06		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
54868-1323-07		Q0170		06/15/2005	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	BO	PO	EA	25	MG	1	06/15/2005	12/31/2013						
54868-1323-08		Q0170		09/21/2005	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	50	EA	BO	PO	EA	25	MG	1	09/21/2005	12/31/2013						
54868-1366-00		J8999		04/06/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MATULANE 50 MG	100	EA	BO	PO	EA	1	EA	1	04/06/2006	99/99/9999						
54868-1367-00		J8999		08/08/2003	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDREA 500 MG	100	EA	BO	PO	EA	1	EA	1	08/08/2003	02/03/2016						
54868-1429-01		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN N 100 U/ML	10	ML	VL	SC	ML	5	U	20	01/01/2003	99/99/9999						
54868-1613-02		J8498		09/11/2006	10/17/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE (USP) 50 MG	6	EA	BX	RC	EA	1	EA	1	09/11/2006	10/17/2016						
54868-1629-00		J8999		10/03/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	100	EA	BO	PO	EA	1	EA	1	10/03/2005	99/99/9999						
54868-1629-01		J8999		10/03/2005	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	14	EA	BO	PO	EA	1	EA	1	10/03/2005	02/03/2016						
54868-1629-02		J8999		07/06/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG	30	EA	BO	PO	EA	1	EA	1	07/06/2007	99/99/9999						
54868-1720-00		J7510		01/01/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PEDIAPRED 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	01/01/2002	99/99/9999						
54868-1729-00		J1000		01/01/2002	99/99/9999	INJECTION, DEPO-ESTRADIOL CYPIONATE, UP TO 5 MG	DEPO-ESTRADIOL (VIAL) 5 MG/ML	5	ML	VL	IM	ML	5	MG	1	01/01/2002	99/99/9999						
54868-1744-00		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 1.5 MG	100	EA	BO	PO	EA	0.25	MG	6	01/01/2006	99/99/9999						
54868-1795-00		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	XYLOCAINE (M.D.V.) 1%	50	ML	VL	EP	ML	10	MG	1	01/01/2004	99/99/9999						
54868-1798-01		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	XYLOCAINE (M.D.V.) 2%	10	ML	VL	IJ	ML	10	MG	2	01/01/2004	99/99/9999						
54868-1854-00		Q0178		01/01/2002	06/30/2012	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	06/30/2012						
54868-1854-01		Q0178		01/01/2002	06/30/2012	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30	EA	BO	PO	EA	50	MG	1	01/01/2002	06/30/2012						
54868-1854-03		Q0178		01/01/2002	06/30/2012	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	60	EA	BO	PO	EA	50	MG	1	01/01/2002	06/30/2012						
54868-1854-04		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013						
54868-1867-00		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120	ML	BO	PO	ML	25	MG	0.05	01/01/2002	12/31/2013						
54868-1932-00		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 12.5 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	02/03/2016						
54868-1932-01		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 12.5 MG	1	EA	BX	RC	EA	1	EA	1	01/01/2006	02/03/2016						
54868-1932-02		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 12.5 MG	6	EA	BX	RC	EA	1	EA	1	01/01/2006	02/03/2016						
54868-1963-00		Q0174		02/11/2003	02/03/2016	THIETHYLPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TORECAN 10 MG	15	EA	BO	PO	EA	10	MG	1	02/11/2003	02/03/2016						
54868-1963-01		Q0174		02/11/2003	02/03/2016	THIETHYLPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TORECAN 10 MG	10	EA	BO	PO	EA	10	MG	1	02/11/2003	02/03/2016						
54868-2048-00		J1200		01/01/2002	02/03/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (VIAL) 50 MG/ML	1	ML	VL	IJ	ML	50	MG	1	01/01/2002	02/03/2016						
54868-2048-01		J1200		01/01/2002	02/03/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (VIAL) 50 MG/ML	1	ML	VL	IJ	ML	50	MG	1	01/01/2002	02/03/2016						
54868-2062-00		J2310		01/01/2002	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (AMP) 0.4 MG/ML	1	ML	AM	IJ	ML	1	MG	0.4	01/01/2002	99/99/9999						
54868-2064-00		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (M.D.V.) 2%	50	ML	VL	IJ	ML	10	MG	2	01/01/2004	99/99/9999						
54868-2064-01		J2001		06/23/2006	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL 2%	1250	ML	VL	IJ	ML	10	MG	2	06/23/2006	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-2088-00		J2550		09/29/2005	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL 50 MG/ML	25 ML	AM UJ	ML			50 MG		1	09/29/2005	99/99/9999						
54868-2184-00		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	100 EA	BO PO	EA			1 EA		1	01/01/2002	02/03/2016						
54868-2184-02		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	30 EA	BO PO	EA			1 EA		1	01/01/2002	02/03/2016						
54868-2184-03		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	25 EA	BO PO	EA			1 EA		1	01/01/2002	02/03/2016						
54868-2184-04		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	50 EA	BO PO	EA			1 EA		1	01/01/2002	02/03/2016						
54868-2219-00		J3490		01/01/2002	02/03/2016	UNCLASSIFIED DRUGS	RECOMBIVAX HB (3 DOSE VIAL, TAX INCL) 10 MCG/ML	3 ML	VL IM	ML			1 EA		1	01/01/2002	02/03/2016						
54868-2219-01		J3490		01/01/2002	02/03/2016	UNCLASSIFIED DRUGS	RECOMBIVAX HB (S.D.V., TAX INCL) 10 MCG/ML	1 ML	VL IM	ML			1 EA		1	01/01/2002	02/03/2016						
54868-2299-00		J1940		09/29/2005	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (ABBOJECT) 10 MG/ML	250 ML	VL UJ	ML			20 MG		0.5	09/29/2005	99/99/9999						
54868-2302-00		Q0172		01/01/2002	12/31/2013	CHLORPROMAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 50 MG	10 EA	BO PO	EA			25 MG		2	01/01/2002	12/31/2013						
54868-2302-02		Q0172		01/01/2002	12/31/2013	CHLORPROMAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 50 MG	100 EA	BO PO	EA			25 MG		2	01/01/2002	12/31/2013						
54868-2320-01		J3360		01/01/2002	02/03/2016	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM 5 MG/ML	2 ML	SR UJ	ML			5 MG		1	01/01/2002	02/03/2016						
54868-2320-02		J3360		01/01/2002	02/03/2016	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (AMP) 5 MG/ML	2 ML	AM UJ	ML			5 MG		1	01/01/2002	02/03/2016						
54868-2347-00		Q0172		01/01/2002	12/31/2013	CHLORPROMAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 100 MG	100 EA	BO PO	EA			25 MG		4	01/01/2002	12/31/2013						
54868-2380-01		J1815		07/16/2007	02/03/2016	INJECTION, INSULIN, PER 5 UNITS	NOVOLIN N 100 U/ML	10 ML	VL SC	ML			5 U		20	07/16/2007	02/03/2016						
54868-2429-01		J0515		01/01/2002	99/99/9999	INJECTION, BENZTROPINE MESYLATE, PER 1 MG	COGENTIN (AMP) 1 MG/ML	2 ML	AM UJ	ML			1 MG		1	01/01/2002	99/99/9999						
54868-2464-00		Q0172		01/01/2002	12/31/2013	CHLORPROMAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 25 MG	30 EA	BO PO	EA			25 MG		1	01/01/2002	12/31/2013						
54868-2464-02		Q0172		08/08/2007	12/31/2013	CHLORPROMAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 25 MG	60 EA	NA PO	EA			25 MG		1	08/08/2007	12/31/2013						
54868-2472-00		J7613		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3 ML	PC IH	ML			1 MG		0.83	04/01/2008	99/99/9999						
54868-2472-00	KO	J7613	KO	04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3 ML	PC IH	ML			1 MG		0.83	04/01/2008	99/99/9999						
54868-2472-01		J7611		04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE 0.5%	3 ML	PC IH	ML			1 MG		5	04/01/2008	99/99/9999						
54868-2489-01		J3411		01/01/2004	99/99/9999	INJECTION, THIAMINE HCL, 100 MG	THIAMINE HCL 100 MG/ML	2 ML	VL UJ	ML			100 MG		1	01/01/2004	99/99/9999						
54868-2522-00		J1440		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 300 MCG	NEUPOGEN (S.D.V.,PF) 300 MCG/ML	1 ML	VL UJ	ML			300 MCG		1	01/01/2002	12/31/2013						
54868-2523-00		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (S.D.V.) 10000 U/ML	1 ML	VL UJ	ML			1000 U		10	01/01/2006	99/99/9999						
54868-2523-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (S.D.V.) 10000 U/ML	1 ML	VL UJ	ML			1000 U		10	01/01/2006	99/99/9999						
54868-2526-00		J1642		01/01/2002	06/30/2015	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEP-LOCK (VIAL,DOSETTE) 100 U/ML	1 ML	VL IV	ML			10 U		10	01/01/2002	06/30/2015						
54868-2527-00		A4216		06/28/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (150X5ML) 0.9%	5 ML	SR IV	ML			10 ML		0.1	06/28/2007	02/03/2016						
54868-2528-00		J2545		01/01/2007	02/03/2016	PENTAMIDINE ISETHIONATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MG	NEBUPENT (S.D.V.,PF) 300 MG	1 EA	VL IH	EA			300 MG		1	01/01/2007	02/03/2016						
54868-2530-00		J3070		01/01/2002	02/03/2016	INJECTION, PENTAZOCINE, 30 MG	TALWIN LACTATE (VIAL) 30 MG/ML	10 ML	VL UJ	ML			30 MG		1	01/01/2002	02/03/2016						
54868-2652-00		J3030		01/01/2002	02/03/2016	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX (S.D.V.) 6 MG/0.5 ML	0.5 ML	VL SC	ML			6 MG		2	01/01/2002	02/03/2016						
54868-2684-01		Q0171		02/01/2007	12/31/2013	CHLORPROMAZINE HYDROCHLORIDE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE 10 MG	30 EA	BO PO	EA			10 MG		1	02/01/2007	12/31/2013						
54868-2686-00		Q0175		01/01/2002	02/03/2016	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 4 MG	30 EA	BO PO	EA			4 MG		1	01/01/2002	02/03/2016						
54868-2687-01		Q0176		01/01/2002	12/31/2013	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	100 EA	BO PO	EA			8 MG		1	01/01/2002	12/31/2013						
67457-0372-99		J1644		05/25/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X1ML) 1000 U/1 ML	1 ML	VL UJ	ML			1000 U		1	05/25/2018	99/99/9999						
54868-2687-02		Q0176		06/12/2007	12/31/2013	PERPHENAZINE, 8MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	60 EA	BO PO	EA			8 MG		1	06/12/2007	12/31/2013						
54868-2746-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN 70/30 (VIAL) 70 U/ML-30 U/ML	10 ML	VL SC	ML			5 U		20	01/01/2003	99/99/9999						
54868-2777-00		J1817		05/07/2007	02/03/2016	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	NOVOLOG 100 U/ML	10 ML	VL SC	ML			50 U		2	05/07/2007	02/03/2016						
54868-2825-00		J1950		03/10/2003	02/03/2016	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT 3.75 MG	1 EA	BX IM	EA			3.75 MG		1	03/10/2003	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-2844-00		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	60	EA	BO	PO	EA	25	MG	2	01/01/2002	12/31/2013						
54868-2844-01		Q0170		04/21/2008	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	30	EA	BO	PO	EA	25	MG	2	04/21/2008	12/31/2013						
54868-2889-00		J1631		01/01/2002	02/03/2016	INJECTION, HALOPERIDOL DECANOATE, PER 50 MG	HALDOL DECANOATE (AMP) 50 MG/ML	1	ML	AM	IM	ML	50	MG	1	01/01/2002	02/03/2016						
54868-2889-01		J1631		01/01/2002	02/03/2016	INJECTION, HALOPERIDOL DECANOATE, PER 50 MG	HALDOL DECANOATE (AMP) 50 MG/ML	1	ML	AM	IM	ML	50	MG	1	01/01/2002	02/03/2016						
54868-2892-00		Q0177		01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	99/99/9999						
54868-2892-03		Q0177		09/19/2005	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	BO	PO	EA	25	MG	1	09/19/2005	99/99/9999						
54868-2892-04		Q0177		10/11/2005	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	15	EA	BO	PO	EA	25	MG	1	10/11/2005	99/99/9999						
54868-2913-00		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	100	EA	BO	PO	EA	4	MG	1	01/01/2002	99/99/9999						
54868-2913-01		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	30	EA	BO	PO	EA	4	MG	1	01/01/2002	99/99/9999						
54868-2913-02		J7509		07/29/2003	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	60	EA	BO	PO	EA	4	MG	1	07/29/2003	99/99/9999						
54868-3004-01		J8999		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 10 MG	120	EA	BO	PO	EA	1	EA	1	01/01/2002	02/03/2016						
54868-3004-02		J8999		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE 10 MG	60	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
54868-3004-03		J8999		02/02/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (USP) 10 MG	180	EA	BO	PO	EA	1	EA	1	02/02/2006	99/99/9999						
54868-3004-04		J8999		04/10/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (USP) 10 MG	100	EA	BO	PO	EA	1	EA	1	04/10/2006	99/99/9999						
54868-3004-05		J8999		04/13/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (USP) 10 MG	30	EA	BO	PO	EA	1	EA	1	04/13/2006	99/99/9999						
54868-3025-00		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 400 MG	15	EA	BO	PO	EA	1	EA	1	01/01/2002	02/03/2016						
54868-3050-00		J1441		08/14/2006	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 480 MCG	NEUPOGEN 480 MCG/0.8 ML	10	ML	SR	IJ	ML	480	MCG	1.25	08/14/2006	12/31/2013						
54868-3084-00		Q0167		01/01/2002	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 2.5 MG	60	EA	BO	PO	EA	2.5	MG	1	01/01/2002	99/99/9999						
54868-3084-01		Q0167		02/11/2004	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL 2.5 MG	30	EA	BO	PO	EA	2.5	MG	1	02/11/2004	99/99/9999						
54868-3084-02		Q0167		01/27/2006	02/03/2016	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 2.5 MG	90	EA	BO	PO	EA	2.5	MG	1	01/27/2006	02/03/2016						
54868-3089-00		J7799		12/11/2006	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXDROSE (10X50ML) 50%	50	ML	SR	IV	ML	1	EA	1	12/11/2006	99/99/9999						
54868-3089-01		J7799		12/05/2007	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXDROSE (1X1250ML) 50%	1250	ML	GC	IV	ML	1	EA	1	12/05/2007	99/99/9999						
54868-3099-01		J8999		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGACE 40 MG/ML	240	ML	BO	PO	ML	1	EA	1	01/01/2002	02/03/2016						
54868-3112-00		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999						
54868-3112-01		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	6	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999						
54868-3134-00		J3490		01/01/2002	02/03/2016	UNCLASSIFIED DRUGS	MARCAINE HCL (S.D.V.) 0.5%	30	ML	VL	IJ	ML	1	EA	1	01/01/2002	02/03/2016						
54868-3134-01		J3490		02/02/2007	99/99/9999	UNCLASSIFIED DRUGS	MARCAINE HCL 0.5%	50	ML	VL	IJ	ML	1	EA	1	02/02/2007	99/99/9999						
54868-3157-00		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 2 MG	10	EA	BO	PO	EA	0.25	MG	8	01/01/2006	99/99/9999						
54868-3157-01		J8540		05/10/2007	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (USP, GLUTEN-FREE) 2 MG	48	EA	BO	PO	EA	0.25	MG	8	05/10/2007	99/99/9999						
54868-3181-00		J3030		01/01/2002	02/03/2016	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX (SRN) 6 MG/0.5 ML	2	ML	BX	SC	ML	6	MG	2	01/01/2002	02/03/2016						
54868-3188-00		J2820		05/23/2006	02/03/2016	INJECTION, SARGRAMOSTIM (GM-CSF), 50 MCG	LEUKINE 500 MCG/ML	5	ML	VL	IV	ML	50	MCG	10	05/23/2006	02/03/2016						
67457-0373-99		J1644		06/14/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV, 25X1ML, LATEX-FREE) 20000 U/1 ML	1	ML	VL	IJ	ML	1000	U	20	06/14/2018	99/99/9999						
54868-3189-00		Q0168		06/07/2005	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	25	EA	BO	PO	EA	5	MG	1	06/07/2005	12/31/2013						
54868-3189-01		Q0168		01/30/2006	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL 5 MG	100	EA	BO	PO	EA	5	MG	1	01/30/2006	12/31/2013						
54868-3189-02		Q0168		02/07/2006	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL 5 MG	60	EA	BO	PO	EA	5	MG	1	02/07/2006	12/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
54868-3859-01		J2560		01/01/2002	02/03/2016	INJECTION, PHENOBARBITAL SODIUM, UP TO 120 MG	PHENOBARBITAL SODIUM (TUBEX) 30 MG/ML	1	ML	SR	IJ	ML	120	MG	0.25	01/01/2002	02/03/2016							
54868-3873-00	J1800			12/11/2006	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL (S.D.V., 10X1ML) 1 MG/ML	1	ML	VL	IV	ML	1	MG	1	12/11/2006	99/99/9999							
54868-3889-00	J2597			01/01/2002	02/03/2016	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DDAVP (VIAL) 4 MCG/ML	10	ML	VL	IJ	ML	1	MCG	4	01/01/2002	02/03/2016							
54868-3890-00	J1790			01/01/2002	02/03/2016	INJECTION, DROPERIDOL, UP TO 5 MG	DROPERIDOL (AMP) 2.5 MG/ML	1	ML	AM	IJ	ML	5	MG	0.5	01/01/2002	02/03/2016							
54868-3894-00	J2001			01/01/2004	02/03/2016	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	XYLOCAINE (AMP) 2%	5	ML	AM	IJ	ML	10	MG	2	01/01/2004	02/03/2016							
54868-3896-01	J1030			05/03/2005	02/03/2016	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL 40 MG/ML	25	ML	VL	IJ	ML	40	MG	1	05/03/2005	02/03/2016							
54868-3896-02	J1030			02/02/2007	02/03/2016	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO-MEDROL 40 MG/ML	5	ML	VL	IJ	ML	40	MG	1	02/02/2007	02/03/2016							
54868-3905-00	A4217			01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION	6000	ML	FC	IV	ML	500	ML	0.002	01/01/2004	99/99/9999							
54868-3975-00	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.)	5	ML	VL	IV	ML	10	ML	0.1	01/01/2004	99/99/9999							
54868-3979-00	J0740			04/12/2006	02/03/2016	INJECTION, CIDOFOVIR, 375 MG	VISTIDE 75 MG/ML	5	ML	VL	IV	ML	375	MG	0.2	04/12/2006	02/03/2016							
67457-0602-99	J1644			05/25/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X1ML) 1000 U/1 ML	1	ML	VL	IJ	ML	1000	U	10	05/25/2018	99/99/9999							
54868-3996-00	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
54868-3996-01	J8499			01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40	EA	BO	PO	EA	1	EA	1	01/01/2002	02/03/2016							
54868-3996-02	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
54868-3996-03	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	50	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
54868-3996-04	J8499			06/17/2004	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	100	EA	BO	PO	EA	1	EA	1	06/17/2004	02/03/2016							
54868-3996-05	J8499			08/06/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	60	EA	BO	PO	EA	1	EA	1	08/06/2007	99/99/9999							
54868-3997-00	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
54868-3997-01	J8499			06/12/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100	EA	BO	PO	EA	1	EA	1	06/12/2003	99/99/9999							
54868-3997-02	J8499			09/25/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	20	EA	BO	PO	EA	1	EA	1	09/25/2003	99/99/9999							
54868-3997-03	J8499			10/20/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	10	EA	BO	PO	EA	1	EA	1	10/20/2003	99/99/9999							
54868-3997-04	J8499			11/03/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	40	EA	BO	PO	EA	1	EA	1	11/03/2003	99/99/9999							
54868-3997-05	J8499			08/01/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	60	EA	BO	PO	EA	1	EA	1	08/01/2005	99/99/9999							
54868-3998-00	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
54868-3998-01	J8499			01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	50	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999							
54868-3998-02	J8499			03/05/2003	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	15	EA	BO	PO	EA	1	EA	1	03/05/2003	02/03/2016							
54868-3998-03	J8499			12/08/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	20	EA	BO	PO	EA	1	EA	1	12/08/2003	99/99/9999							
54868-3998-04	J8499			01/28/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	40	EA	BO	PO	EA	1	EA	1	01/28/2004	99/99/9999							
54868-3998-05	J8499			06/09/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100	EA	BO	PO	EA	1	EA	1	06/09/2004	99/99/9999							
54868-3998-06	J8499			07/06/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	35	EA	BO	PO	EA	1	EA	1	07/06/2004	99/99/9999							
54868-3998-07	J8499			07/23/2004	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	500	EA	BO	PO	EA	1	EA	1	07/23/2004	02/03/2016							
54868-3998-08	J8499			04/22/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	60	EA	BO	PO	EA	1	EA	1	04/22/2005	99/99/9999							
54868-4021-00	J2550			01/01/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (AMP) 25 MG/ML	1	ML	AM	IJ	ML	50	MG	0.5	01/01/2002	99/99/9999							
54868-4047-00	J0290			01/01/2002	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (VIAL) 500 MG	1	EA	VL	IJ	EA	500	MG	1	01/01/2002	99/99/9999							
54868-4050-00	J2271			01/01/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE	1	EA	JR	NA	GM	100	MG	10	01/01/2002	12/31/2014							
54868-4076-00	Q0144			01/01/2002	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	01/01/2002	02/03/2016							
54868-4078-00	Q0144			01/01/2002	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	22.5	ML	BO	PO	ML	1	GM	0.04	01/01/2002	02/03/2016							
54868-4078-01	Q0144			01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.04	01/01/2002	99/99/9999							
54868-4078-02	Q0144			01/01/2002	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	30	ML	BO	PO	ML	1	GM	0.04	01/01/2002	02/03/2016							
54868-4082-00	J7644			01/01/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (VIAL) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/01/2002	99/99/9999							
54868-4082-01	KO J7644 KO			01/01/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (VIAL) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/01/2002	99/99/9999							
54868-4082-02	J7644			01/01/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (VIAL) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/01/2002	99/99/9999							
54868-4082-03	KO J7644 KO			01/01/2002	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (VIAL) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	01/01/2002	99/99/9999							
54868-4096-00	J7506			11/27/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (6 DAY DOSEPAK) 5 MG	21	EA	BX	PO	EA	5	MG	1	11/27/2002	12/31/2015							
54868-4100-00	J1055			01/01/2002	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	DEPO-PROVERA CONTRACEPTIVE (SRN,PREFILLED) 150 MG/ML	1	ML	SR	IM	ML	150	MG	1	01/01/2002	12/31/2012							
54868-4100-01	J1055			02/11/2002	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	DEPO-PROVERA CONTRACEPTIVE (SRN,PREFILLED) 150 MG/ML	1	ML	SR	IM	ML	150	MG	1	02/11/2002	12/31/2012							
54868-4103-00	J1580			02/12/2003	02/03/2016	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (FLIPTOP VIAL) 40 MG/ML	2	ML	VL	IJ	ML	80	MG	0.5	02/12/2003	02/03/2016							
54868-4106-00	J3260			01/01/2002	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (M.D.V.) 40 MG/ML	2	ML	VL	IJ	ML	80	MG	0.5	01/01/2002	99/99/9999							
54868-4109-00	Q0178			01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED																		

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-4137-00		J0780		01/01/2002	02/03/2016	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE (CARPUJECT) 5 MG/ML	2 ML	SR	IJ	ML		10 MG		0.5	01/01/2002	02/03/2016						
54868-4138-00		Q0180		02/10/2005	02/03/2016	DOLASETRON MESYLATE, 100 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	ANZEMET 100 MG	5 EA	BO	PO	EA		100 MG		1	02/10/2005	02/03/2016						
54868-4139-00		Q0166		06/03/2005	02/03/2016	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	KYTRIL 1 MG	2 EA	BO	PO	EA		1 MG		1	06/03/2005	02/03/2016						
54868-4139-01		Q0166		06/28/2005	02/03/2016	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	KYTRIL 1 MG	10 EA	BO	PO	EA		1 MG		1	06/28/2005	02/03/2016						
54868-4139-02		Q0166		09/07/2005	02/03/2016	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	KYTRIL 1 MG	6 EA	BO	PO	EA		1 MG		1	09/07/2005	02/03/2016						
54868-4139-03		Q0166		10/14/2005	02/03/2016	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	KYTRIL 1 MG	8 EA	BO	PO	EA		1 MG		1	10/14/2005	02/03/2016						
54868-4139-04		Q0166		09/22/2005	02/03/2016	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	KYTRIL 1 MG	3 EA	BO	PO	EA		1 MG		1	09/22/2005	02/03/2016						
54868-4139-05		Q0166		01/05/2006	02/03/2016	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	KYTRIL 1 MG	20 EA	BO	PO	EA		1 MG		1	01/05/2006	02/03/2016						
54868-4139-06		Q0166		06/07/2006	02/03/2016	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	KYTRIL 1 MG	30 EA	BO	PO	EA		1 MG		1	06/07/2006	02/03/2016						
54868-4142-00	None			06/29/2005	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	5 EA	BO	PO	EA		20 MG		1	06/29/2005	99/99/9999						
54868-4142-01	None			08/03/2006	02/03/2016	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	25 EA	BO	PO	EA		20 MG		1	08/03/2006	02/03/2016						
54868-4142-02	None			01/26/2006	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	10 EA	BO	PO	EA		20 MG		1	01/26/2006	99/99/9999						
54868-4142-03	None			03/16/2006	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	60 EA	BO	PO	EA		20 MG		1	03/16/2006	99/99/9999						
54868-4142-04	None			03/23/2006	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	40 EA	BO	PO	EA		20 MG		1	03/23/2006	99/99/9999						
54868-4142-05	None			03/23/2006	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	30 EA	BO	PO	EA		20 MG		1	03/23/2006	99/99/9999						
54868-4142-06	None			05/16/2006	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR 20 MG	20 EA	BO	PO	EA		20 MG		1	05/16/2006	99/99/9999						
54868-4143-00	None			02/10/2005	99/99/9999	CAPECITABINE, 150 MG, ORAL	XELODA 150 MG	60 EA	BO	PO	EA		150 MG		1	02/10/2005	99/99/9999						
54868-4143-01	None			08/08/2007	02/03/2016	CAPECITABINE, 150 MG, ORAL	XELODA 150 MG	120 EA	BO	PO	EA		150 MG		1	08/08/2007	02/03/2016						
54868-4143-02	None			10/19/2005	02/03/2016	CAPECITABINE, 150 MG, ORAL	XELODA 150 MG	30 EA	BO	PO	EA		150 MG		1	10/19/2005	02/03/2016						
54868-4143-03	None			05/19/2006	99/99/9999	CAPECITABINE, 150 MG, ORAL	XELODA 150 MG	28 EA	BO	PO	EA		150 MG		1	05/19/2006	99/99/9999						
54868-4154-00		J3490		01/01/2002	02/03/2016	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE (S.D.V.) 150 MG/ML	4 ML	VL	IJ	ML		1 EA		1	01/01/2002	02/03/2016						
54868-4167-00		J2765		01/01/2002	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (S.D.V.) 5 MG/ML	2 ML	VL	IV	ML		10 MG		0.5	01/01/2002	99/99/9999						
54868-4169-00		J3490		03/02/2004	02/03/2016	UNCLASSIFIED DRUGS	CLEOCIN PHOSPHATE (S.D.V.) 150 MG/ML	2 ML	VL	IJ	ML		1 EA		1	03/02/2004	02/03/2016						
54868-4189-00		J2270		01/01/2002	02/03/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (AMP,DOSETTE) 10 MG/ML	1 ML	AM	IJ	ML		10 MG		1	01/01/2002	02/03/2016						
54868-4194-00		J3490		01/01/2002	06/30/2013	UNCLASSIFIED DRUGS	BREVITAL SODIUM (VIAL) 5 GM	1 EA	VL	IV	EA		1 EA		1	01/01/2002	06/30/2013						
54868-4287-00		J8999		01/17/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	30 EA	BO	PO	EA		1 EA		1	01/17/2005	99/99/9999						
54868-4287-01		J8999		01/17/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	10 EA	BO	PO	EA		1 EA		1	01/17/2005	99/99/9999						
54868-4287-02		J8999		02/14/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	100 EA	BO	PO	EA		1 EA		1	02/14/2005	99/99/9999						
54868-4287-03		J8999		09/22/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	90 EA	BO	PO	EA		1 EA		1	09/22/2005	99/99/9999						
54868-4287-04		J8999		01/18/2008	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (FILM COATED) 20 MG	60 EA	BO	PO	EA		1 EA		1	01/18/2008	99/99/9999						
54868-4296-00		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR IRRIGATION	500 ML	VL	IR	ML		500 ML		0.002	01/01/2004	99/99/9999						
54868-4311-00		A4217		01/01/2004	99/99/9999	STERILE WATER/SALINE, 500 ML	WATER FOR INJECTION	500 ML	NA	IV	ML		500 ML		0.002	01/01/2004	99/99/9999						
54868-4339-00	None			08/16/2005	02/03/2016	MELPHALAN 2 MG, ORAL	ALKERAN (FILM-COATED) 2 MG	4 EA	BO	PO	EA		2 MG		1	08/16/2005	02/03/2016						
54868-4339-01	None			11/22/2005	02/03/2016	MELPHALAN, 2 MG, ORAL	ALKERAN 2 MG	50 EA	BO	PO	EA		2 MG		1	11/22/2005	02/03/2016						
54868-4339-02	None			02/03/2006	02/03/2016	MELPHALAN, 2 MG, ORAL	ALKERAN 2 MG	24 EA	BO	PO	EA		2 MG		1	02/03/2006	02/03/2016						
54868-4339-03	None			04/03/2006	02/03/2016	MELPHALAN, 2 MG, ORAL	ALKERAN 2 MG	28 EA	BO	PO	EA		2 MG		1	04/03/2006	02/03/2016						
54868-4339-04	None			02/05/2008	02/03/2016	MELPHALAN, 2 MG, ORAL	ALKERAN 2 MG	32 EA	BO	PO	EA		2 MG		1	02/05/2008	02/03/2016						
54868-4381-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG MIX 75/25 (VIAL) 75 U/ML-25 U/ML	10 ML	VL	SC	ML		5 U		20	01/01/2003	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-4409-00		J7614		04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.021%	3	ML	PC	IH	ML	0.5	MG	0.42	04/01/2008	99/99/9999						
54868-4409-00	KO	J7614	KO	04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.021%	3	ML	PC	IH	ML	0.5	MG	0.42	04/01/2008	99/99/9999						
54868-4419-00		J1885		01/01/2002	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 30 MG/ML	2	ML	VL	IM	ML	15	MG	2	01/01/2002	99/99/9999						
54868-4419-01		J1885		10/17/2005	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/ML	2	ML	VL	IM	ML	15	MG	2	10/17/2005	99/99/9999						
54868-4464-00		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (PF) 0.9%	10	ML	VL	IV	ML	10	ML	0.1	01/01/2004	99/99/9999						
54868-4488-00		J2540		01/01/2002	99/99/9999	INJECTION, PENICILLIN G POTASSIUM, UP TO 600,000 UNITS	PENICILLIN G POTASSIUM (VIAL,PHARMACY BOTTLE) 20 Million U	1	EA	VL	IV	EA	600000	U	33.33333	01/01/2002	99/99/9999						
54868-4508-00		J1720		01/01/2002	02/03/2016	INJECTION, HYDROCORTISONE SODIUM SUCCINATE, UP TO 100 MG	SOLU-CORTEF (ACT-Q-VIAL) 1 GM	1	EA	VL	U	EA	100	MG	10	01/01/2002	02/03/2016						
54868-4527-00		J0456		01/01/2002	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	ZITHROMAX (VIAL) 500 MG	1	EA	VL	IV	EA	500	MG	1	01/01/2002	99/99/9999						
54868-4547-00		J0744		01/01/2002	07/29/2013	INJECTION, CIPROFLOXACIN FOR INTRAVENOUS INFUSION, 200 MG	CIPRO IV (VIAL) 10 MG/ML	40	ML	VL	IV	ML	200	MG	0.05	01/01/2002	07/29/2013						
54868-4580-00		J2250		01/01/2002	02/03/2016	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 5 MG/ML DIAZEPAM (22GX1 1/4",CARPUJECT) 5 MG/ML	5	ML	VL	U	ML	1	MG	5	01/01/2002	02/03/2016						
54868-4586-00		J3360		01/23/2002	02/03/2016	INJECTION, DIAZEPAM, UP TO 5 MG	LANTUS (VIAL) 100 U/ML	2	ML	SR	IJ	ML	5	MG	1	01/23/2002	02/03/2016						
54868-4626-00		J1815		01/01/2003	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	FLUTAMIDE 125 MG	180	EA	BO	PO	EA	1	EA	1	06/12/2002	02/03/2016						
54868-4628-00		J8999		06/12/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	PROPOFOL (S.D.V.) 10 MG/ML	20	ML	VL	IV	ML	1	EA	1	10/07/2003	02/03/2016						
54868-4629-00		J3490		10/07/2003	02/03/2016	UNCLASSIFIED DRUGS	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	4	EA	BO	PO	EA	1	GM	0.25	07/26/2002	02/03/2016						
54868-4644-00		Q0144		07/26/2002	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6	EA	BO	PO	EA	1	GM	0.25	02/21/2005	99/99/9999						
54868-4644-01		Q0144		02/21/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6	EA	BO	PO	EA	1	GM	0.25	02/21/2005	99/99/9999						
54868-4644-02		Q0144		06/01/2005	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	30	EA	BO	PO	EA	1	GM	0.25	06/01/2005	02/03/2016						
54868-4651-00		J0690		09/15/2003	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (VIAL,PF) 500 MG	1	EA	VL	U	EA	500	MG	1	09/15/2003	99/99/9999						
54868-4686-00		J8498		01/01/2006	02/03/2016	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHEGAN 25 MG	6	EA	BX	RC	EA	1	EA	1	01/01/2006	02/03/2016						
54868-4686-01		J8498		04/26/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHEGAN 25 MG	12	EA	NA	RC	EA	1	EA	1	04/26/2006	99/99/9999						
54868-4716-00		J9250		12/16/2002	02/03/2016	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (P.F.V.,PF) 25 MG/ML	10	ML	VL	U	ML	5	MG	5	12/16/2002	02/03/2016						
54868-4721-00		Q0164		02/10/2003	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	30	EA	BO	PO	EA	5	MG	1	02/10/2003	99/99/9999						
54868-4721-01		Q0164		04/08/2003	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	15	EA	BO	PO	EA	5	MG	1	04/08/2003	99/99/9999						
54868-4721-02		Q0164		06/09/2005	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	60	EA	BO	PO	EA	5	MG	1	06/09/2005	99/99/9999						
54868-4721-03		Q0164		06/04/2007	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	100	EA	BO	PO	EA	5	MG	1	06/04/2007	99/99/9999						
54868-4748-00		J7510		02/28/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	02/28/2003	99/99/9999						
54868-4749-00		J7510		02/28/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240	ML	BO	PO	ML	5	MG	0.6	02/28/2003	99/99/9999						
54868-4749-01		J7510		05/25/2004	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (CHERRY) 15 MG/5 ML	480	ML	BO	PO	ML	5	MG	0.6	05/25/2004	99/99/9999						
54868-4751-00		J2175		03/11/2003	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE (CARPUJECT) 100 MG/ML	1	ML	AM	IJ	ML	100	MG	1	03/11/2003	99/99/9999						
54868-4751-01		J2175		07/03/2003	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMOROL HYDROCHLORIDE 100 MG/ML	1	ML	AM	IJ	ML	100	MG	1	07/03/2003	99/99/9999						
54868-4752-00		J2270		03/11/2003	02/03/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE 10 MG/ML	1	ML	VL	IJ	ML	10	MG	1	03/11/2003	02/03/2016						
54868-4773-00		J8999		04/10/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDROXYUREA 500 MG	30	EA	BO	PO	EA	1	EA	1	04/10/2003	99/99/9999						
54868-4773-01		J8999		08/06/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDROXYUREA 500 MG	100	EA	BO	PO	EA	1	EA	1	08/06/2003	99/99/9999						
54868-4773-02		J8999		07/07/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDROXYUREA 500 MG	50	EA	BO	PO	EA	1	EA	1	07/07/2005	99/99/9999						
54868-4773-03		J8999		07/14/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	HYDROXYUREA 500 MG	60	EA	BO	PO	EA	1	EA	1	07/14/2005	99/99/9999						
54868-4781-00		J3490		04/24/2003	02/03/2016	UNCLASSIFIED DRUGS	ENGERIX-B PEDIATRIC (PEDIATRIC,PF) 10 MCG/0.5 ML	0.5	ML	VL	IM	ML	1	EA	1	04/24/2003	02/03/2016						
67457-0603-99		J1644		06/14/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X1ML) 10000 U/1 ML	4	ML	VL	IJ	ML	1000	U	10	06/14/2018	99/99/9999						
54868-4782-00		J1438		04/30/2003	06/30/2012	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (PF) 25 MG	4	EA	BX	SC	EA	25	MG	1	04/30/2003	06/30/2012						
54868-4794-02		J8498		08/08/2007	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE 12.5 MG	2	EA	BX	RC	EA	1	EA	1	08/08/2007	99/99/9999						
54868-4804-00		J2270		05/30/2003	06/30/2015	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (22G,SLIM PK,LATEX FREE) 10 MG/ML	1	ML	EA	IJ	ML	10	MG	1	05/30/2003	06/30/2015						
54868-4809-00		J9250		06/03/2003	02/03/2016	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (VIAL, L.P.P.) 25 MG/ML	10	ML	EA	IJ	ML	5	MG	5	06/03/2003	02/03/2016						
67457-0794-10		J3489		06/05/2018	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SINGLE USE,PF) 5 MG/100 ML	100	ML	BG	IV	ML	1	MG	0.05	06/05/2018	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-5350-02		None		11/22/2005	99/99/9999	TEMODAR, 100 MG, ORAL	TEMODAR 100 MG	5 EA	BO PO EA	EA			100 MG			1	11/22/2005	99/99/9999					
54868-5350-03		None		02/08/2006	99/99/9999	TEMODAR, 100 MG, ORAL	TEMODAR 100 MG	10 EA	BO PO EA	EA			100 MG			1	02/08/2006	99/99/9999					
54868-5350-04		None		03/23/2006	99/99/9999	TEMODAR, 100 MG, ORAL	TEMODAR 100 MG	30 EA	BO PO EA	EA			100 MG			1	03/23/2006	99/99/9999					
54868-5354-00		None		04/13/2006	99/99/9999	TEMODAR, 250 MG, ORAL	TEMODAR 250 MG	5 EA	BO PO EA	EA			250 MG			1	04/13/2006	99/99/9999					
54868-5355-00		None		12/20/2005	02/03/2016	ETOPOSIDE, 50 MG, ORAL	ETOPOSIDE 50 MG	20 EA	BX PO EA	EA			50 MG			1	12/20/2005	02/03/2016					
54868-5355-01		None		01/30/2006	99/99/9999	ETOPOSIDE, 50 MG, ORAL	ETOPOSIDE 50 MG	7 EA	NA PO EA	EA			50 MG			1	01/30/2006	99/99/9999					
54868-5355-02		None		01/30/2006	02/03/2016	ETOPOSIDE, 50 MG, ORAL	ETOPOSIDE 50 MG	1 EA	BO PO EA	EA			50 MG			1	01/30/2006	02/03/2016					
54868-5389-00		J8999		09/01/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG/ML	240 ML	BO PO ML	ML			1 EA			1	09/01/2005	99/99/9999					
54868-5389-01		J8999		12/14/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG/ML	480 ML	BO PO ML	ML			1 EA			1	12/14/2005	99/99/9999					
54868-5404-00		Q0144		09/02/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZMAX (CHERRY-BANANA) 2 GM/60 ML	1 EA	BO PO EA	EA			1 GM			2	09/02/2005	99/99/9999					
54868-5406-00		J3110		09/06/2005	02/03/2016	INJECTION, TERIPARATIDE, 10 MCG	FORTEO (RDNA ORIGIN) 250 MCG/ML	3 ML	SR SC ML	ML			10 MCG			25	09/06/2005	02/03/2016					
54868-5428-00		J0881		08/10/2007	06/30/2013	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP 0.2 MG/0.4 ML	0.4 ML	SR IJ ML	ML			1 MCG			500	08/10/2007	06/30/2013					
54868-5429-00		J0881		03/20/2008	06/30/2013	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (1X0.6ML, PREFILLED,PF) 0.3 MG/0.6 ML	0.6 ML	SR IJ ML	ML			1 MCG			500	03/20/2008	06/30/2013					
54868-5440-00		J1650		09/29/2005	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX 40 MG/0.4 ML	0.4 ML	SR SC ML	ML			10 MG			10	09/29/2005	99/99/9999					
54868-5440-01		J1650		11/01/2005	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX 40 MG/0.4 ML	0.4 ML	SR SC ML	ML			10 MG			10	11/01/2005	99/99/9999					
54868-5444-00		J1438		03/18/2008	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (4X0.98ML,PF) 50 MG/ML	0.98 ML	SR SC ML	ML			25 MG			2	03/18/2008	99/99/9999					
54868-5459-00		J7614		04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.042%	3 ML	PC IH ML	ML			0.5 MG			0.84	04/01/2008	99/99/9999					
54868-5459-00	KO	J7614	KO	04/01/2008	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.042%	3 ML	PC IH ML	ML			0.5 MG			0.84	04/01/2008	99/99/9999					
54868-5471-00		Q0144		11/16/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (PAK) 250 MG	6 EA	DP PO EA	EA			1 GM			0.25	11/16/2005	99/99/9999					
54868-5478-00		Q0144		11/23/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6 EA	BO PO EA	EA			1 GM			0.25	11/23/2005	99/99/9999					
54868-5478-01		Q0144		12/13/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	30 EA	BO PO EA	EA			1 GM			0.25	12/13/2005	99/99/9999					
54868-5478-02		Q0144		02/07/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	10 EA	BO PO EA	EA			1 GM			0.25	02/07/2006	99/99/9999					
54868-5487-00		Q0144		12/13/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	6 EA	BO PO EA	EA			1 GM			0.5	12/13/2005	99/99/9999					
54868-5487-01		Q0144		08/10/2007	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	60 EA	BO PO EA	EA			1 GM			0.5	08/10/2007	99/99/9999					
54868-5501-00		J1652		01/11/2006	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA 7.5 MG/0.6 ML	0.6 ML	SR SC ML	ML			0.5 MG			25	01/11/2006	99/99/9999					
54868-5501-01		J1652		01/11/2006	02/03/2016	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA 7.5 MG/0.6 ML	0.6 ML	SR SC ML	ML			0.5 MG			25	01/11/2006	02/03/2016					
54868-5501-02		J1652		11/13/2006	02/03/2016	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA 7.5 MG/0.6 ML	0.6 ML	SR SC ML	ML			0.5 MG			25	11/13/2006	02/03/2016					
54868-5522-00		J7502		02/10/2006	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE 100 MG	30 EA	BO PO EA	EA			100 MG			1	02/10/2006	99/99/9999					
54868-5533-00		J0696		02/17/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 500 MG	1 EA	VL IJ EA	EA			250 MG			2	02/17/2006	99/99/9999					
54868-5551-00		J0150		03/16/2006	12/31/2014	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS, INSTEAD USE A9270)	ADENOSINE 3 MG/ML	2 ML	VL IV ML	ML			6 MG			0.5	03/16/2006	12/31/2014					
54868-5568-00		J9217		04/12/2006	02/03/2016	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	LUPRON DEPOT 30 MG	1 EA	BX IM EA	EA			7.5 MG			4	04/12/2006	02/03/2016					
54868-5569-00		J2355		04/13/2006	02/03/2016	INJECTION, OPRELVEKIN, 5 MG	NEUMEGA 5 MG	1 EA	VL SC EA	EA			5 MG			1	04/13/2006	02/03/2016					
54868-5587-00		J1650		05/17/2006	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX 60 MG/0.6 ML	0.6 ML	SR SC ML	ML			10 MG			10	05/17/2006	99/99/9999					
54868-5587-01		J1650		09/25/2007	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX 60 MG/0.6 ML	6 ML	SR SC ML	ML			10 MG			10	09/25/2007	99/99/9999					
54868-5589-00		J0696		05/12/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 250 MG	1 EA	VL IJ EA	EA			250 MG			1	05/12/2006	99/99/9999					
54868-5596-00		J9015		05/22/2006	02/03/2016	INJECTION, ALDESLEUKIN, PER SINGLE USE VIAL	PROLEKIN 22 Million IU	1 EA	VL IV EA	EA			1 VIAL			1	05/22/2006	02/03/2016					
54868-5612-00		J0770		06/12/2006	02/03/2016	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE 150 MG	1 EA	VL IJ EA	EA			150 MG			1	06/12/2006	02/03/2016					
68001-0355-25		J2469		06/15/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL 0.05 MG/1 ML	5 ML	VL IV ML	ML			25 MCG			2	06/15/2018	99/99/9999					
54868-5621-00		J7626		07/17/2007	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES 0.5 MG/2 ML	60 ML	PC IH ML	ML			0.5 MG			0.5	07/17/2007	99/99/9999					
54868-5621-00	KO	J7626	KO	07/17/2007	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES 0.5 MG/2 ML	60 ML	PC IH ML	ML			0.5 MG			0.5	07/17/2007	99/99/9999					
54868-5634-00		J2941		06/30/2006	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK 0.4 MG	7 EA	CT SC EA	EA			1 MG			0.4	06/30/2006	99/99/9999					
54868-5647-00		Q0144		08/01/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 100 MG/5 ML	15 ML	BO PO ML	ML			1 GM			0.02	08/01/2006	99/99/9999					
54868-5648-00		Q0144		08/01/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	30 ML	BO PO ML	ML			1 GM			0.04	08/01/2006	99/99/9999					
54868-5648-01		Q0144		08/01/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	23 ML	BO PO ML	ML			1 GM			0.04	08/01/2006	99/99/9999					
54868-5648-02		Q0144		08/03/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	15 ML	BO PO ML	ML			1 GM			0.04	08/03/2006	99/99/9999					
54868-5670-00		J7608		08/10/2007	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 20%	30 ML	VL IH ML	ML			1 GM			0.2	08/10/2007	99/99/9999					
54868-5670-00	KO	J7608	KO	08/10/2007	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE 20%	30 ML	VL IH ML	ML			1 GM			0.2	08/10/2007	99/99/9999					
54868-5670-01		J7608		08/10/2007	02/03/2016	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (3X30ML) 20%	30 ML	VL IH ML	ML			1 GM			0.2	08/10/2007	02/03/2016					
54868-5670-01	KO	J7608	KO	08/10/2007	02/03/2016	ACETYLCYSTEINE, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (3X30ML) 20%	30 ML	VL IH ML	ML			1 GM			0.2	08/10/2007	02/03/2016					
54868-5673-01		J0885		03/24/2008	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (M.D.V.1X4ML) 200																

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-5716-00		J2370		12/11/2006	01/15/2013	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	PHENYLEPHRINE HYDROCHLORIDE (SDV,25X1ML) 10 MG/ML	1 ML	VL	IJ	ML		1 ML		1	12/11/2006	01/15/2013						
54868-5717-00		J1250		12/11/2006	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE 12.5 MG/ML	20 ML	VL	IV	ML		250 MG		0.05	12/11/2006	99/99/9999						
54868-5717-01		J1250		01/02/2007	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE (10X40ML) 12.5 MG/ML	40 ML	VL	IV	ML		250 MG		0.05	01/02/2007	99/99/9999						
54868-5717-02		J1250		06/28/2007	99/99/9999	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE 12.5 MG/ML	200 ML	VL	IV	ML		250 MG		0.05	06/28/2007	99/99/9999						
54868-5722-00		J0282		12/11/2006	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE (SDV,10X3ML) 50 MG/ML	3 ML	VL	IV	ML		30 MG		1.666666	12/11/2006	99/99/9999						
54868-5724-00		J3475		12/12/2006	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNES SULF (25X10ML) 500 MG/ML	10 ML	SR	IJ	ML		500 MG		1	12/12/2006	99/99/9999						
68180-0962-56		J7682		06/12/2018	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5 ML	AM	IH	ML		300 MG		0.2	06/12/2018	99/99/9999						
68180-0962-56	KO	J7682	KO	06/12/2018	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5 ML	AM	IH	ML		300 MG		0.2	06/12/2018	99/99/9999						
54868-5741-00		Q0173		01/05/2007	99/99/9999	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE 300 MG	100 EA	BO	PO	EA		250 MG		1.2	01/05/2007	99/99/9999						
69101-0410-01		J7510		06/14/2018	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (AF,DYE-FREE, GRAPE) 20 MG/5 ML	237 ML	BO	PO	ML		5 MG		0.8	06/14/2018	99/99/9999						
54868-5752-00		J0285		01/25/2007	02/03/2016	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B 50 MG	1 EA	VL	IV	EA		50 MG		1	01/25/2007	02/03/2016						
54868-5760-00		J2941		08/17/2007	99/99/9999	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN MINIQUICK 0.8 MG	1 EA	CT	SC	EA		1 MG		0.8	08/17/2007	99/99/9999						
54868-5765-00		J1815		04/04/2007	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	LANTUS 100 U/ML	15 ML	CT	SC	ML		5 U		20	04/04/2007	99/99/9999						
54868-5774-00		J7626		06/01/2007	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES 0.25 MG/2 ML	2 ML	PC	IH	ML		0.25 MG		0.5	06/01/2007	99/99/9999						
54868-5774-00	KO	J7626	KO	06/01/2007	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	PULMICORT RESPULES 0.25 MG/2 ML	2 ML	PC	IH	ML		0.25 MG		0.5	06/01/2007	99/99/9999						
54868-5775-00		J2780		06/06/2007	02/03/2016	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC 25 MG/ML	40 ML	VL	IJ	ML		25 MG		1	06/06/2007	02/03/2016						
54868-5802-00		J0885		08/13/2007	99/99/9999	INJECTION, EPINETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (SDV,1MLX4) 40000 U/ML	1 ML	VL	IJ	ML		1000 U		40	08/13/2007	99/99/9999						
54868-5808-00		J2175		08/21/2007	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	DEMEROL HYDROCHLORIDE (1MLX10) 50 MG/ML	1 ML	SR	IJ	ML		100 MG		0.5	08/21/2007	99/99/9999						
54868-5825-00		J0152		10/18/2007	12/31/2013	INJECTION, ADENOSINE FOR DIAGNOSTIC USE, 30 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOSCAN 3 MG/ML	30 ML	VL	IV	ML		30 MG		0.1	10/18/2007	12/31/2013						
54868-5825-01		J0152		10/18/2007	12/31/2013	INJECTION, ADENOSINE FOR DIAGNOSTIC USE, 30 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS; INSTEAD USE A9270)	ADENOSCAN 3 MG/ML	20 ML	VL	IV	ML		30 MG		0.1	10/18/2007	12/31/2013						
54868-5835-00		J1650		11/29/2007	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (10X1ML) 100 MG/ML	1 ML	SR	IJ	ML		10 MG		10	11/29/2007	99/99/9999						
54868-5836-00		J1817		12/03/2007	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	INSULIN-HUMALOG (1X15ML) 100 U/ML	15 ML	CT	SC	ML		50 U		2	12/03/2007	99/99/9999						
54868-5837-00		J1650		12/04/2007	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX (8X0.8ML) 120 MG/0.8 ML	0.8 ML	SR	IJ	ML		10 MG		15	12/04/2007	99/99/9999						
54868-5867-00		J0881		03/20/2008	06/30/2013	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (1X1ML, PREFILLED,PF) 0.5 MG/ML	1 ML	SR	IJ	ML		1 MCG		500	03/20/2008	06/30/2013						
54868-5888-00		J2405		05/09/2008	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (1X10ML) 2 MG/ML	10 ML	NA	IJ	ML		1 MG		2	05/09/2008	99/99/9999						
54868-5899-00		J1815		05/12/2008	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG PEN (1X15ML) 100 U/ML	15 ML	CT	SC	ML		5 U		20	05/12/2008	99/99/9999						
54868-6624-01		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (DOSE PACK) 4 MG	21 EA	DP	PO	EA		4 MG		1	01/01/2002	99/99/9999						
54888-1082-03		Q0165		10/20/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100 EA	NA	PO	EA		10 MG		1	10/20/2004	12/31/2013						
55045-1124-00		Q0163		05/01/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100 EA	BO	PO	EA		50 MG		1	05/01/2004	06/01/2014						
55045-1124-01		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	3 EA	BO	PO	EA		50 MG		1	12/06/2004	06/01/2014						
55045-1124-02		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	60 EA	NA	PO	EA		50 MG		1	12/06/2004	06/01/2014						
55045-1124-03		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	90 EA	NA	PO	EA		50 MG		1	12/06/2004	06/01/2014						
55045-1124-04		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	120 EA	NA	PO	EA		50 MG		1	12/06/2004	06/01/2014						
55045-1124-05		Q0163		01/01/2003	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15 EA	BO	PO	EA		50 MG		1	01/01/2003	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55045-1124-06		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	6 EA	NA	PO	EA		50 MG		1	12/06/2004	06/01/2014						
55045-1124-07		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20 EA	NA	PO	EA		50 MG		1	12/06/2004	06/01/2014						
55045-1124-08		Q0163		01/01/2003	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30 EA	BO	PO	EA		50 MG		1	01/01/2003	06/01/2014						
55045-1124-09		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	50 EA	NA	PO	EA		50 MG		1	12/06/2004	06/01/2014						
55045-1125-00		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	120 EA	NA	PO	EA		50 MG		0.5	12/06/2004	06/01/2014						
55045-1125-01		Q0163		07/01/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100 EA	NA	PO	EA		50 MG		0.5	07/01/2004	06/01/2014						
55045-1125-02		Q0163		02/01/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	6 EA	NA	PO	EA		50 MG		0.5	02/01/2004	06/01/2014						
55045-1125-03		Q0163		12/06/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	90 EA	NA	PO	EA		50 MG		0.5	12/06/2004	06/01/2014						
55045-1125-04		Q0163		01/01/2003	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	12 EA	BO	PO	EA		50 MG		0.5	01/01/2003	06/01/2014						
55045-1125-05		Q0163		01/02/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15 EA	NA	PO	EA		50 MG		0.5	01/02/2004	06/01/2014						
55045-1125-06		Q0163		01/01/2003	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20 EA	BO	PO	EA		50 MG		0.5	01/01/2003	06/01/2014						
55045-1125-08		Q0163		01/01/2003	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30 EA	BO	PO	EA		50 MG		0.5	01/01/2003	06/01/2014						
55045-1125-09		Q0163		02/01/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60 EA	NA	PO	EA		50 MG		0.5	02/01/2004	06/01/2014						
55045-1126-02		Q0165		04/01/2005	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10 EA	BO	PO	EA		10 MG		1	04/01/2005	12/31/2013						
55045-1126-03		Q0165		07/01/2003	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	5 EA	BO	PO	EA		10 MG		1	07/01/2003	12/31/2013						
55045-1126-04		Q0165		01/01/2003	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	12 EA	BO	PO	EA		10 MG		1	01/01/2003	12/31/2013						
55045-1126-06		Q0165		11/10/2005	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60 EA	BO	PO	EA		10 MG		1	11/10/2005	12/31/2013						
55045-1126-07		Q0165		07/01/2005	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20 EA	BO	PO	EA		10 MG		1	07/01/2005	12/31/2013						
55045-1126-08		Q0165		07/01/2003	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30 EA	BO	PO	EA		10 MG		1	07/01/2003	12/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
55045-1252-02		Q0163		01/01/2003	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (AF) 12.5 MG/5 ML	118	ML	BO	PO	ML	50	MG	0.05	01/01/2003	06/01/2014							
55045-1259-09	J7509			01/01/2003	06/01/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (DOSEPAK) 4 MG	21	EA	DP	PO	EA	4	MG	1	01/01/2003	06/01/2014							
55045-1260-00	J7506			12/06/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE (DOSEPACK) 5 MG	48	EA	DP	PO	EA	5	MG	1	12/06/2004	06/01/2014							
55045-1260-09	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE (DOSEPACK) 5 MG	21	EA	DP	PO	EA	5	MG	1	01/01/2003	06/01/2014							
55045-1308-01	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	100	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014							
55045-1308-02	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	60	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014							
55045-1308-03	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	90	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014							
55045-1308-06	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	6	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014							
55045-1308-07	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	20	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014							
55045-1308-08	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	30	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014							
55045-1308-09	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	36	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014							
55045-1444-01	J7506			12/06/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	35	EA	NA	PO	EA	5	MG	4	12/06/2004	06/01/2014							
55045-1444-02	J7506			05/01/2005	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	42	EA	BO	PO	EA	5	MG	4	05/01/2005	06/01/2014							
55045-1444-03	J7506			01/01/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	18	EA	BO	PO	EA	5	MG	4	01/01/2004	06/01/2014							
55045-1444-04	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	12	EA	BO	PO	EA	5	MG	4	01/01/2003	06/01/2014							
55045-1444-07	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	21	EA	BO	PO	EA	5	MG	4	01/01/2003	06/01/2014							
55045-1444-08	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	01/01/2003	06/01/2014							
55045-1480-01	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2003	06/01/2014							
55045-1480-02	J7506			12/06/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	60	EA	NA	PO	EA	5	MG	1	12/06/2004	06/01/2014							
55045-1480-05	J7506			12/06/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	15	EA	NA	PO	EA	5	MG	1	12/06/2004	06/01/2014							
55045-1480-06	J7506			12/06/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	20	EA	NA	PO	EA	5	MG	1	12/06/2004	06/01/2014							
55045-1480-07	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	21	EA	BO	PO	EA	5	MG	1	01/01/2003	06/01/2014							
55045-1480-08	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	30	EA	BO	PO	EA	5	MG	1	01/01/2003	06/01/2014							
55045-1480-09	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	40	EA	BO	PO	EA	5	MG	1	01/01/2003	06/01/2014							
55045-1533-01	J7506			05/01/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100	EA	NA	PO	EA	5	MG	2	05/01/2004	06/01/2014							
55045-1533-03	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	20	EA	BO	PO	EA	5	MG	2	01/01/2003	06/01/2014							
55045-1533-06	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	42	EA	BO	PO	EA	5	MG	2	01/01/2003	06/01/2014							
55045-1533-07	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21	EA	BO	PO	EA	5	MG	2	01/01/2003	06/01/2014							
55045-1533-08	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2003	06/01/2014							
55045-1533-09	J7506			01/01/2003	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	40	EA	BO	PO	EA	5	MG	2	01/01/2003	06/01/2014							
55045-1596-00		Q0170		12/06/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	25	MG	1	12/06/2004	12/31/2013							
55045-1596-01		Q0170		12/06/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	120	EA	BO	PO	EA	25	MG	1	12/06/2004	12/31/2013							
55045-1596-02		Q0170		08/09/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	NA	PO	EA	25	MG	1	08/09/2004	12/31/2013							
55045-1596-03		Q0170		01/01/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	25	MG	1	01/01/2003	12/31/2013							
55045-1596-04		Q0170		02/09/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	NA	PO	EA	25	MG	1	02/09/2003	12/31/2013							
55045-1596-05		Q0170		01/01/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	25	MG	1	01/01/2003	12/31/2013							
55045-1596-06		Q0170		01/01/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	25	MG	1	01/01/2003	12/31/2013							
55045-1596-08		Q0170		01/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	25	MG	1	05/23/2005	12/31/2013	01/01/2004	05/22/2005	1				
55045-1596-09		Q0170		12/06/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	90	EA	BO	PO	EA	25	MG	1	12/06/2004	12/31/2013							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55045-1628-03		Q0173		01/01/2003	06/01/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	10	EA	BO	PO	EA	250	MG	1	01/01/2003	06/01/2014						
55045-1643-09		Q0170		01/01/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (FRUIT,TROPICAL) 6.25 MG/5 ML	118	ML	BO	PO	ML	25	MG	0.05	01/01/2003	12/31/2013						
55045-1661-00		Q0178		12/06/2004	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	NA	PO	EA	50	MG	1	12/06/2004	12/31/2013						
55045-1661-01		Q0178		12/06/2004	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	120	EA	NA	PO	EA	50	MG	1	12/06/2004	12/31/2013						
55045-1661-02		Q0178		12/06/2004	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20	EA	NA	PO	EA	50	MG	1	12/06/2004	12/31/2013						
55045-1661-03		Q0178		09/01/2004	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	40	EA	NA	PO	EA	50	MG	1	09/01/2004	12/31/2013						
55045-1661-06		Q0178		09/01/2004	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	60	EA	NA	PO	EA	50	MG	1	09/01/2004	12/31/2013						
55045-1661-08		Q0178		06/01/2003	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30	EA	BO	PO	EA	50	MG	1	06/01/2003	12/31/2013						
55045-1661-09		Q0178		12/06/2004	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	90	EA	NA	PO	EA	50	MG	1	12/06/2004	12/31/2013						
55045-1696-02		Q0164		12/06/2004	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (FILM-COATED) 5 MG	10	EA	NA	PO	EA	5	MG	1	12/06/2004	06/01/2014						
55045-1749-02		J8498		01/01/2006	06/01/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 25 MG	4	EA	BO	RC	EA	1	EA	1	01/01/2006	06/01/2014						
55045-1811-03		J7509		12/06/2004	06/01/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	40	EA	NA	PO	EA	4	MG	1	12/06/2004	06/01/2014						
55045-1811-08		J7509		12/06/2004	06/01/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	30	EA	NA	PO	EA	4	MG	1	12/06/2004	06/01/2014						
55045-1970-05		J8540		01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	8	EA	BO	PO	EA	0.25	MG	16	01/01/2006	06/01/2014						
55045-2043-07		J7613		04/01/2008	06/01/2014	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (3MLX25) 0.083%	3	ML	NA	IH	ML	1	MG	0.83	04/01/2008	06/01/2014						
55045-2043-07	KO	J7613	KO	04/01/2008	06/01/2014	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (3MLX25) 0.083%	3	ML	NA	IH	ML	1	MG	0.83	04/01/2008	06/01/2014						
55045-2133-03		J3360		03/24/2003	06/01/2014	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM 5 MG/ML	10	ML	VL	IJ	ML	5	MG	1	03/24/2003	06/01/2014						
55045-2195-02		Q0177		12/06/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	120	EA	NA	PO	EA	25	MG	1	12/06/2004	06/01/2014						
55045-2195-04		Q0177		07/01/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	9	EA	BO	PO	EA	25	MG	1	07/01/2004	06/01/2014						
55045-2195-05		Q0177		03/24/2003	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	15	EA	BO	PO	EA	25	MG	1	03/24/2003	06/01/2014						
55045-2195-06		Q0177		12/06/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	60	EA	NA	PO	EA	25	MG	1	12/06/2004	06/01/2014						
55045-2195-07		Q0177		03/01/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20	EA	NA	PO	EA	25	MG	1	03/01/2004	06/01/2014						
55045-2195-08		Q0177		02/01/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	NA	PO	EA	25	MG	1	02/01/2004	06/01/2014						
55045-2195-09		Q0177		12/06/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	90	EA	NA	PO	EA	25	MG	1	12/06/2004	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55045-2372-05		Q0144		01/19/2005	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 100 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.02	01/19/2005	06/01/2014						
55045-2373-05		Q0144		01/19/2005	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.04	01/19/2005	06/01/2014						
55045-2373-06		Q0144		01/01/2003	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	22.5 ML	BO	PO	ML		1 GM		0.04	01/01/2003	06/01/2014						
55045-2373-08		Q0144		01/19/2005	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 200 MG/5 ML	30 ML	BO	PO	ML		1 GM		0.04	01/19/2005	06/01/2014						
55045-2400-02		J8498		01/01/2006	06/01/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12 EA	BX	RC	EA		1 EA		1	01/01/2006	06/01/2014						
55045-2470-02	J7611			04/01/2008	06/01/2014	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE 0.5%	20 ML	NA	IH	ML		1 MG		5	04/01/2008	06/01/2014						
55045-2492-06	Q0144			07/03/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX Z-PAK 250 MG	6 EA	BX	PO	EA		1 GM		0.25	07/03/2006	06/01/2014						
55045-2533-00	J0595			01/01/2004	06/01/2014	INJECTION, BUTORPHANOL TARTRATE, 1 MG	STADOL 2 MG/ML	10 ML	VL	IJ	ML		1 MG		2	01/01/2004	06/01/2014						
55045-2565-00	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	100 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2565-02	J8499			12/06/2004	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25 EA	BO	PO	EA		1 EA		1	12/06/2004	06/01/2014						
55045-2565-04	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	50 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2565-05	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	15 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2565-08	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2571-00	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2571-02	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2571-04	J8499			01/01/2003	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15 EA	BO	PO	EA		1 EA		1	01/01/2003	06/01/2014						
55045-2571-05	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	50 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2571-06	J8499			03/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	60 EA	NA	PO	EA		1 EA		1	03/01/2005	06/01/2014						
55045-2571-08	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2648-00	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2648-02	J8499			07/01/2003	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	15 EA	BO	PO	EA		1 EA		1	07/01/2003	06/01/2014						
55045-2648-03	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	25 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2648-05	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	50 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2648-06	J8499			01/01/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	60 EA	NA	PO	EA		1 EA		1	01/01/2005	06/01/2014						
55045-2665-02	J8540			01/01/2006	06/01/2014	Dexamethasone, oral, 0.25 MG	DEXAMETHASONE 0.5 MG	12 EA	BO	PO	EA		0.25 MG		2	01/01/2006	06/01/2014						
55045-2781-06	Q0163			07/01/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	24 EA	NA	PO	EA		50 MG		0.5	07/01/2004	06/01/2014						
55045-2857-01	J2250			12/01/2005	06/01/2014	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HYDROCHLORIDE 5 MG/ML	1 ML	VL	IJ	ML		1 MG		5	12/01/2005	06/01/2014						
55045-2885-00	J7510			01/02/2006	06/01/2014	PREDNISOLONE ORAL, PER 5 MG	ORAPRED (10X20ML) 15 MG/5 ML	20 ML	BO	PO	ML		5 MG		0.6	01/02/2006	06/01/2014						
55045-2885-08	J7510			07/05/2006	06/01/2014	PREDNISOLONE ORAL, PER 5 MG	ORAPRED 15 MG/5 ML	237 ML	BO	PO	ML		5 MG		0.6	07/05/2006	06/01/2014						
55045-2887-02	J2250			08/27/2003	06/01/2014	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (10X2ML) 1 MG/ML	2 ML	EA	IJ	ML		1 MG		1	08/27/2003	06/01/2014						
55045-2963-01	J7506			12/06/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE (DOSEPACK) 10 MG	21 EA	DP	PO	EA		5 MG		2	12/06/2004	06/01/2014						
55045-2963-02	J7506			12/06/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE (DOSEPACK) 10 MG	48 EA	DP	PO	EA		5 MG		2	12/06/2004	06/01/2014						
55045-2968-01	J0595			01/01/2005	06/01/2014	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (10X1ML) 2 MG/ML	1 ML	NA	IJ	ML		1 MG		2	01/01/2005	06/01/2014						
55045-2968-02	J0595			04/11/2006	06/01/2014	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE 2 MG/ML	1 ML	NA	IJ	ML		1 MG		2	04/11/2006	06/01/2014						
55045-3011-02	J8498			01/01/2006	06/01/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 25 MG	4 EA	BX	RC	EA		1 EA		1	01/01/2006	06/01/2014						
55045-3011-03	J8498			01/01/2006	06/01/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 25 MG	12 EA	NA	RC	EA		1 EA		1	01/01/2006	06/01/2014						
55045-3029-02	J1080			01/01/2003	06/01/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DEPO-TESTOSTERONE 200 MG/ML	10 ML	VL	IM	ML		200 MG		1	01/01/2003	06/01/2014						
55045-3203-03	Q0173			05/01/2005	06/01/2014	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	TRIMETHOBENZAMIDE 300 MG	10 EA	NA	PO	EA		250 MG		1.2	05/01/2005	06/01/2014						
55045-3212-03	J1100			07/01/2006	06/01/2014	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE 4 MG/ML	30 ML	NA	IJ	ML		1 MG		4	07/01/2006	06/01/2014						
55045-3231-01	J2001			07/01/2006	06/01/2014	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HYDROCHLORIDE 1%	50 ML	NA	IJ	ML		10 MG		1	07/01/2006	06/01/2014						
55045-3232-01	J0690			09/01/2004	06/01/2014	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM 1 GM	1 EA	NA	IJ	EA		500 MG		2	09/01/2004	06/01/2014						
55045-3242-02	J1030			07/01/2006	06/01/2014	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO MEDROL 40 MG/ML	10 ML	NA	IJ	ML		40 MG		1	07/01/2006	06/01/2014						
55045-3242-05	J1030			07/01/2006	06/01/2014	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	DEPO MEDROL 40 MG/ML	5 ML	NA	IJ	ML		40 MG		1	07/01/2006	06/01/2014						
55045-3243-01	J1040			07/20/2006	06/01/2014	INJECTION, METHYLPREDNISOLONE ACETATE, 80 MG	DEPO MEDROL 80 MG/ML	1 ML	VL	IJ	ML		80 MG		1	07/20/2006	06/01/2014						
55045-3248-01	J3301			07/21/2006	06/01/2014	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG 40 40 MG/ML	1 ML	VL	IJ	ML		10 MG		4	07/21/2006	06/01/2014						
55045-3249-05	J2001			07/01/2006	06/01/2014	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HYDROCHLORIDE 2%	50 ML	NA	IJ	ML		10 MG		2	07/01/2006	06/01/2014						
55045-3251-05	J3490			07/01/2006	06/01/2014	UNCLASSIFIED DRUGS	MARCAINE HYDROCHLORIDE 0.5%	50 ML	NA	IJ	ML		1 EA		1	07/01/2006	06/01/2014						
55045-3252-02	J3490			07/01/2006	06/01/2014	UNCLASSIFIED DRUGS	MARCAINE HYDROCHLORIDE 0.25%	50 ML	NA	IJ	ML		1 EA		1	07/01/2006	06/01/2014						
55045-3281-03	J7506			12/20/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	15 EA	NA	PO	EA		5 MG		2	12/20/2004	06/01/2014						
55045-3281-04	J7506			02/11/2005	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	18 EA	NA	PO	EA		5 MG		2	02/11/2005	06/01/2014						
55045-3298-01	J1200			01/01/200																			

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55045-3511-01		J0696		07/11/2006	06/01/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 1 GM	1 EA	VL	U	EA		250 MG		4	07/11/2006	06/01/2014						
55045-3511-02		J0696		07/14/2006	06/01/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 1 GM	1 EA	VL	U	EA		250 MG		4	07/14/2006	06/01/2014						
55045-3512-01		J3030		07/11/2006	06/01/2014	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX (5X0.5ML) 6 MG/0.5 ML	0.5 ML	VL	SC	ML		6 MG		2	07/11/2006	06/01/2014						
55045-3513-01		J7509		06/23/2006	06/01/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 8 MG	25 EA	BO	PO	EA		4 MG		2	06/23/2006	06/01/2014						
55045-3514-01		J2550		07/12/2006	06/01/2014	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HYDROCHLORIDE (25X1ML) 25 MG/ML	1 ML	AM	IJ	ML		50 MG		0.5	07/12/2006	06/01/2014						
55045-3515-01		J2310		07/12/2006	06/01/2014	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HYDROCHLORIDE 0.4 MG/ML	1 ML	AM	IJ	ML		1 MG		0.4	07/12/2006	06/01/2014						
55045-3516-01		J0696		07/12/2006	06/01/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 250 MG	1 EA	VL	U	EA		250 MG		1	07/12/2006	06/01/2014						
55045-3685-01		J1815		11/15/2006	06/01/2014	INJECTION, INSULIN, PER 5 UNITS	LANTUS 100 U/ML	10 ML	VL	SC	ML		5 U		20	11/15/2006	06/01/2014						
55045-3693-01		Q0144		12/06/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	3 EA	NA	PO	EA		1 GM		0.5	12/06/2006	06/01/2014						
55045-3698-03		Q0144		12/26/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	30 ML	BO	PO	ML		1 GM		0.04	12/26/2006	06/01/2014						
55045-3710-01		A4216		01/01/2007	06/01/2014	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (10MLX25) 0.9%	10 ML	NA	IJ	ML		10 ML		0.1	01/01/2007	06/01/2014						
55045-3725-01		Q0144		12/26/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 100 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.02	12/26/2006	06/01/2014						
72205-0007-92		None		10/01/2018	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (FILM COATED) 500 MG	120 EA	BO	PO	EA		500 MG		1	10/01/2018	99/99/9999						
55045-3726-02		Q0144		12/26/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	22.5 ML	BO	PO	ML		1 GM		0.04	12/26/2006	06/01/2014						
55045-3727-01		Q0144		12/26/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 200 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.04	12/26/2006	06/01/2014						
55045-3773-05		J3490		04/06/2007	06/01/2014	UNCLASSIFIED DRUGS	BACITRACIN 50000 U	1 EA	NA	IM	EA		1 EA		1	04/06/2007	06/01/2014						
55289-0006-10		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 200 MG	10 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0006-25		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 200 MG	25 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0006-35		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 200 MG	35 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0006-50		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 200 MG	50 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0100-01		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100 EA	BO	PO	EA		50 MG		1	01/01/2002	02/03/2016						
55289-0100-10		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	10 EA	BO	PO	EA		50 MG		1	01/01/2002	02/03/2016						
55289-0100-15		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15 EA	BO	PO	EA		50 MG		1	01/01/2002	02/03/2016						
55289-0100-20		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20 EA	BO	PO	EA		50 MG		1	01/01/2002	02/03/2016						
55289-0100-30		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30 EA	BO	PO	EA		50 MG		1	01/01/2002	02/03/2016						
55289-0100-40		Q0163		09/09/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	40 EA	BO	PO	EA		50 MG		1	09/09/2002	02/03/2016						
55289-0119-02		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	2 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
55289-0119-06		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	6 EA	BX	RC	EA		1 EA		1	01/01/2006	99/99/9999						
55289-0224-04		Q0165		05/21/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	4 EA	BO	PO	EA		10 MG		1	05/21/2002	12/31/2013						
55289-0224-06		Q0165		03/07/2008	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10 EA	BO	PO	EA		10 MG		1	03/07/2008	12/31/2013						
55289-0224-12		Q0165		04/02/2008	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	12 EA	BO	PO	EA		10 MG		1	04/02/2008	12/31/2013						
55289-0226-10		Q0177		01/01/2002	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	10 EA	BO	PO	EA		25 MG		1	01/01/2002	99/99/9999						
55289-0226-15		Q0177		03/06/2008	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	15 EA	BO	PO	EA		25 MG		1	03/06/2008	99/99/9999						
55289-0273-10		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	10 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0273-25		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0273-30		J8499		08/01/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30 EA	BO	PO	EA		1 EA		1	08/01/2006	99/99/9999						
55289-0273-35		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	35 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55289-0273-50		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	50 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0274-02		Q0144		10/16/2007	03/08/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	2 EA	BO	PO	EA		1 GM		0.5	10/16/2007	03/08/2017						
55289-0274-03		Q0144		04/02/2008	03/08/2017	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	3 EA	BO	PO	EA		1 GM		0.5	04/02/2008	03/08/2017						
55289-0310-04		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	4 EA	BO	PO	EA		1 GM		0.25	01/01/2002	99/99/9999						
55289-0310-06		Q0144		01/15/2004	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6 EA	BO	PO	EA		1 GM		0.25	01/15/2004	99/99/9999						
55289-0310-14		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	14 EA	BO	PO	EA		1 GM		0.25	01/01/2002	99/99/9999						
55289-0330-05		J7506		04/25/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (USP) 50 MG	5 EA	BO	PO	EA		5 MG		10	04/25/2008	12/31/2015						
55289-0330-10		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 50 MG	10 EA	BO	PO	EA		5 MG		10	01/01/2002	12/31/2015						
55289-0352-05		J7506		05/01/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (USP) 20 MG	5 EA	BO	PO	EA		5 MG		4	05/01/2008	12/31/2015						
55289-0352-07		J7506		03/01/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	7 EA	BO	PO	EA		5 MG		4	03/01/2004	12/31/2015						
55289-0352-09		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	9 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
55289-0352-10		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	10 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
55289-0352-12		J7506		05/01/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (USP) 20 MG	12 EA	BO	PO	EA		5 MG		4	05/01/2008	12/31/2015						
55289-0352-14		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	14 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
55289-0352-15		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	15 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
55289-0352-20		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	20 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
55289-0352-21		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	21 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
55289-0352-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	30 EA	BO	PO	EA		5 MG		4	01/01/2002	12/31/2015						
55289-0354-10		Q0178		10/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	10 EA	BO	PO	EA		50 MG		1	10/01/2002	12/31/2013						
55289-0373-01		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	100 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0373-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	30 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0373-36		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	36 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0373-42		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	42 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0373-46		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	46 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0373-55		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	50 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0373-60		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	60 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0373-72		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	72 EA	BO	PO	EA		5 MG		1	01/01/2002	12/31/2015						
55289-0438-20		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
55289-0438-21		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	21 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
55289-0438-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
55289-0438-36		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	36 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
55289-0438-38		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	38 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
55289-0438-40		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	40 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
55289-0438-42		J7506		03/18/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (USP) 10 MG	42 EA	BO	PO	EA		5 MG		2	03/18/2008	12/31/2015						
55289-0438-50		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	50 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
55289-0438-60		J7506		03/05/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	60 EA	BO	PO	EA		5 MG		2	03/05/2002	12/31/2015						
55289-0462-05		J8499		01/15/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	5 EA	BO	PO	EA		1 EA		1	01/15/2004	99/99/9999						
55289-0462-12		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	12 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0462-15		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0462-21		J8499		08/17/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	21 EA	BO	PO	EA		1 EA		1	08/17/2006	99/99/9999						
55289-0462-25		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0462-30		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30 EA	BO	PO	EA		1 EA		1	01/01/2002	99/99/9999						
55289-0462-35		J8499		04/21/2008	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (USP) 400 MG	35 EA	BO	PO	EA		1 EA		1	04/21/2008	99/99/9999						
55289-0462-60		J8499		03/01/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (USP) 400 MG	60 EA	BO	PO	EA		1 EA		1	03/01/2006	99/99/9999						
55289-0464-15		Q0170		12/01/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15 EA	BO	PO	EA		25 MG		1	12/01/2003	12/31/2013						
55289-0464-79		Q0170		02/01/2005	12/31/2015	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	1 EA	BO	PO	EA		25 MG		1	05/24/2005	12/31/2013	02/01/2005	05/23/2005				
55289-0479-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
55289-0479-10		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	10 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
55289-0479-12		Q0163		07/01/2006	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	12 EA	BO	PO	EA		50 MG		0.5	07/01/2006	99/99/9999						
55289-0479-15		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55289-0479-20		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
55289-0479-24		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	24	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
55289-0479-30		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
55289-0531-04		Q0170		02/26/2008	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 50 MG	4	EA	BO	PO	EA	25	MG	2	02/26/2008	12/31/2013						
55289-0564-15		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	15	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
55289-0564-20		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	20	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
55289-0564-48		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 800 MG	48	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
55289-0568-10		Q0164		07/01/2005	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	10	EA	BO	PO	EA	5	MG	1	07/01/2005	99/99/9999						
55289-0568-12		Q0164		10/01/2002	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	12	EA	BO	PO	EA	5	MG	1	10/01/2002	99/99/9999						
55289-0568-30		Q0164		11/15/2007	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	30	EA	BO	PO	EA	5	MG	1	11/15/2007	99/99/9999						
55289-0582-04		J8540		10/01/2007	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	4	EA	BO	PO	EA	0.25	MG	16	10/01/2007	99/99/9999						
55289-0582-10		J8540		04/10/2008	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	10	EA	BO	PO	EA	0.25	MG	16	04/10/2008	99/99/9999						
55289-0629-10		J8499		08/26/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	10	EA	BO	PO	EA	1	EA	1	08/26/2002	99/99/9999						
55289-0629-30		J8499		06/05/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	06/05/2007	99/99/9999						
55289-0629-50		J8499		04/23/2008	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (USP) 800 MG	50	EA	BO	PO	EA	1	EA	1	04/23/2008	99/99/9999						
55289-0649-30		J7509		10/15/2003	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	30	EA	BO	PO	EA	4	MG	1	10/15/2003	99/99/9999						
55289-0649-98		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	120	EA	BO	PO	EA	4	MG	1	01/01/2002	99/99/9999						
55289-0691-12		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 400 MG	12	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
55289-0691-15		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 400 MG	15	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
55289-0691-25		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ZOVIRAX 400 MG	25	EA	BO	PO	EA	1	EA	1	01/01/2002	99/99/9999						
55289-0924-30		None		11/01/2005	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	30	EA	BO	PO	EA	2.5	MG	1	11/01/2005	99/99/9999						
55289-0928-02		J8498		03/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE (USP) 25 MG	2	EA	BX	RC	EA	1	EA	1	03/01/2006	99/99/9999						
55289-0928-04		J8498		05/09/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE 25 MG	4	EA	BX	RC	EA	1	EA	1	05/09/2006	99/99/9999						
55289-0928-06		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE 25 MG	6	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999						
55289-0928-79		J8498		01/01/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE 25 MG	1	EA	BX	RC	EA	1	EA	1	01/01/2006	99/99/9999						
55289-0940-02		J8498		03/01/2006	02/05/2018	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE (USP) 12.5 MG	2	EA	BX	RC	EA	1	EA	1	03/01/2006	02/05/2018						
55289-0940-06		J8498		05/09/2006	02/05/2018	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HYDROCHLORIDE 12.5 MG	6	EA	BX	RC	EA	1	EA	1	05/09/2006	02/05/2018						
55289-0948-02		Q0169		05/09/2006	99/99/9999	PROMETHAZINE 12.5 MG	PROMETHAZINE 12.5 MG	2	EA	BO	PO	EA	12.5	MG	1	05/09/2006	99/99/9999						
55289-0953-06		Q0173		05/09/2006	99/99/9999	TRIMETHOBENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOBENZAMIDE 300 MG	6	EA	BO	PO	EA	250	MG	1.2	05/09/2006	99/99/9999						
55289-0964-04		Q0144		11/01/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	4	EA	BO	PO	EA	1	GM	0.25	11/01/2005	99/99/9999						
55289-0964-14		Q0144		02/01/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	14	EA	BO	PO	EA	1	GM	0.25	02/01/2006	99/99/9999						
55390-0003-10		J1800		01/01/2002	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (S.D.V.) 1 MG/ML	1	ML	VL	IV	ML	1	MG	1	01/01/2002	99/99/9999						
55390-0004-01		J1610		01/01/2002	04/08/2015	INJECTION, GLUCAGON HYDROCHLORIDE, PER 1 MG	GLUCAGEN DIAGNOSTIC KIT (VIAL W/STERILE WATER) 1 MG	1	EA	VL	IJ	EA	1	MG	1	01/01/2002	04/08/2015						
55390-0004-10		J1610		01/01/2002	04/08/2015	INJECTION, GLUCAGON HYDROCHLORIDE, PER 1 MG	GLUCAGEN (VIAL) 1 MG	1	EA	VL	IJ	EA	1	MG	1	01/01/2002	04/08/2015						
55390-0009-01		J0640		01/01/2002	09/05/2014	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (S.D.V.,PF) 10 MG/ML	50	ML	VL	IJ	ML	50	MG	0.2	01/01/2002	09/05/2014						
55390-0012-01		J1450		07/29/2004	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE 200 MG/100 ML DIHYDROERGOTAMINE MESYLATE (VIAL) 1 MG/ML	100	ML	VL	IV	ML	200	MG	0.01	07/29/2004	99/99/9999						
55390-0013-10		J1110		09/03/2003	11/09/2016	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG	DEXRAZOXANE 250 MG	1	ML	VL	IJ	ML	1	MG	1	09/03/2003	11/09/2016						
55390-0014-02		J1190		04/08/2005	09/05/2014	INJECTION, DEXRAZOXANE HYDROCHLORIDE, PER 250 MG	DEXRAZOXANE 250 MG	1	EA	VL	IV	EA	250	MG	1	04/08/2005	09/05/2014						
55390-0019-10		J2260		05/31/2002	05/31/2012	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (S.D.V.) 1 MG/ML	10	ML	VL	IV	ML	5	MG	0.2	05/31/2002	05/31/2012						
55390-0020-10		J2260		05/31/2002	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (S.D.V.) 1 MG/ML	20	ML	VL	IV	ML	5	MG	0.2	05/31/2002	99/99/9999						
55390-0021-01		J2260		05/31/2002	05/31/2015	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (S.D.V.) 1 MG/ML	50	ML	VL	IV	ML	5	MG	0.2	05/31/2002	05/31/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55390-0026-01		J3490		01/01/2002	09/30/2012	UNCLASSIFIED DRUGS	FAMOTIDINE (BULK VIAL) 10 MG/ML	50 ML	VL	IV	ML		1 EA		1	01/01/2002	09/30/2012						
55390-0027-01		J3490		01/01/2002	08/31/2012	UNCLASSIFIED DRUGS	FAMOTIDINE (M.D.V.,PF) 10 MG/ML	20 ML	VL	IV	ML		1 EA		1	01/01/2002	08/31/2012						
55390-0028-10		J3490		01/01/2002	08/31/2012	UNCLASSIFIED DRUGS	FAMOTIDINE (M.D.V.,PF) 10 MG/ML	4 ML	VL	IV	ML		1 EA		1	01/01/2002	08/31/2012						
55390-0029-10		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	FAMOTIDINE (S.D.V.,PF) 10 MG/ML	2 ML	VL	IV	ML		1 EA		1	01/01/2002	99/99/9999						
55390-0030-10		J9340		01/01/2002	09/05/2014	INJECTION, THIOTEPA, 15 MG	THIOTEPA (S.D.V.) 15 MG	1 EA	VL	IJ	EA		15 MG		1	01/01/2002	09/05/2014						
55390-0031-10		J9250		01/01/2002	09/05/2014	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	2 ML	VL	IJ	ML		5 MG		5	01/01/2002	09/05/2014						
55390-0032-10		J9250		01/01/2002	09/05/2014	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	4 ML	VL	IJ	ML		5 MG		5	01/01/2002	09/05/2014						
55390-0033-10		J9250		01/01/2002	09/05/2014	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	8 ML	VL	IJ	ML		5 MG		5	01/01/2002	09/05/2014						
55390-0034-10		J9250		01/01/2002	09/05/2014	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	10 ML	VL	IJ	ML		5 MG		5	01/01/2002	09/05/2014						
55390-0045-01		J9209		02/24/2004	09/05/2014	INJECTION, MESNA, 200 MG	MESNA (M.D.V.) 100 MG/ML	10 ML	VL	IV	ML		200 MG		0.5	02/24/2004	09/05/2014						
55390-0046-01		J1450		07/29/2004	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE 400 MG/200 ML	200 ML	VL	IV	ML		200 MG		0.01	07/29/2004	99/99/9999						
55390-0051-10		J0640		01/01/2002	09/05/2014	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (VIAL) 50 MG	1 EA	VL	IJ	EA		50 MG		1	01/01/2002	09/05/2014						
55390-0052-10		J0640		01/01/2002	09/05/2014	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (VIAL) 100 MG	1 EA	VL	IJ	EA		50 MG		2	01/01/2002	09/05/2014						
55390-0053-01		J0640		01/01/2002	09/05/2014	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (VIAL) 200 MG	1 EA	VL	IJ	EA		50 MG		4	01/01/2002	09/05/2014						
55390-0054-01		J0640		01/01/2002	09/05/2014	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (S.D.V.,PF) 350 MG	1 EA	VL	IJ	EA		50 MG		7	01/01/2002	09/05/2014						
55390-0059-10		J2360		04/28/2003	09/05/2014	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORPHENADRINE CITRATE (S.D.V.) 30 MG/ML	2 ML	VL	IJ	ML		60 MG		0.5	04/28/2003	09/05/2014						
55390-0060-02		J1190		04/08/2005	09/05/2014	INJECTION, DEXRAZOXANE HYDROCHLORIDE, PER 250 MG	DEXRAZOXANE 500 MG	1 EA	VL	IV	EA		250 MG		2	04/08/2005	09/05/2014						
55390-0067-10		J0150		06/16/2004	12/31/2014	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS, INSTEAD USE A9270)	ADENOSINE (S.D.V.) 3 MG/ML	2 ML	VL	IV	ML		6 MG		0.5	06/16/2004	12/31/2014						
55390-0069-01		J9390		02/03/2004	12/31/2012	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (S.D.V.,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	02/03/2004	12/31/2012						
55390-0070-01		J9390		02/03/2004	12/31/2012	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (S.D.V.,PF) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	02/03/2004	12/31/2012						
50268-0154-11	None			03/12/2018	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE AVPAK (INNER PACK FILM COATED) 500 MG	1 EA	ST	PO	EA		500 MG		1	03/12/2018	99/99/9999						
55390-0074-10		J2260		05/31/2002	04/18/2013	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE NOVAPLUS (S.D.V.) 1 MG/ML	10 ML	VL	IV	ML		5 MG		0.2	05/31/2002	04/18/2013						
55390-0075-10		J2260		05/31/2002	04/18/2013	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE NOVAPLUS (S.D.V.) 1 MG/ML	20 ML	VL	IV	ML		5 MG		0.2	05/31/2002	04/18/2013						
55390-0076-01		J2260		05/31/2002	04/18/2013	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE NOVAPLUS (S.D.V.) 1 MG/ML	50 ML	VL	IV	ML		5 MG		0.2	05/31/2002	04/18/2013						
55390-0077-01		J0780		07/22/2004	06/14/2016	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE (U.S.P., M.D.V.) 5 MG/ML	10 ML	VL	IJ	ML		10 MG		0.5	07/22/2004	06/14/2016						
55390-0077-10		J0780		07/22/2004	99/99/9999	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE (U.S.P., M.D.V.) 5 MG/ML	2 ML	VL	IJ	ML		10 MG		0.5	07/22/2004	99/99/9999						
50268-0154-13	None			03/12/2018	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE AVPAK (FILM COATED) 500 MG	30 EA	ST	PO	EA		500 MG		1	03/12/2018	99/99/9999						
55390-0091-10		J9360		01/01/2002	09/05/2014	INJECTION, VINBLASTINE SULFATE, 1 MG	VINBLASTINE SULFATE (VIAL) 10 MG	1 EA	VL	IV	EA		1 MG		10	01/01/2002	09/05/2014						
55390-0100-10		J0592		06/03/2005	09/05/2014	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE 0.3 MG/ML	1 ML	VL	IJ	ML		0.1 MG		3.24	06/03/2005	09/05/2014						
55390-0101-10		J3105		04/28/2004	99/99/9999	INJECTION, TERBUTALINE SULFATE, UP TO 1 MG	TERBUTALINE SULFATE 1 MG/ML	2 ML	VL	SC	ML		1 MG		1	04/28/2004	99/99/9999						
54569-1818-02	None			02/08/2018	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	100 EA	BO	PO	EA		2.5 MG		1	02/08/2018	99/99/9999						
00074-0243-02		J0135		05/01/2018	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PF,LATEX-FREE) 40 MG/0.4 ML	2 EA	BX	SC	EA		20 MG		2	05/01/2018	99/99/9999						
00074-0554-02		J0135		05/01/2018	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PF,LATEX-FREE) 40 MG/0.4 ML	2 EA	BX	SC	EA		20 MG		2	05/01/2018	99/99/9999						
55390-0106-01		J9999		09/01/2004	09/05/2014	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	ALLOPURINOL SODIUM (S.D.V.,PF) 500 MG	1 EA	VL	IV	EA		1 EA		1	09/01/2004	09/05/2014						
55390-0108-01		J9150		01/01/2002	09/05/2014	INJECTION, DAUNORUBICIN, 10 MG	DAUNORUBICIN HCL (S.D.V.,PF) 5 MG/ML	10 ML	VL	IV	ML		10 MG		0.5	01/01/2002	09/05/2014						
55390-0108-10		J9150		01/01/2002	09/05/2014	INJECTION, DAUNORUBICIN, 10 MG	DAUNORUBICIN HCL (S.D.V.,PF) 5 MG/ML	4 ML	VL	IV	ML		10 MG		0.5	01/01/2002	09/05/2014						
55390-0113-01		J2760		01/01/2002	01/05/2015	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (S.D.V.) 5 MG	1 EA	VL	IJ	EA		5 MG		1	01/01/2002	01/05/2015						
00074-0616-02		J0135		05/01/2018	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PF,LATEX-FREE) 20 MG/0.2 ML	2 EA	BX	SC	EA		20 MG		1	05/01/2018	99/99/9999						
00074-0817-02		J0135		05/01/2018	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PF,LATEX-FREE) 10 MG/0.1 ML	2 EA	BX	SC	EA		20 MG		0.5	05/01/2018	99/99/9999						
55390-0114-05		J9265		01/01/2002	09/05/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (M.D.V.) 6 MG/ML	5 ML	VL	IV	ML		30 MG		0.2	01/01/2002	09/05/2014						
55390-0114-20		J9265		01/01/2002	09/05/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (M.D.V.) 6 MG/ML	16.7 ML	VL	IV	ML		30 MG		0.2	01/01/2002	09/05/2014						
55390-0114-50		J9265		01/01/2002	09/05/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (M.D.V.) 6 MG/ML	50 ML	VL	IV	ML		30 MG		0.2	01/01/2002	09/05/2014						
55390-0115-01		J9065		01/01/2002	04/18/2013	INJECTION, CLADRIBINE, PER 1 MG	CLADRIBINE NOVAPLUS (S.D.V.,PF) 1 MG/ML	10 ML	VL	IV	ML		1 MG		1	01/01/2002	04/18/2013						
55390-0121-01		J2405		12/26/2006	03/14/2016	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV,USP) 2 MG/ML	20 ML	VL	IJ	ML		1 MG		2	12/26/2006	03/14/2016						
00074-2540-03		J0135		05/01/2018	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA PEDIATRIC CROHN'S DISEASE STARTER PACK (PF,LATEX-FREE) 80 MG/0.8 ML	3 EA	BX	SC	EA		20 MG		4	05/01/2018	99/99/9999						
55390-0122-10		J7516		01/01/2002	09/05/2014	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORINE (S.D.V.) 50 MG/ML	5 ML	VL	IV	ML		250 MG		0.2	01/01/2002	09/05/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55390-0123-01		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	RIFAMPIN (VIAL,30 ML) 600 MG	1 EA	VL	IV	EA		1 EA		1	01/01/2002	99/99/9999						
55390-0124-01		J9065		01/01/2002	09/05/2014	INJECTION, CLADRIBINE, PER 1 MG	CLADRIBINE (S.D.V.,PF) 1 MG/ML	10 ML	VL	IV	ML		1 MG		1	01/01/2002	09/05/2014						
55390-0125-10		J2250		01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 1 MG/ML	10 ML	VL	U	ML		1 MG		1	01/01/2002	99/99/9999						
55390-0126-05		J2250		01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 5 MG/ML	5 ML	VL	U	ML		1 MG		5	01/01/2002	99/99/9999						
55390-0126-10		J2250		01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 5 MG/ML	10 ML	VL	U	ML		1 MG		5	01/01/2002	99/99/9999						
55390-0127-01		J2430		01/01/2002	09/05/2014	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (VIAL) 30 MG	1 EA	VL	IV	EA		30 MG		1	01/01/2002	09/05/2014						
55390-0129-01		J2430		01/01/2002	09/05/2014	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (VIAL) 90 MG	1 EA	VL	IV	EA		30 MG		3	01/01/2002	09/05/2014						
55390-0131-10		J9100		01/01/2002	09/05/2014	INJECTION, CYTARABINE, 100 MG	CYTARABINE (VIAL) 100 MG	1 EA	VL	U	EA		100 MG		1	01/01/2002	09/05/2014						
55390-0135-01		J9200		01/01/2002	09/05/2014	INJECTION, FLOXURIDINE, 500 MG	FLOXURIDINE (VIAL) 0.5 GM	1 EA	VL	U	EA		500 MG		1	01/01/2002	09/05/2014						
55390-0136-05		J1955		01/01/2002	09/05/2014	INJECTION, LEVOCARNITINE, PER 1 GM	LEVOCARNITINE (S.D.V.) 200 MG/ML	5 ML	VL	IV	ML		1 GM	0.2	01/01/2002	09/05/2014							
55390-0137-02		J2250		01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 1 MG/ML	2 ML	VL	U	ML		1 MG		1	01/01/2002	99/99/9999						
55390-0137-05		J2250		01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 1 MG/ML	5 ML	VL	U	ML		1 MG		1	01/01/2002	99/99/9999						
55390-0138-01		J2250		01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 5 MG/ML	1 ML	VL	U	ML		1 MG		5	01/01/2002	99/99/9999						
55390-0138-02		J2250		01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (VIAL,PF) 5 MG/ML	2 ML	VL	U	ML		1 MG		5	01/01/2002	99/99/9999						
55390-0142-10		J9150		01/01/2002	04/18/2013	INJECTION, DAUNORUBICIN, 10 MG	DAUNORUBICIN HCL NOVAPLUS (S.D.V.,PF) 5 MG/ML	4 ML	VL	IV	ML		10 MG		0.5	01/01/2002	04/18/2013						
55390-0143-01		J9260		09/07/2005	09/05/2014	METHOTREXATE SODIUM, 50 MG	METHOTREXATE SODIUM (S.D.V.,30ML VIAL,PF) 1 GM	1 EA	VL	U	EA		50 MG		20	09/07/2005	09/05/2014						
55390-0147-01		J1630		01/01/2002	09/05/2014	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL LACTATE (M.D.V.) 5 MG/ML	10 ML	VL	IM	ML		5 MG		1	01/01/2002	09/05/2014						
55390-0147-10		J1630		01/01/2002	09/05/2014	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL LACTATE (S.D.V.) 5 MG/ML	1 ML	VL	IM	ML		5 MG		1	01/01/2002	09/05/2014						
00143-9240-01		J9040		05/16/2018	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN (USP LYOPHILIZED) 15 U	1 EA	VL	U	EA		15 U		1	05/16/2018	99/99/9999						
00143-9241-01		J9040		05/16/2018	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN (USP LYOPHILIZED) 30 U	1 EA	VL	U	EA		15 U		2	05/16/2018	99/99/9999						
00338-9572-24		J0583		05/01/2018	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN-SODIUM CHLORIDE 250 MG/50 ML-0.9%	50 ML	BG	IV	ML		1 MG		5	05/01/2018	99/99/9999						
00338-9576-12		J0583		05/01/2018	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN-SODIUM CHLORIDE 500 MG/100 ML-0.9%	100 ML	BG	IV	ML		1 MG		5	05/01/2018	99/99/9999						
00517-1133-01		J2710		05/11/2018	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (INNER PACK,LATEX-FREE) 0.5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		1	05/11/2018	99/99/9999						
55390-0157-01		J2430		01/01/2003	04/18/2013	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (LYOPHILIZED) 30 MG	1 EA	VL	IV	EA		30 MG		1	01/01/2003	04/18/2013						
55390-0159-01		J2430		01/01/2003	04/18/2013	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (LYOPHILIZED) 90 MG	1 EA	VL	IV	EA		30 MG		3	01/01/2003	04/18/2013						
55390-0160-10		J2354		05/04/2005	09/05/2014	INJECTION, 25 MCG	OCTREOTIDE 50 MCG/ML	1 ML	VL	U	ML		25 MCG		2	05/04/2005	09/05/2014						
55390-0161-10		J2354		04/04/2005	09/05/2014	INJECTION, 25 MCG	OCTREOTIDE 100 MCG/ML	1 ML	VL	U	ML		25 MCG		4	04/04/2005	09/05/2014						
00517-1133-05		J2710		05/11/2018	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 0.5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		1	05/11/2018	99/99/9999						
55390-0162-10		J2354		04/04/2005	09/05/2014	INJECTION, 25 MCG	OCTREOTIDE 500 MCG/ML	1 ML	VL	U	ML		25 MCG		20	04/04/2005	09/05/2014						
55390-0163-01		J2354		05/25/2005	09/05/2014	INJECTION, 25 MCG	OCTREOTIDE ACETATE (MDV) 200 MCG/ML	5 ML	VL	U	ML		25 MCG		8	05/25/2005	09/05/2014						
55390-0164-01		J2354		05/25/2005	01/14/2016	INJECTION, 25 MCG	OCTREOTIDE ACETATE (MDV) 1000 MCG/ML	5 ML	VL	U	ML		25 MCG		40	05/25/2005	01/14/2016						
00517-1134-01		J2710		05/11/2018	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (INNER PACK,LATEX-FREE) 1 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		2	05/11/2018	99/99/9999						
00517-1134-05		J2710		05/11/2018	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 1 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		2	05/11/2018	99/99/9999						
55390-0183-01		J0595		01/01/2004	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (S.D.V.) 1 MG/ML	1 ML	VL	U	ML		1 MG		1	01/01/2004	99/99/9999						
55390-0184-01		J0595		01/01/2004	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (S.D.V.) 2 MG/ML	1 ML	VL	U	ML		1 MG		2	01/01/2004	99/99/9999						
25021-0184-82		J1450		04/23/2018	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (10X100ML,PF,LATEX-FREE) 200 MG/100 ML	100 ML	FC	IV	ML		200 MG		0.01	04/23/2018	99/99/9999						
25021-0184-87		J1450		04/23/2018	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (10X200ML,PF,LATEX-FREE) 400 MG/200 ML	200 ML	FC	IV	ML		200 MG		0.01	04/23/2018	99/99/9999						
55390-0193-10		J3105		11/19/2004	04/18/2013	INJECTION, TERBUTALINE SULFATE, UP TO 1 MG	TERBUTALINE SULFATE NOVAPLUS 1 MG/ML	1 ML	VL	SC	ML		1 MG		1	11/19/2004	04/18/2013						
25021-0186-20		J0295		04/23/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM (USP,SDV,PF,LATEX-FREE) 1 GM-0.5 GM	10 EA	VL	U	EA		1.5 GM		1	04/23/2018	99/99/9999						
25021-0187-30		J0295		04/23/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM (USP,SDV,PF,LATEX-FREE) 2 GM-1 GM	10 EA	VL	U	EA		1.5 GM		2	04/23/2018	99/99/9999						
25021-0188-99		J0295		04/23/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM (PHARMACY BULK,USP,PF) 10 GM-5 GM	1 EA	VL	IV	EA		1.5 GM		10	04/23/2018	99/99/9999						
47781-0622-22		J9209		04/24/2018	99/99/9999	INJECTION, MESNA, 200 MG	MESNA 100 MG/1 ML	10 ML	VL	IV	ML		200 MG		0.5	04/24/2018	99/99/9999						
47781-0622-91		J9209		04/24/2018	99/99/9999	INJECTION, MESNA, 200 MG	MESNA 100 MG/1 ML	10 ML	VL	IV	ML		200 MG		0.5	04/24/2018	99/99/9999						
55390-0220-01		J9045		11/19/2004	06/07/2012	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN AMERINET CHOICE (M.D.V.,PF) 10 MG/ML	5 ML	VL	IV	ML		50 MG		0.2	11/19/2004	06/07/2012						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55390-0221-01	J9045			11/19/2004	06/07/2012	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN AMERINET CHOICE (M.D.V.,PF) 10 MG/ML	15 ML	VL	IV	ML		50 MG		0.2	11/19/2004	06/07/2012						
55390-0222-01	J9045			11/19/2004	06/07/2012	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN AMERINET CHOICE (M.D.V.,PF) 10 MG/ML	45 ML	VL	IV	ML		50 MG		0.2	11/19/2004	06/07/2012						
47781-0623-07	J0895			04/26/2018	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE (USP,PF,LATEX-FREE) 500 MG	1 EA	VL	IJ	EA		500 MG		1	04/26/2018	99/99/9999						
47781-0624-07	J0895			04/26/2018	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE (USP,PF,LATEX-FREE) 2 GM	1 EA	VL	IJ	EA		500 MG		4	04/26/2018	99/99/9999						
52609-4505-06	J0895			04/16/2018	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE (USP,SINGLE USE) 500 MG	4 EA	VL	IJ	EA		500 MG		1	04/16/2018	99/99/9999						
55390-0226-02	J0278			01/01/2006	01/14/2016	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (S.D.V.,PF) 250 MG/ML	2 ML	VL	IJ	ML		100 MG		2.5	01/01/2006	01/14/2016						
55390-0226-04	J0278			01/01/2006	09/05/2014	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (PF) 250 MG/ML	4 ML	VL	IJ	ML		100 MG		2.5	01/01/2006	09/05/2014						
55390-0231-10	J9000			01/01/2002	09/05/2014	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 10 MG	1 EA	VL	IV	EA		10 MG		1	01/01/2002	09/05/2014						
55150-0180-03	J0282			05/04/2018	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL 50 MG/1 ML	3 ML	VL	IV	ML		30 MG	1.66666		05/04/2018	99/99/9999						
55150-0181-09	J0282			05/04/2018	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL 50 MG/1 ML	9 ML	VL	IV	ML		30 MG	1.66666		05/04/2018	99/99/9999						
55150-0182-18	J0282			05/04/2018	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL 50 MG/1 ML	18 ML	VL	IV	ML		30 MG	1.66666		05/04/2018	99/99/9999						
55390-0232-10	J9000			01/01/2002	09/05/2014	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 20 MG	1 EA	VL	IV	EA		10 MG		2	01/01/2002	09/05/2014						
55390-0233-01	J9000			01/01/2002	09/05/2014	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 50 MG	1 EA	VL	IV	EA		10 MG		5	01/01/2002	09/05/2014						
55390-0235-10	J9000			01/01/2002	09/05/2014	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.) 2 MG/ML	5 ML	VL	IV	ML		10 MG		0.2	01/01/2002	09/05/2014						
55390-0236-10	J9000			01/01/2002	09/05/2014	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 2 MG/ML	10 ML	VL	IV	ML		10 MG		0.2	01/01/2002	09/05/2014						
55390-0237-01	J9000			01/01/2002	09/05/2014	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.) 2 MG/ML	25 ML	VL	IV	ML		10 MG		0.2	01/01/2002	09/05/2014						
55390-0238-01	J9000			01/01/2002	09/05/2014	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (M.D.V.) 2 MG/ML	100 ML	VL	IV	ML		10 MG		0.2	01/01/2002	09/05/2014						
55390-0241-10	J9000			01/01/2002	04/18/2013	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL NOVAPLUS (S.D.V.,PF) 10 MG	1 EA	VL	IV	EA		10 MG		1	01/01/2002	04/18/2013						
55390-0243-01	J9000			01/01/2002	04/18/2013	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL NOVAPLUS (S.D.V.,PF) 50 MG	1 EA	VL	IV	EA		10 MG		5	01/01/2002	04/18/2013						
55390-0244-01	J9268			08/08/2007	09/05/2014	INJECTION, PENTOSTATIN, 10 MG	PENTOSTATIN (SDV) 10 MG	1 EA	VL	IV	EA		10 MG		1	08/08/2007	09/05/2014						
55390-0245-10	J9000			01/01/2002	04/18/2013	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL NOVAPLUS (S.D.V.,PF) 2 MG/ML	5 ML	VL	IV	ML		10 MG		0.2	01/01/2002	04/18/2013						
55390-0246-10	J9000			01/01/2002	04/18/2013	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL NOVAPLUS (S.D.V.,PF) 2 MG/ML	10 ML	VL	IV	ML		10 MG		0.2	01/01/2002	04/18/2013						
55390-0247-01	J9000			01/01/2002	04/18/2013	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL NOVAPLUS (S.D.V.,PF) 2 MG/ML	25 ML	VL	IV	ML		10 MG		0.2	01/01/2002	04/18/2013						
55390-0248-01	J9000			01/01/2002	04/18/2013	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL NOVAPLUS (M.D.V.) 2 MG/ML	100 ML	VL	IV	ML		10 MG		0.2	01/01/2002	04/18/2013						
55390-0248-10	J0895			06/18/2007	09/05/2014	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE (USP) 500 MG	1 EA	VL	IJ	EA		500 MG		1	06/18/2007	09/05/2014						
55390-0265-01	J0895			06/18/2007	09/05/2014	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFEROXAMINE MESYLATE (USP) 2 GM	1 EA	VL	IJ	EA		500 MG		4	06/18/2007	09/05/2014						
55150-0267-05	J2680			04/21/2018	99/99/9999	INJECTION, FLUPHENAZINE DECANOATE, UP TO 25 MG	FLUPHENAZINE DECANOATE (MDV,LATEX-FREE) 25 MG/1 ML	5 ML	VL	IJ	ML		25 MG		1	04/21/2018	99/99/9999						
58463-0010-08	J8540			04/18/2018	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DECADRON (RASPBERRY) 0.5 MG/5 ML	237 ML	BO	PO	ML		0.25 MG		0.4	04/18/2018	99/99/9999						
60842-0021-01	J0171			04/18/2018	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	AUVI-Q 0.1 MG/0.1 ML	2 EA	SR	IJ	EA		0.1 MG		1	04/18/2018	99/99/9999						
55390-0266-01	J9209			09/21/2005	06/07/2012	INJECTION, MESNA, 200 MG	MESNA AMERINET CHOICE (M.D.V.) 100 MG/ML	10 ML	VL	IV	ML		200 MG		0.5	09/21/2005	06/07/2012						
55390-0267-01	J9390			09/21/2005	06/07/2012	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE AMERINET CHOICE (S.D.V.,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	09/21/2005	06/07/2012						
55390-0268-01	J9390			09/21/2005	06/07/2012	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE AMERINET CHOICE (S.D.V.,PF) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	09/21/2005	06/07/2012						
55390-0281-10	J9150			01/01/2002	09/05/2014	INJECTION, DAUNORUBICIN, 10 MG	CERUBIDINE (S.D.V.) 20 MG	1 EA	VL	IV	EA		10 MG		2	01/01/2002	09/05/2014						
55390-0291-01	J9181			01/01/2002	09/05/2014	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V.) 20 MG/ML	5 ML	VL	IV	ML		10 MG		2	01/01/2002	09/05/2014						
55390-0292-01	J9181			01/01/2002	09/05/2014	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V.) 20 MG/ML	25 ML	VL	IV	ML		10 MG		2	01/01/2002	09/05/2014						
55390-0293-01	J9181			01/01/2002	09/05/2014	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V.) 20 MG/ML	50 ML	VL	IV	ML		10 MG		2	01/01/2002	09/05/2014						
55390-0304-05	J9265			12/04/2006	04/18/2013	INJECTION, PACLITAXEL, 30 MG	NOVAPLUS PACLITAXEL (MDV,USP) 6 MG/ML	5 ML	VL	IV	ML		30 MG		0.2	12/04/2006	04/18/2013						
55390-0304-20	J9265			12/04/2006	04/18/2013	INJECTION, PACLITAXEL, 30 MG	NOVAPLUS PACLITAXEL (MDV,USP) 6 MG/ML	16.7 ML	VL	IV	ML		30 MG		0.2	12/04/2006	04/18/2013						
55390-0304-50	J9265			12/04/2006	04/18/2013	INJECTION, PACLITAXEL, 30 MG	NOVAPLUS PACLITAXEL (MULTIPLE-DOSE,USP) 6 MG/ML	50 ML	VL	IV	ML		30 MG		0.2	12/04/2006	04/18/2013						
61314-0318-01	Q5101			05/04/2018	99/99/9999	INJECTION, FILGRASTIM-SNDZ, BIOSIMILAR, (ZARXIO), 1 MICROGRAM	ZARXIO (PF) 300 MCG/0.5 ML	0.5 ML	SR	IJ	ML		1 MCG		600	05/04/2018	99/99/9999						
61314-0326-01	Q5101			05/04/2018	99/99/9999	INJECTION, FILGRASTIM-SNDZ, BIOSIMILAR, (ZARXIO), 1 MICROGRAM	ZARXIO (PF) 480 MCG/0.8 ML	0.8 ML	SR	IJ	ML		1 MCG		600	05/04/2018	99/99/9999						
55390-0308-03	J0207			04/08/2008	12/31/2016	INJECTION, AMIFOSTINE, 500 MG	AMIFOSTINE (3X10ML,LYOPHILIZED) 500 MG	1 EA	VL	IV	EA		500 MG		1	04/08/2008	12/31/2016						
55390-0314-05	J9265			01/14/2004	06/07/2012	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL AMERINET CHOICE (M.D.V.) 6 MG/ML	5 ML	VL	IV	ML		30 MG		0.2	01/14/2004	06/07/2012						
55390-0314-20	J9265			01/14/2004	06/07/2012	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL AMERINET CHOICE (M.D.V.) 6 MG/ML	16.7 ML	VL	IV	ML		30 MG		0.2	01/14/2004	06/07/2012						
55390-0314-50	J9265			01/14/2004	06/07/2012	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL AMERINET CHOICE (M.D.V.) 6 MG/ML	50 ML	VL	IV	ML		30 MG		0.2	01/14/2004	06/07/2012						
55390-0347-01	J9209			03/05/2008	04/18/2013	INJECTION, MESNA, 200 MG	NOVAPLUS MESNA (1X10ML,M.D.V) 100 MG/ML	10 ML	VL	IV	ML		200 MG		0.5	03/05/2008	04/18/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
55390-0403-20	J2400			01/01/2002	09/05/2014	INJECTION, CHLOROPROCAINE HYDROCHLORIDE, PER 30 ML	CHLOROPROCAINE HCL (S.D.V.,PF) 2%	20	ML	VL	IJ	ML	30	ML	0.03333	01/01/2002	09/05/2014							
55390-0404-20	J2400			01/01/2002	09/05/2014	INJECTION, CHLOROPROCAINE HYDROCHLORIDE, PER 30 ML	CHLOROPROCAINE HCL (S.D.V.,PF) 3%	20	ML	VL	IJ	ML	30	ML	0.03333	01/01/2002	09/05/2014							
63323-0203-26	J3370			05/02/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	PREMIERPRO RX VANCOMYCIN HCL 750 MG	10	EA	VL	IV	EA	500	MG	1.5	05/02/2018	99/99/9999							
63323-0651-20	J0153			05/02/2018	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (SDV,PF,LATEX-FREE) 3 MG/1 ML	20	ML	VL	IV	ML	1	MG	3	05/02/2018	99/99/9999							
63323-0651-30	J0153			05/02/2018	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (SDV,PF,LATEX-FREE) 3 MG/1 ML	30	ML	VL	IV	ML	1	MG	3	05/02/2018	99/99/9999							
64011-0301-03	J1726			02/14/2018	99/99/9999	INJECTION, HYDROXYPROGESTERONE CAPROATE, (MAKENA), 10 MG	MAKENA (PF) 275 MG/1.1 ML	1.1	ML	VL	SC	ML	10	MG	25	02/14/2018	99/99/9999							
55390-0460-01	J1120			01/01/2002	09/05/2014	INJECTION, ACETAZOLAMIDE SODIUM, UP TO 500 MG	ACETAZOLAMIDE SODIUM (S.D.V.,PF) 500 MG	1	EA	VL	IV	EA	500	MG	1	01/01/2002	09/05/2014							
55390-0480-01	J1885			01/01/2002	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 15 MG/ML	1	ML	VL	IJ	ML	15	MG	1	01/01/2002	99/99/9999							
55390-0481-01	J1885			01/01/2002	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 30 MG/ML	1	ML	VL	IJ	ML	15	MG	2	01/01/2002	99/99/9999							
55390-0481-02	J1885			01/01/2002	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 30 MG/ML	2	ML	VL	IM	ML	15	MG	2	01/01/2002	99/99/9999							
67457-0854-04	J0153			05/08/2018	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (10X4ML,SDV,PF) 3 MG/1 ML	4	ML	VL	IV	ML	1	MG	3	05/08/2018	99/99/9999							
55390-0481-10	J1885			01/01/2002	11/30/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (M.D.V.) 30 MG/ML	10	ML	VL	IJ	ML	15	MG	2	01/01/2002	11/30/2013							
55390-0491-01	J9181			01/01/2002	04/18/2013	INJECTION, ETOPOSID, 10 MG	ETOPOSID NOVAPLUS (M.D.V.) 20 MG/ML	5	ML	VL	IV	ML	10	MG	2	01/01/2002	04/18/2013							
55390-0492-01	J9181			01/01/2002	04/18/2013	INJECTION, ETOPOSID, 10 MG	ETOPOSID NOVAPLUS (M.D.V.) 20 MG/ML	25	ML	VL	IV	ML	10	MG	2	01/01/2002	04/18/2013							
55390-0493-01	J9181			01/01/2002	04/18/2013	INJECTION, ETOPOSID, 10 MG	ETOPOSID NOVAPLUS (M.D.V.) 20 MG/ML	50	ML	VL	IV	ML	10	MG	2	01/01/2002	04/18/2013							
55390-0500-02	J3490			01/01/2002	04/30/2013	UNCLASSIFIED DRUGS	BUMETANIDE (S.D.V.) 0.25 MG/ML	2	ML	VL	IJ	ML	1	EA	1	01/01/2002	04/30/2013							
55390-0500-05	J3490			01/01/2002	04/30/2013	UNCLASSIFIED DRUGS	BUMETANIDE (S.D.V.) 0.25 MG/ML	4	ML	VL	IJ	ML	1	EA	1	01/01/2002	04/30/2013							
55390-0500-10	J3490			01/01/2002	04/30/2013	UNCLASSIFIED DRUGS	BUMETANIDE (M.D.V.) 0.25 MG/ML	10	ML	VL	IJ	ML	1	EA	1	01/01/2002	04/30/2013							
67457-0855-02	J0153			05/08/2018	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (10X2ML,SDV,PF) 3 MG/1 ML	2	ML	VL	IV	ML	1	MG	3	05/08/2018	99/99/9999							
68001-0246-04	Q0162			04/24/2018	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,3X10,STRAWBERRY) 4 MG	30	EA	ST	PO	EA	1	MG	4	04/24/2018	99/99/9999							
68001-0247-04	Q0162			04/24/2018	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP, 3X10,STRAWBERRY) 8 MG	30	EA	ST	PO	EA	1	MG	8	04/24/2018	99/99/9999							
68001-0342-34	J9201			05/01/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 100 MG/1 ML	2	ML	VL	IV	ML	200	MG	0.5	05/01/2018	99/99/9999							
55390-0560-90	J1250			01/01/2002	09/05/2014	INJECTION, DOBUTAMINE HYDROCHLORIDE, PER 250 MG	DOBUTAMINE HCL (S.D.V.,PF) 12.5 MG/ML	20	ML	VL	IV	ML	250	MG	0.05	01/01/2002	09/05/2014							
55390-0600-20	J7501			01/01/2002	09/05/2014	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE SODIUM (PF) 100 MG	1	EA	VL	IV	EA	100	MG	1	01/01/2002	09/05/2014							
55390-0612-10	J0133			01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR SODIUM (PF) 500 MG	1	EA	VL	IV	EA	5	MG	100	01/01/2006	99/99/9999							
55390-0613-20	J0133			01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR SODIUM (PF) 1000 MG	1	EA	VL	IV	EA	5	MG	200	01/01/2006	99/99/9999							
55390-0616-01	J2780			11/22/2004	09/05/2014	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE (M.D.V.) 25 MG/ML	6	ML	VL	IJ	ML	25	MG	1	11/22/2004	09/05/2014							
68001-0347-36	J0894			05/01/2018	99/99/9999	INJECTION, DECTABINE, 1 MG	DECTABINE (LYOPHILIZED) 50 MG	1	EA	VL	IV	EA	1	MG	50	05/01/2018	99/99/9999							
55390-0616-10	J2780			11/22/2004	09/05/2014	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE (S.D.V.) 25 MG/ML	2	ML	VL	IJ	ML	25	MG	1	11/22/2004	09/05/2014							
55390-0618-01	J2780			03/29/2006	09/05/2014	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE (PHARMACY BULK PACKAGE) 25 MG/ML	40	ML	VL	IJ	ML	25	MG	1	03/29/2006	09/05/2014							
55390-0805-10	J9150			01/01/2002	04/18/2013	INJECTION, DAUNORUBICIN, 10 MG	DAUNORUBICIN HCL NOVAPLUS (S.D.V.) 20 MG	1	EA	VL	IV	EA	10	MG	2	01/01/2002	04/18/2013							
55390-0806-10	J9100			01/01/2002	04/18/2013	INJECTION, CYTARABINE, 100 MG	CYTARABINE NOVAPLUS (VIAL) 100 MG	1	EA	VL	IJ	EA	100	MG	1	01/01/2002	04/18/2013							
55390-0818-10	J0640			01/01/2002	04/18/2013	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM NOVAPLUS (VIAL) 100 MG	1	EA	VL	IJ	EA	50	MG	2	01/01/2002	04/18/2013							
55390-0824-01	J0640			01/01/2002	04/18/2013	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM NOVAPLUS (VIAL) 200 MG	1	EA	VL	IJ	EA	50	MG	4	01/01/2002	04/18/2013							
55390-0825-01	J0640			01/01/2002	04/18/2013	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM NOVAPLUS (S.D.V.,PF) 350 MG	1	EA	VL	IJ	EA	50	MG	7	01/01/2002	04/18/2013							
55390-0826-01	J0640			01/01/2002	04/18/2013	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM NOVAPLUS (S.D.V.,PF) 10 MG/ML	50	ML	VL	IJ	ML	50	MG	0.2	01/01/2002	04/18/2013							
55513-0002-01	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.025 MG/ML	1	ML	VL	IJ	ML	1	MCG	25	09/11/2006	99/99/9999							
55513-0002-04	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (4X1ML,PF) 0.025 MG/ML	1	ML	VL	IJ	ML	1	MCG	25	09/11/2006	99/99/9999							
55513-0003-01	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.04 MG/ML	1	ML	VL	IJ	ML	1	MCG	40	09/11/2006	99/99/9999							
55513-0003-04	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (1MLX4,PF) 0.04 MG/ML	1	ML	VL	IJ	ML	1	MCG	40	09/11/2006	99/99/9999							
55513-0004-01	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.06 MG/ML	1	ML	VL	IJ	ML	1	MCG	60	09/11/2006	99/99/9999							
55513-0004-04	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (1MLX4,PF) 0.06 MG/ML	1	ML	VL	IJ	ML	1	MCG	60	09/11/2006	99/99/9999							
55513-0005-01	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.1 MG/ML	1	ML	VL	IJ	ML	1	MCG	100	09/11/2006	99/99/9999							
55513-0005-04	J0881			09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (1MLX4,PF) 0.1 MG/ML	1	ML	VL	IJ	ML	1	MCG	100	09/11/2006	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55513-0006-01		J0881		09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.2 MG/ML	1 ML	VL	IJ	ML		1 MCG		200	09/11/2006	99/99/9999						
55513-0021-01		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.04 MG/0.4 ML	0.4 ML	SR	IJ	ML		1 MCG		100	08/14/2006	99/99/9999						
55513-0021-04		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.04 MG/0.4 ML	0.4 ML	SR	IJ	ML		1 MCG		100	08/14/2006	99/99/9999						
55513-0023-01		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.06 MG/0.3 ML	0.3 ML	SR	IJ	ML		1 MCG		200	08/14/2006	99/99/9999						
55513-0023-04		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.06 MG/0.3 ML	0.3 ML	SR	IJ	ML		1 MCG		200	08/14/2006	99/99/9999						
55513-0025-01		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.1 MG/0.5 ML	0.5 ML	SR	IJ	ML		1 MCG		200	08/14/2006	99/99/9999						
55513-0025-04		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.1 MG/0.5 ML	0.5 ML	SR	IJ	ML		1 MCG		200	08/14/2006	99/99/9999						
55513-0027-01		J0881		09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.15 MG/0.3 ML	0.3 ML	SR	IJ	ML		1 MCG		500	09/11/2006	99/99/9999						
55513-0027-04		J0881		09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (0.3MLX4,PF) 0.15 MG/0.3 ML	0.3 ML	SR	IJ	ML		1 MCG		500	09/11/2006	99/99/9999						
55513-0028-01		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.2 MG/0.4 ML	0.4 ML	SR	IJ	ML		1 MCG		500	08/14/2006	99/99/9999						
55513-0032-01		J0881		06/07/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (SINGLEJECT,G27,1/2",PF) 0.5 MG/ML	1 ML	SR	IJ	ML		1 MCG		500	06/07/2006	99/99/9999						
55513-0053-01		J0881		09/11/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.15 MG/0.75 ML	1 ML	VL	IJ	ML		1 MCG		200	09/11/2006	99/99/9999						
55513-0053-04		J0881		09/11/2006	12/02/2014	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (1MLX4,PF) 0.15 MG/0.75 ML	1 ML	VL	IJ	ML		1 MCG		200	09/11/2006	12/02/2014						
55513-0057-01		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.025 MG/0.42 ML	0.42 ML	SR	IJ	ML		1 MCG	59.52381	200	08/14/2006	99/99/9999						
55513-0057-04		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.025 MG/0.42 ML	0.42 ML	SR	IJ	ML		1 MCG	59.52381	200	08/14/2006	99/99/9999						
55513-0110-01		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF STERILE) 0.3 MG/ML	1 ML	VL	IJ	ML		1 MCG		300	08/14/2006	99/99/9999						
55513-0111-01		J0881		08/14/2006	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	ARANESP (PF) 0.3 MG/0.5 ML	0.6 ML	SR	IJ	ML		1 MCG		500	08/14/2006	99/99/9999						
55513-0126-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S2,PF) 2000 U/ML	1 ML	VL	IJ	ML		1000 U		2	01/01/2006	99/99/9999						
55513-0126-10		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S2,PF) 2000 U/ML	1 ML	VL	IJ	ML		1000 U		2	01/01/2006	99/99/9999						
55513-0144-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S10,PF) 10000 U/ML	1 ML	VL	IJ	ML		1000 U		10	01/01/2006	99/99/9999						
55513-0144-10		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S10,PF) 10000 U/ML	1 ML	VL	IJ	ML		1000 U		10	01/01/2006	99/99/9999						
55513-0148-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S4,PF) 4000 U/ML	1 ML	VL	IJ	ML		1000 U		4	01/01/2006	99/99/9999						
55513-0148-10		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S4,PF) 4000 U/ML	1 ML	VL	IJ	ML		1000 U		4	01/01/2006	99/99/9999						
55513-0177-01		J3490		01/01/2002	04/10/2013	UNCLASSIFIED DRUGS	MG/0.67 ML KINERET (SRN,W/27G NDL,PF) 100	0.67 ML	SR	SC	ML		1 EA		1	01/01/2002	04/10/2013						
55513-0177-28		J3490		02/23/2004	04/10/2013	UNCLASSIFIED DRUGS	MG/0.67 ML KINERET (SRN,W/27G NDL,PF) 100	0.67 ML	SR	SC	ML		1 EA		1	02/23/2004	04/10/2013						
55513-0190-01		J2505		01/01/2004	99/99/9999	INJECTION, PEGFILGRASTIM, 6 MG	NEULASTA (SRN,PREFILLED,PF,4X0.6ML) 6 MG/0.6 ML	0.6 ML	SR	SC	ML		6 MG	1.66666		01/01/2004	99/99/9999						
55513-0209-01		J1441		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 480 MCG	NEUPOGEN (26GX5/8",PF,SINGLEJECT) 480 MCG/0.8 ML	0.8 ML	SR	IJ	ML		480 MCG		1.25	01/01/2002	12/31/2013						
55513-0209-10		J1441		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 480 MCG	NEUPOGEN (26GX5/8",10X0.8ML,PF,SINGLEJECT) 480 MCG/0.8 ML	0.8 ML	SR	IJ	ML		480 MCG		1.25	01/01/2002	12/31/2013						
55513-0267-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S3,PF) 3000 U/ML	1 ML	VL	IJ	ML		1000 U		3	01/01/2006	99/99/9999						
55513-0267-10		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (S.D.V.,S3,PF) 3000 U/ML	1 ML	VL	IJ	ML		1000 U		3	01/01/2006	99/99/9999						
55513-0283-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (M.D.V.,M10) 10000 U/ML	2 ML	VL	IJ	ML		1000 U		10	01/01/2006	99/99/9999						
55513-0283-10		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (M.D.V.,M10) 10000 U/ML	2 ML	VL	IJ	ML		1000 U		10	01/01/2006	99/99/9999						
55513-0478-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (M.D.V.,M20) 20000 U/ML	1 ML	VL	IJ	ML		1000 U		20	01/01/2006	99/99/9999						
55513-0478-10		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	EPOGEN (M.D.V.,M20) 20000 U/ML	1 ML	VL	IJ	ML		1000 U		20	01/01/2006	99/99/9999						
55513-0530-01		J1440		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 300 MCG	NEUPOGEN (S.D.V.,PF) 300 MCG/ML NEUPOGEN (S.D.V.,1MLX10,PF) 300 MCG/ML	1 ML	VL	IJ	ML		300 MCG		1	01/01/2002	12/31/2013						
55513-0530-10		J1440		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 300 MCG	NEUPOGEN (S.D.V.,PF) 300 MCG/ML	1 ML	VL	IJ	ML		300 MCG		1	01/01/2002	12/31/2013						
55513-0546-01		J1441		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 480 MCG	NEUPOGEN (S.D.V.,PF) 480 MCG/1.6 ML NEUPOGEN (S.D.V.,1.6MLX10,PF) 480 MCG/1.6 ML	1.6 ML	VL	IJ	ML		480 MCG	0.625		01/01/2002	12/31/2013						
55513-0546-10		J1441		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 480 MCG	NEUPOGEN (26GX5/8",SINGLE USE) 300 MCG/0.5 ML	0.5 ML	SR	IJ	ML		480 MCG	0.625		01/01/2002	12/31/2013						
55513-0924-01		J1440		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 300 MCG	NEUPOGEN ((26GX5/8"),0.5MLX10,PF) 300 MCG/0.5 ML	0.5 ML	SR	IJ	ML		300 MCG		2	01/01/2002	12/31/2013						
55513-0924-10		J1440		01/01/2002	12/31/2013	INJECTION, FILGRASTIM (G-CSF), 300 MCG	NEUPOGEN ((26GX5/8"),0.5MLX10,PF) 300 MCG/0.5 ML	0.5 ML	SR	IJ	ML		300 MCG		2	01/01/2002	12/31/2013						
55513-0954-01		J9303		01/01/2008	99/99/9999	INJECTION, PANITUMUMAB, 10 MG	VECTIBX 20 MG/ML	5 ML	VL	IV	ML		10 MG		2	01/01/2008	99/99/9999						
55513-0956-01		J9303		01/01/2008	99/99/9999	INJECTION, PANITUMUMAB, 10 MG	VECTIBX 20 MG/ML	20 ML	VL	IV	ML		10 MG		2	01/01/2008	99/99/9999						
55553-0042-05		J3302		01/01/2002	05/15/2016	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	CLINACORT (VIAL) 40 MG/ML	5 ML	VL	IJ	ML		5 MG		8	01/01/2002	05/15/2016						
55553-0055-50		J2001		01/01/2004	02/10/2016	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	ANESTACAINE (VIAL) 1%	50 ML	VL	EP	ML		10 MG		1	01/01/2004	02/10/2016						
55553-0056-50		J2001		01/01/2004	02/10/2016	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	ANESTACAINE (VIAL) 2%	50 ML	VL	IJ	ML		10 MG		2	01/01/2004	02/10/2016						
55553-0091-10		J3420		01/01/2002	02/03/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	VITA #12 (VIAL) 1000 MCG/ML	10 ML	VL	IM	ML		1000 MCG		1	01/01/2002	02/03/2016						
55553-0091-30		J3420		01/01/2002	02/03/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	VITA #12 (VIAL) 1000 MCG/ML	30 ML	VL	IM	ML		1000 MCG		1	01/01/2002	02/03/2016						
55553-0092-05		J1094		01/01/2003	02/03/2016	INJECTION, DEXAMETHASONE ACETATE, 1 MG	CORTASTAT LA (VIAL) 8 MG/ML	5 ML	VL	IJ	ML		1 MG		8	01/01/2003	0						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
58016-0217-07		J7506		01/01/2007	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	7	EA	NA	PO	EA	5	MG	4	01/01/2007	01/31/2014						
58016-0217-10		J7506		03/21/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	10	EA	BO	PO	EA	5	MG	4	03/21/2002	01/31/2014						
58016-0217-12		J7506		01/01/2007	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	12	EA	NA	PO	EA	5	MG	4	01/01/2007	01/31/2014						
58016-0217-15		J7506		03/21/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	15	EA	BO	PO	EA	5	MG	4	03/21/2002	01/31/2014						
58016-0217-16		J7506		03/21/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	16	EA	BO	PO	EA	5	MG	4	03/21/2002	01/31/2014						
58016-0217-18		J7506		03/21/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	18	EA	BO	PO	EA	5	MG	4	03/21/2002	01/31/2014						
58016-0217-20		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	20	EA	BO	PO	EA	5	MG	4	01/01/2002	01/31/2014						
58016-0217-21		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	21	EA	BO	PO	EA	5	MG	4	01/01/2002	01/31/2014						
58016-0217-22		J7506		03/21/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	22	EA	BO	PO	EA	5	MG	4	03/21/2002	01/31/2014						
58016-0217-23		J7506		01/01/2007	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	23	EA	NA	PO	EA	5	MG	4	01/01/2007	01/31/2014						
58016-0217-24		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	24	EA	BO	PO	EA	5	MG	4	01/01/2002	01/31/2014						
58016-0217-28		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	28	EA	BO	PO	EA	5	MG	4	01/01/2002	01/31/2014						
58016-0217-30		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	30	EA	BO	PO	EA	5	MG	4	01/01/2002	01/31/2014						
58016-0217-40		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	40	EA	BO	PO	EA	5	MG	4	01/01/2002	01/31/2014						
58016-0217-60		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 20 MG	60	EA	BO	PO	EA	5	MG	4	01/01/2002	01/31/2014						
58016-0218-00		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	100	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-20		J7506		03/22/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	20	EA	BO	PO	EA	5	MG	1	03/22/2002	01/31/2014						
58016-0218-21		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	21	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-24		J7506		03/22/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	24	EA	BO	PO	EA	5	MG	1	03/22/2002	01/31/2014						
58016-0218-30		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	30	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-33		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	33	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-36		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	36	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-40		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	40	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-50		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	50	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-55		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	55	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-60		J7506		01/01/2002	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	60	EA	BO	PO	EA	5	MG	1	01/01/2002	01/31/2014						
58016-0218-69		J7506		01/01/2007	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	69	EA	NA	PO	EA	5	MG	1	01/01/2007	01/31/2014						
58016-0218-90		J7506		05/31/2005	01/31/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	90	EA	BO	PO	EA	5	MG	1	05/31/2005	01/31/2014						
58016-0259-00		Q0177		01/01/2002	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	01/31/2014						
58016-0259-02		Q0177		01/01/2007	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	120	EA	NA	PO	EA	25	MG	1	01/01/2007	01/31/2014						
58016-0259-10		Q0177		01/01/2002	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	10	EA	BO	PO	EA	25	MG	1	01/01/2002	01/31/2014						
58016-0259-20		Q0177		01/01/2002	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20	EA	BO	PO	EA	25	MG	1	01/01/2002	01/31/2014						
58016-0259-30		Q0177		01/01/2002	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	BO	PO	EA	25	MG	1	01/01/2002	01/31/2014						
58016-0259-50		Q0177		01/01/2002	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	50	EA	BO	PO	EA	25	MG	1	01/01/2002	01/31/2014						
58016-0259-60		Q0177		01/01/2002	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	60	EA	BO	PO	EA	25	MG	1	01/01/2002	01/31/2014						
58016-0259-90		Q0177		01/01/2007	01/31/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	90	EA	NA	PO	EA	25	MG	1	01/01/2007	01/31/2014						
58016-0290-00		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	100	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-02		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	120	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-03		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	150	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-12		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	12	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-15		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	15	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-20		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	20	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-30		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	30	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-73		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	300	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0290-89		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	200	EA	BO	PO	EA	0.25	MG	2	01/01/2006	01/31/2014						
58016-0291-60		J8540		01/01/2007	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.5 MG	60	EA	BO	PO	EA	0.25	MG	2	01/01/2007	01/31/2014						
58016-0293-00		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	100	EA	BO	PO	EA	0.25	MG	3	01/01/2006	01/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
58016-0293-06		J8540		01/01/2007	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	6 EA	NA	PO	EA		0.25 MG		3	01/01/2007	01/31/2014						
58016-0293-12		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	12 EA	BO	PO	EA		0.25 MG		3	01/01/2006	01/31/2014						
58016-0293-15		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	15 EA	BO	PO	EA		0.25 MG		3	01/01/2006	01/31/2014						
58016-0293-20		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	20 EA	BO	PO	EA		0.25 MG		3	01/01/2006	01/31/2014						
58016-0293-30		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	30 EA	BO	PO	EA		0.25 MG		3	01/01/2006	01/31/2014						
58016-0326-00		Q0164		03/01/2007	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	100 EA	BO	PO	EA		5 MG		1	03/01/2007	01/31/2014						
58016-0326-12		Q0164		09/15/2003	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	12 EA	BO	PO	EA		5 MG		1	09/15/2003	01/31/2014						
58016-0326-30		Q0164		03/01/2007	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	30 EA	BO	PO	EA		5 MG		1	03/01/2007	01/31/2014						
58016-0326-60		Q0164		03/01/2007	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	60 EA	BO	PO	EA		5 MG		1	03/01/2007	01/31/2014						
58016-0326-90		Q0164		03/01/2007	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	90 EA	BO	PO	EA		5 MG		1	03/01/2007	01/31/2014						
58016-0391-00		Q0144		01/15/2004	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	100 EA	BO	PO	EA		1 GM	0.25	01/15/2004	01/31/2014							
58016-0391-01		Q0144		04/03/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX Z-PAK 250 MG	6 EA	BX	PO	EA		1 GM	0.25	04/03/2002	01/31/2014							
58016-0391-06		Q0144		01/01/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6 EA	BO	PO	EA		1 GM	0.25	01/01/2002	01/31/2014							
58016-0391-10		Q0144		01/01/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	10 EA	BO	PO	EA		1 GM	0.25	01/01/2002	01/31/2014							
58016-0391-15		Q0144		01/01/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	15 EA	BO	PO	EA		1 GM	0.25	01/01/2002	01/31/2014							
58016-0391-18		Q0144		01/01/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	18 EA	BX	PO	EA		1 GM	0.25	01/01/2002	01/31/2014							
58016-0391-20		Q0144		01/01/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	20 EA	BO	PO	EA		1 GM	0.25	01/01/2002	01/31/2014							
58016-0391-28		Q0144		01/01/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	28 EA	BO	PO	EA		1 GM	0.25	01/01/2002	01/31/2014							
58016-0391-30		Q0144		01/01/2002	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	30 EA	BO	PO	EA		1 GM	0.25	01/01/2002	01/31/2014							
58016-0391-60		Q0144		01/15/2004	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	60 EA	BO	PO	EA		1 GM	0.25	01/15/2004	01/31/2014							
58016-0391-90		Q0144		01/15/2004	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	90 EA	BO	PO	EA		1 GM	0.25	01/15/2004	01/31/2014							
58016-0408-00		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-06		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	6 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-09		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	9 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-10		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	10 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-12		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	12 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-14		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	14 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-15		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-20		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						
58016-0408-21		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	21 EA	BO	PO	EA		50 MG		0.5	01/01/2002	01/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
58016-0408-24		Q0163		01/01/2007	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	24	EA	NA	PO	EA	50	MG	0.5	01/01/2007	01/31/2014						
58016-0408-25		Q0163		01/01/2007	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	25	EA	NA	PO	EA	50	MG	0.5	01/01/2007	01/31/2014						
58016-0408-28		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	28	EA	BO	PO	EA	50	MG	0.5	01/01/2002	01/31/2014						
58016-0408-30		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30	EA	BO	PO	EA	50	MG	0.5	01/01/2002	01/31/2014						
58016-0408-40		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	40	EA	BO	PO	EA	50	MG	0.5	01/01/2002	01/31/2014						
58016-0408-60		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60	EA	BO	PO	EA	50	MG	0.5	01/01/2002	01/31/2014						
58016-0409-00		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	01/31/2014						
58016-0409-10		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	10	EA	BO	PO	EA	50	MG	1	01/01/2002	01/31/2014						
58016-0409-12		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	12	EA	BO	PO	EA	50	MG	1	01/01/2002	01/31/2014						
58016-0409-15		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15	EA	BO	PO	EA	50	MG	1	01/01/2002	01/31/2014						
58016-0409-20		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20	EA	BO	PO	EA	50	MG	1	01/01/2002	01/31/2014						
58016-0409-21		Q0163		01/01/2007	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	21	EA	BO	PO	EA	50	MG	1	01/01/2007	01/31/2014						
58016-0409-24		Q0163		03/26/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	24	EA	BO	PO	EA	50	MG	1	03/26/2002	01/31/2014						
58016-0409-30		Q0163		01/01/2002	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30	EA	BO	PO	EA	50	MG	1	01/01/2002	01/31/2014						
58016-0409-40		Q0163		01/01/2007	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	40	EA	NA	PO	EA	50	MG	1	01/01/2007	01/31/2014						
58016-0409-60		Q0163		08/01/2006	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	60	EA	BO	PO	EA	50	MG	1	08/01/2006	01/31/2014						
58016-0409-90		Q0163		08/01/2006	01/31/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	90	EA	BO	PO	EA	50	MG	1	08/01/2006	01/31/2014						
58016-0424-00		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
58016-0424-02		Q0170		09/15/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	120	EA	BO	PO	EA	25	MG	1	09/15/2003	12/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
58016-0424-03		Q0170		09/15/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	150	EA	BO	PO	EA	25	MG	1	09/15/2003	12/31/2013						
58016-0424-10		Q0170		03/26/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	25	MG	1	03/26/2002	12/31/2013						
58016-0424-12		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
58016-0424-15		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
58016-0424-20		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
58016-0424-30		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
58016-0424-40		Q0170		01/01/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	40	EA	NA	PO	EA	25	MG	1	01/01/2007	12/31/2013						
58016-0424-48		Q0170		01/01/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	48	EA	NA	PO	EA	25	MG	1	01/01/2007	12/31/2013						
58016-0424-50		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	50	EA	BO	PO	EA	25	MG	1	01/01/2002	12/31/2013						
58016-0424-60		Q0170		07/13/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	BO	PO	EA	25	MG	1	07/13/2003	12/31/2013						
58016-0424-73		Q0170		09/15/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	300	EA	BO	PO	EA	25	MG	1	09/15/2003	12/31/2013						
58016-0424-89		Q0170		09/15/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	200	EA	BO	PO	EA	25	MG	1	09/15/2003	12/31/2013						
58016-0424-90		Q0170		09/15/2003	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	90	EA	BO	PO	EA	25	MG	1	09/15/2003	12/31/2013						
58016-0464-10		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	10	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013						
58016-0464-15		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	15	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013						
58016-0464-20		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013						
58016-0464-30		Q0178		01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30	EA	BO	PO	EA	50	MG	1	01/01/2002	12/31/2013						
58016-0603-01	A4216			01/01/2006	01/31/2014	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE 0.9%	3	ML	EA	IH	ML	10	ML	0.1	01/01/2006	01/31/2014						
58016-0627-00	J8499			01/29/2002	01/31/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100	EA	BO	PO	EA	1	EA	1	01/29/2002	01/31/2014						
58016-0627-20	J8499			01/29/2002	01/31/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	20	EA	BO	PO	EA	1	EA	1	01/29/2002	01/31/2014						
58016-0627-30	J8499			01/29/2002	01/31/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	01/29/2002	01/31/2014						
58016-0627-60	J8499			01/29/2002	01/31/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	60	EA	BO	PO	EA	1	EA	1	01/29/2002	01/31/2014						
58016-0627-90	J8499			01/29/2002	01/31/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	90	EA	BO	PO	EA	1	EA	1	01/29/2002	01/31/2014						
58016-0673-12	J7510			01/01/2002	01/31/2014	PREDNISOLONE ORAL, PER 5 MG	PRELONE 15 MG/5 ML	60	ML	EA	PO	ML	5	MG	0.6	01/01/2002	01/31/2014						
58016-0673-24	J7510			01/01/2002	01/31/2014	PREDNISOLONE ORAL, PER 5 MG	PRELONE 15 MG/5 ML	120	ML	EA	PO	ML	5	MG	0.6	01/01/2002	01/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
58016-0673-48		J7510		01/01/2002	01/31/2014	PREDNISOLONE ORAL, PER 5 MG	PRELONE 15 MG/5 ML	240	ML	EA	PO	ML	5	MG	0.6	01/01/2002	01/31/2014						
58016-0706-00		Q0165		09/23/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	10	MG	1	09/23/2004	12/31/2013						
58016-0706-02		Q0165		09/23/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	120	EA	BO	PO	EA	10	MG	1	09/23/2004	12/31/2013						
58016-0706-03		Q0165		09/23/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	150	EA	BO	PO	EA	10	MG	1	09/23/2004	12/31/2013						
58016-0706-08		Q0165		01/01/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	8	EA	NA	PO	EA	10	MG	1	01/01/2007	12/31/2013						
58016-0706-30		Q0165		09/23/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	10	MG	1	09/23/2004	12/31/2013						
58016-0706-60		Q0165		09/23/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	10	MG	1	09/23/2004	12/31/2013						
58016-0706-90		Q0165		09/23/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	90	EA	BO	PO	EA	10	MG	1	09/23/2004	12/31/2013						
58016-0781-00		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	100	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-08		J8540		01/01/2007	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	8	EA	NA	PO	EA	0.25	MG	16	01/01/2007	01/31/2014						
58016-0781-10		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	10	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-12		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	12	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-14		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	14	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-15		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	15	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-20		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	20	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-21		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	21	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-24		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	24	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-28		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	28	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-30		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	30	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-40		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	40	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0781-50		J8540		01/01/2006	01/31/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	50	EA	BO	PO	EA	0.25	MG	16	01/01/2006	01/31/2014						
58016-0951-00		Q0168		04/01/2004	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	100	EA	BO	PO	EA	5	MG	1	04/01/2004	12/31/2013						
58016-0951-30		Q0168		04/01/2004	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	30	EA	BO	PO	EA	5	MG	1	04/01/2004	12/31/2013						
58016-0951-60		Q0168		04/01/2004	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	60	EA	BO	PO	EA	5	MG	1	04/01/2004	12/31/2013						
58016-0951-90		Q0168		04/01/2004	12/31/2013	DRONABINOL, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	90	EA	BO	PO	EA	5	MG	1	04/01/2004	12/31/2013						
58016-0973-00		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	100	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014						
58016-0973-02		Q0173		09/15/2003	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	120	EA	BO	PO	EA	250	MG	1	09/15/2003	01/31/2014						
58016-0973-03		Q0173		09/15/2003	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	150	EA	BO	PO	EA	250	MG	1	09/15/2003	01/31/2014						
58016-0973-08		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	8	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
58016-0973-10		Q0173		09/15/2003	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	10	EA	BO	PO	EA	250	MG	1	09/15/2003	01/31/2014							
58016-0973-12		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	12	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014							
58016-0973-15		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	15	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014							
58016-0973-20		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	20	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014							
58016-0973-24		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	24	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014							
58016-0973-30		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	30	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014							
58016-0973-50		Q0173		01/01/2002	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	50	EA	BO	PO	EA	250	MG	1	01/01/2002	01/31/2014							
58016-0973-60		Q0173		09/15/2003	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	60	EA	BO	PO	EA	250	MG	1	09/15/2003	01/31/2014							
58016-0973-73		Q0173		09/15/2003	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	300	EA	BO	PO	EA	250	MG	1	09/15/2003	01/31/2014							
58016-0973-89		Q0173		09/15/2003	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	200	EA	BO	PO	EA	250	MG	1	09/15/2003	01/31/2014							
58016-0973-90		Q0173		09/15/2003	01/31/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	90	EA	BO	PO	EA	250	MG	1	09/15/2003	01/31/2014							
58016-2001-01		J7509		10/01/2006	01/31/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21	EA	DP	PO	EA	4	MG	1	10/01/2006	01/31/2014							
58016-2004-01		J7509		01/01/2002	01/31/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (DOSE PACK) 4 MG	21	EA	DP	PO	EA	4	MG	1	01/01/2002	01/31/2014							
58016-3018-03		J8498		01/01/2006	01/31/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	COMPazine 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	01/31/2014							
58016-3066-01		J8498		01/01/2006	01/31/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 12.5 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	01/31/2014							
58016-3067-01		J8498		01/01/2006	01/31/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PHENERGAN 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	01/31/2014							
58016-3222-01		J8498		01/01/2006	01/31/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	COMPazine 5 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	01/31/2014							
58016-4008-01		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120	ML	NA	PO	ML	25	MG	0.05	01/01/2002	12/31/2013							
58016-4144-01		J7510		01/01/2002	01/31/2014	PREDNISOLONE ORAL, PER 5 MG	PEDIAPRED 5 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.2	01/01/2002	01/31/2014							
58016-4719-01		J7509		02/16/2005	01/31/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 8 MG	25	EA	BO	PO	EA	4	MG	2	02/16/2005	01/31/2014							
58016-4770-01		J2300		02/01/2006	01/31/2014	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL (10X1ML AMP) 10 MG/ML	1	ML	AM	IJ	ML	10	MG	1	02/01/2006	01/31/2014							
58016-4771-01		J2941		02/01/2006	01/31/2014	INJECTION, SOMATROPIN, 1 MG	GENOTROPIN 13.8 MG	1	EA	CT	SC	EA	1	MG	13.8	02/01/2006	01/31/2014							
58016-4786-01		J0696		02/01/2006	01/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 1 G	1	EA	VL	IJ	EA	250	MG	4	02/01/2006	01/31/2014							
58016-4788-01		J1815		02/01/2006	01/31/2014	INJECTION, INSULIN, PER 5 UNITS	HUMULIN N 100 U/ML	10	ML	VL	SC	ML	5	U	20	02/01/2006	01/31/2014							
58016-4790-01		J0696		02/01/2006	01/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 250 MG	1	EA	VL	IJ	EA	250	MG	1	02/01/2006	01/31/2014							
58016-4811-01		J2765		02/01/2006	01/31/2014	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	REGLAN (25X2ML) 5 MG/ML	2	ML	VL	IV	ML	10	MG	0.5	02/01/2006	01/31/2014							
58016-4814-01		Q0144		12/20/2005	01/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6	EA	DP	PO	EA	1	MG	0.25	12/20/2005	01/31/2014							
58016-4832-01		J7506		02/01/2006	01/31/2014	PREDNISOLONE, ORAL, PER 5MG	PREDNISOLONE 5 MG	21	EA	DP	PO	EA	5	MG	1	02/01/2006	01/31/2014							
58016-4834-01		J0696		02/01/2006	01/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 2 G	1	EA	VL	IJ	EA	250	MG	8	02/01/2006	01/31/2014							
58016-4838-01		A4216		02/01/2006	01/31/2014	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	BRONCHO SALINE 0.9% AEROSOL 0.9%	240	ML	BO	IH	ML	10	ML	0.1	02/01/2006	01/31/2014							
58016-4840-01		J2001		02/01/2006	01/31/2014	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE (SDA) 1%	5	ML	AM	EP	ML	10	MG	1	02/01/2006	01/31/2014							
58016-4843-01		J7510		02/01/2006	01/31/2014	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	240	ML	BO	PO	ML	5	MG	0.6	02/01/2006	01/31/2014							
58016-4849-01		J7644		02/01/2006	01/31/2014	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM (2.5MLX25) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	02/01/2006	01/31/2014							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
58016-4849-01	KO	J7644	KO	02/01/2006	01/31/2014	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM (2.5MLX25) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	02/01/2006	01/31/2014							
58016-4855-01		J3303		02/01/2006	01/31/2014	INJECTION, TRIAMCINOLONE HEXACETONIDE, PER 5MG	ARISTOSPAN 20 MG/ML	5	ML	VL	IJ	ML	5	MG	4	02/01/2006	01/31/2014							
58016-4868-01		J0595		03/15/2006	01/31/2014	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (10X1ML) 2 MG/ML	1	ML	VL	IJ	ML	1	MG	2	03/15/2006	01/31/2014							
58016-4872-01		J1650		04/01/2006	01/31/2014	INJECTION, ENOXAPARIN SODIUM, 10 MG	LOVENOX 40 MG/0.4 ML	0.4	ML	SR	SC	ML	10	MG	10	04/01/2006	01/31/2014							
58016-4893-01		J1040		06/01/2006	01/31/2014	INJECTION, METHYLPREDNISOLONE ACETATE, 80 MG	METHYLPREDNISOLONE ACETATE 80 MG/ML	1	ML	VL	IJ	ML	80	MG	1	06/01/2006	01/31/2014							
58016-4897-01		J2920		07/01/2006	01/31/2014	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 40 MG	SOLU-MEDROL (SDV) 40 MG	1	EA	VL	IJ	EA	40	MG	1	07/01/2006	01/31/2014							
58016-4995-01		A4216		01/01/2007	01/31/2014	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (10MLX100) 0.9%	10	ML	SR	IJ	ML	10	ML	0.1	01/01/2007	01/31/2014							
58016-5009-01		J8498		01/01/2006	01/31/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROMETHAZINE HCL 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	01/31/2014							
58016-6404-01		J7611		04/01/2008	01/31/2014	COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG	ALBUTEROL SULFATE 0.5%	20	ML	NA	IH	ML	1	MG	5	04/01/2008	01/31/2014							
58016-6506-01		J8498		01/01/2006	01/31/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12	EA	BX	RC	EA	1	EA	1	01/01/2006	01/31/2014							
58016-9191-01		J0702		01/01/2002	01/31/2014	INJECTION, BETAMETHASONE ACETATE 3MG AND BETAMETHASONE SODIUM PHOSPHATE 3MG	CELESTONE SOLUSPAN (M.D.V.) 3 MG/ML	5	ML	VL	IJ	ML	3	MG	1	01/01/2002	01/31/2014							
58016-9299-01		J3410		01/01/2002	01/31/2014	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL 50 MG/ML	10	ML	VL	IM	ML	25	MG	2	01/01/2002	01/31/2014							
58016-9331-01		J2001		08/01/2004	01/31/2014	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (M.D.V.) 1%	50	ML	VL	EP	ML	10	MG	1	08/01/2004	01/31/2014							
58016-9343-01		J3490		01/01/2002	01/31/2014	UNCLASSIFIED DRUGS	MARCAINE HCL (M.D.V.) 0.5%	50	ML	VL	IJ	ML	1	EA	1	01/01/2002	01/31/2014							
58016-9384-01		J2300		01/01/2002	01/31/2014	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL (M.D.V.) 10 MG/ML	10	ML	VL	IJ	ML	10	MG	1	01/01/2002	01/31/2014							
58016-9413-01		J1885		01/01/2002	01/31/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV) 30 MG/ML	2	ML	VL	IM	ML	15	MG	2	01/01/2002	01/31/2014							
58016-9438-01		J0696		02/22/2002	01/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	ROCEPHIN 1 GM	1	EA	VL	IJ	EA	250	MG	4	02/22/2002	01/31/2014							
58016-9452-01		J2930		01/01/2002	01/31/2014	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	SOLU-MEDROL 125 MG	1	EA	VL	IJ	EA	125	MG	1	01/01/2002	01/31/2014							
58016-9453-01		J0696		01/01/2002	01/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	ROCEPHIN 250 MG	1	EA	VL	IJ	EA	250	MG	1	01/01/2002	01/31/2014							
58016-9464-01		A4216		01/01/2004	01/31/2014	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION	50	ML	VL	IJ	ML	10	ML	0.1	01/01/2004	01/31/2014							
58016-9551-01		J0696		01/01/2002	01/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	ROCEPHIN 500 MG	1	EA	VL	IJ	EA	250	MG	2	01/01/2002	01/31/2014							
58160-0815-11		J3490		08/06/2007	08/07/2017	UNCLASSIFIED DRUGS	TWINRIX (TAX INCLUDED,1MLX10,PF) 720 EL U/ML-20 MCG/ML	1	ML	VL	IM	ML	1	EA	1	08/06/2007	08/07/2017							
58160-0820-11		J3490		02/01/2007	10/03/2017	UNCLASSIFIED DRUGS	ENGERIX-B PEDIATRIC (10X0.5ML,SDV,TAXINCL,PF) 10 MCG/0.5 ML	0.5	ML	VL	IM	ML	1	EA	1	02/01/2007	10/03/2017							
58160-0821-11		J3490		02/01/2007	99/99/9999	UNCLASSIFIED DRUGS	ENGERIX-B (SDV,TAXINCL,PF) 20 MCG/ML	1	ML	VL	IM	ML	1	EA	1	02/01/2007	99/99/9999							
58160-0856-35		J3490		01/01/2002	02/03/2016	UNCLASSIFIED DRUGS	ENGERIX-B PEDIATRIC (TIPLOK,23GX1,TAX INC,PF) 10 MCG/0.5 ML	0.5	ML	SR	IM	ML	1	EA	1	01/01/2002	02/03/2016							
58281-0560-01		J0475		01/01/2002	01/24/2018	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT (1X20 ML AMP) 0.5 MG/ML	20	ML	BX	IN	EA	10	MG	1	01/01/2002	01/24/2018							
58281-0560-02		J0475		04/02/2004	01/24/2018	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT (2X20ML AMP) 0.5 MG/ML	20	ML	BX	MR	EA	10	MG	2	04/02/2004	01/24/2018							
58281-0561-02		J0475		01/01/2002	01/24/2018	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT (2X5 ML AMP) 2 MG/ML	5	ML	BX	IN	EA	10	MG	2	01/01/2002	01/24/2018							
58281-0562-01		J0476		01/01/2002	07/10/2017	INJECTION, BACLOFEN, 50 MCG FOR INTRATHECAL TRIAL	LIORESAL INTRATHECAL SCREENING KIT (1X1 ML AMP) 0.05 MG/ML	1	ML	AM	IN	EA	50	MCG	1	01/01/2002	07/10/2017							
58281-0563-01		J0475		10/21/2003	07/23/2017	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT (1X20ML AMP) 2 MG/ML	20	ML	BX	MR	EA	10	MG	4	10/21/2003	07/23/2017							
58281-0563-02		J0475		04/02/2004	07/23/2017	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT (2X20ML AMP) 2 MG/ML	20	ML	BX	MR	EA	10	MG	8	04/02/2004	07/23/2017							
58406-0425-34		J1438		01/01/2002	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (S.D. TRAY,PF) 25 MG	4	EA	BX	SC	EA	25	MG	1	01/01/2002	99/99/9999							
58406-0425-41		J1438		01/01/2002	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (S.D. TRAY,PF) 25 MG	1	EA	BX	SC	EA	25	MG	1	01/01/2002	99/99/9999							
58406-0435-01		J1438		11/17/2004	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (ACTUAL FILL 50MG/0.98ML) 50 MG/ML	0.98	ML	SR	SC	ML	25	MG	2	11/17/2004	99/99/9999							
58406-0435-04		J1438		11/17/2004	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (ACTUAL FILL 50MG/0.98ML) 50 MG/ML	0.98	ML	SR	SC	ML	25	MG	2	11/17/2004	99/99/9999							
58406-0445-01		J1438		07/17/2006	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (SURECLICK AUTOINJECTOR) 50 MG/ML	0.98	ML	SR	SC	ML	25	MG	2	07/17/2006	99/99/9999							
58406-0445-04		J1438		07/17/2006	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (SURECLICK AUTOINJECTOR) 50 MG/ML	0.98	ML	SR	SC	ML	25	MG	2	07/17/2006	99/99/9999							
58406-0455-01		J1438		04/30/2007	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (27G,1/2",PF) 50 MG/ML	0.51	ML	SR	SC	ML	25	MG	2	04/30/2007	99/99/9999							
58406-0455-04		J1438		04/30/2007	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (4X0.51ML,27G,1/2",PF) 50 MG/ML	0.51	ML	SR	SC	ML	25	MG	2	04/30/2007	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
58468-0040-01		J0180		01/01/2005	99/99/9999	INJECTION, AGALSIDASE BETA, 1 MG	FABRAZYME (PF) 35 MG	1 EA	VL	IV	EA		1 MG		35	01/01/2005	99/99/9999						
58468-0041-01		J0180		01/01/2005	99/99/9999	INJECTION, AGALSIDASE BETA, 1 MG	FABRAZYME (PF) 5 MG	1 EA	VL	IV	EA		1 MG		5	01/01/2005	99/99/9999						
58468-0070-01		J1931		01/01/2005	99/99/9999	INJECTION, LARONIDASE, 0.1 MG LYMPHOCYTE IMMUNE GLOBULIN, ANTITHYMOCYTE GLOBULIN, RABBIT, PARENTERAL, 25MG	ALDURAZYME (PF) 0.58 MG/ML	5 ML	VL	IV	ML		0.1 MG		5.8	01/01/2005	99/99/9999						
58468-0080-01		J7511		12/01/2005	99/99/9999		THYMOGLOBULIN (VIAL,DILUENT) 25 MG	1 EA	VL	IV	EA		25 MG		1	12/01/2005	99/99/9999						
58468-0100-01		J9027		01/01/2006	12/14/2014	INJECTION, CLOFARABINE, 1 MG	CLOLAR (SINGLE-USE VIAL,PF) 1 MG/ML	20 ML	VL	IV	ML		1 MG		1	01/01/2006	12/14/2014						
58468-0218-02		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	120 EA	NA	PO	EA		0.25 MG		16	01/01/2006	99/99/9999						
58468-1849-04		J3240		01/01/2002	05/31/2016	INJECTION, THYROTROPIN ALPHA, 0.9 MG, PROVIDED IN 1.1 MG VIAL	THYROGEN (W/2 VIALS DILUENT) 1.1 MG	1 EA	VL	IJ	EA		1.1 MG		1	01/01/2002	05/31/2016						
58864-0162-30		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (REDI-SCRIPT) 25 MG	30 EA	BO	PO	EA		50 MG		0.5	01/01/2002	99/99/9999						
58864-0162-56		Q0163		03/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL (REDI-SCRIPT) 25 MG	56 EA	BO	PO	EA		50 MG		0.5	03/01/2004	99/99/9999						
58864-0191-25		J8499		03/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (REDI-SCRIPT) 800 MG	25 EA	BO	PO	EA		1 EA		1	03/01/2004	99/99/9999						
58864-0191-35		J8499		03/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (REDI-SCRIPT) 800 MG	35 EA	BO	PO	EA		1 EA		1	03/01/2004	99/99/9999						
58864-0362-20		J7506		03/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,REDI-SCRIPT) 5 MG	20 EA	BO	PO	EA		5 MG		1	03/01/2004	12/31/2015						
58864-0362-56		J7506		03/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,REDI-SCRIPT) 5 MG	56 EA	BO	PO	EA		5 MG		1	03/01/2004	12/31/2015						
58864-0423-15		J7506		01/01/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	15 EA	BO	PO	EA		5 MG		2	01/01/2005	12/31/2015						
58864-0423-20		J7506		06/01/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	06/01/2005	12/31/2015						
58864-0423-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (REDI-SCRIPT) 10 MG	30 EA	BO	PO	EA		5 MG		2	01/01/2002	12/31/2015						
58864-0423-40		J7506		07/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (REDI-SCRIPT) 10 MG	40 EA	BO	PO	EA		5 MG		2	07/01/2004	12/31/2015						
58864-0424-14		J7506		03/02/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (REDI-SCRIPT) 20 MG	14 EA	BO	PO	EA		5 MG		4	03/02/2004	12/31/2015						
58864-0424-20		J7506		01/01/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (REDI-SCRIPT) 20 MG	20 EA	BO	PO	EA		5 MG		4	01/01/2005	12/31/2015						
58864-0424-30		J7506		03/02/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30 EA	BO	PO	EA		5 MG		4	03/02/2004	12/31/2015						
58864-0602-01		J8499		06/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (REDI-SCRIPT) 400 MG	100 EA	BO	PO	EA		1 EA		1	06/01/2004	99/99/9999						
58864-0602-30		J8499		03/02/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (REDI-SCRIPT) 400 MG	30 EA	BO	PO	EA		1 EA		1	03/02/2004	99/99/9999						
58864-0644-42		Q0165		03/01/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (REDI-SCRIPT) 10 MG	42 EA	BO	PO	EA		10 MG		1	03/01/2004	12/31/2013						
58864-0655-04		Q0144		07/01/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	4 EA	BO	PO	EA		1 GM		0.25	07/01/2005	99/99/9999						
58864-0655-06		Q0144		09/10/2003	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX (REDI-SCRIPT) 250 MG	6 EA	BO	PO	EA		1 GM		0.25	09/10/2003	99/99/9999						
58864-0655-14		Q0144		02/01/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	14 EA	BO	PO	EA		1 GM		0.25	02/01/2005	99/99/9999						
58864-0655-30		Q0144		06/01/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6 EA	BO	PO	EA		1 GM		0.25	06/01/2006	99/99/9999						
58864-0702-01		Q0164		06/15/2006	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	15 EA	BO	PO	EA		5 MG		1	06/15/2006	99/99/9999						
58864-0761-10		Q0170		08/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (REDI-SCRIPT) 25 MG	10 EA	BO	PO	EA		25 MG		1	08/01/2004	12/31/2013						
58864-0761-30		Q0170		05/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30 EA	BO	PO	EA		25 MG		1	05/01/2004	12/31/2013						
58864-0761-42		Q0170		08/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	42 EA	BO	PO	EA		25 MG		1	08/01/2004	12/31/2013						
58864-0791-06		Q0144		07/01/2004	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN DIHYDRATE 250 MG	6 EA	BO	PO	EA		1 GM		0.25	07/01/2004	99/99/9999						
58864-0876-35		J8499		01/01/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	35 EA	BO	PO	EA		1 EA		1	01/01/2005	99/99/9999						
58914-0080-52		J0500		06/22/2004	99/99/9999	INJECTION, DICYCLIMINE HCL, UP TO 20 MG	BENTYL (AMP) 10 MG/ML	2 ML	AM	IM	ML		20 MG		0.5	03/23/2007	99/99/9999	06/22/2004	11/14/2004	0.5			
59075-0730-15		J2323		01/01/2008	04/01/2014	INJECTION, NATALIZUMAB, 1 MG	TYABRI 20 MG/ML	15 ML	VL	IV	ML		1 MG		20	01/01/2008	04/01/2014						
59148-0016-65		J0400		01/01/2008	06/15/2015	INJECTION, ARIPIPRAZOLE, INTRAMUSCULAR, 0.25 MG	ABILIFY (SDV) 9.75 MG/1.3 ML	1.3 ML	VL	IM	ML		0.25 MG		30	01/01/2008	06/15/2015						
59618-0199-33		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENYL ELIXIR 12.5 MG/5 ML	120 ML	EA	PO	ML		50 MG		0.05	01/01/2002	02/03/2016						
59618-0200-06		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENYL 25 MG	24 EA	NA	PO	EA		50 MG		0.5	01/01/2002	02/03/2016						
59676-0302-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VIAL) 2000 U/ML	1 ML	VL	IJ	ML		1000 U		2	01/01/2006	99/99/9999						
61314-0326-10		Q5101		07/20/2018	99/99/9999	INJECTION, FILGRASTIM-SNDZ, BIOSIMILAR, (ZARXIO), 1 MICROGRAM	ZARXIO (PF) 480 MCG/0.8 ML	0.8 ML	SR	IJ	ML		1 MCG		600	07/20/2018	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
59676-0302-02		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VOLUME PACK VIAL) 2000 U/ML	1 ML	VL	IJ	ML		1000 U		2	01/01/2006	99/99/9999						
59676-0303-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VIAL) 3000 U/ML	1 ML	VL	IJ	ML		1000 U		3	01/01/2006	99/99/9999						
59676-0303-02		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VOLUME PACK VIAL) 3000 U/ML	1 ML	VL	IJ	ML		1000 U		3	01/01/2006	99/99/9999						
59676-0304-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VIAL) 4000 U/ML	1 ML	VL	IJ	ML		1000 U		4	01/01/2006	99/99/9999						
59676-0304-02		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VOLUME PACK VIAL) 4000 U/ML	1 ML	VL	IJ	ML		1000 U		4	01/01/2006	99/99/9999						
59676-0310-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VIAL) 10000 U/ML	1 ML	VL	IJ	ML		1000 U		10	01/01/2006	99/99/9999						
59676-0310-02		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (VOLUME PACK VIAL) 10000 U/ML	1 ML	VL	IJ	ML		1000 U		10	01/01/2006	99/99/9999						
59676-0312-04		J0885		01/18/2008	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (6X2ML MDV) 10000 U/ML	2 ML	VL	IJ	ML		1000 U		10	01/18/2008	99/99/9999						
59676-0320-04		J0886		10/15/2007	12/31/2015	INJECTION, EPOETIN ALFA, 1000 UNITS, (FOR ESRD ON DIALYSIS)	PROCRIT (MULTIDOSE) 20000 U/ML	1 ML	VL	IJ	ML		1000 U		20	10/15/2007	12/31/2015						
59676-0340-01		J0885		01/01/2006	99/99/9999	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (PF) 40000 U/ML	1 ML	VL	IJ	ML		1000 U		40	01/01/2006	99/99/9999						
59741-0119-04		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	120 ML	BO	PO	ML		50 MG		0.05	01/01/2002	02/03/2016						
59741-0119-08		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	240 ML	BO	PO	ML		50 MG		0.05	01/01/2002	02/03/2016						
59741-0119-16		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	480 ML	BO	PO	ML		50 MG		0.05	01/01/2002	02/03/2016						
59741-0119-20		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	3840 ML	BO	PO	ML		50 MG		0.05	01/01/2002	02/03/2016						
59746-0001-03		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21 EA	DP	PO	EA		4 MG		1	01/01/2002	99/99/9999						
59746-0001-06		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	100 EA	BO	PO	EA		4 MG		1	01/01/2002	99/99/9999						
59746-0002-04		J7509		09/24/2007	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (USP) 8 MG	25 EA	BO	PO	EA		4 MG		2	09/24/2007	99/99/9999						
59746-0003-14		J7509		07/20/2007	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (USP) 16 MG	50 EA	BO	PO	EA		4 MG		4	07/20/2007	99/99/9999						
59746-0007-06		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	100 EA	NA	PO	EA		5 MG		1	01/01/2002	12/31/2015						
59746-0007-10		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	1000 EA	NA	PO	EA		5 MG		1	01/01/2002	12/31/2015						
59746-0008-06		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	100 EA	NA	PO	EA		5 MG		2	01/01/2002	12/31/2015						
59746-0008-10		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	1000 EA	NA	PO	EA		5 MG		2	01/01/2002	12/31/2015						
59746-0015-04		J7509		07/20/2007	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (USP) 32 MG	25 EA	BO	PO	EA		4 MG		8	07/20/2007	99/99/9999						
59746-0113-06		Q0164		01/01/2002	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	100 EA	BO	PO	EA		5 MG		1	01/01/2002	99/99/9999						
59746-0115-06		Q0165		01/01/2002	12/31/2015	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100 EA	BO	PO	EA		10 MG		1	01/01/2002	12/31/2015						
59746-0171-06		J7506		10/21/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 1 MG	100 EA	BO	PO	EA		5 MG		0.2	10/21/2005	12/31/2015						
59746-0171-10		J7506		10/21/2005	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 1 MG	1000 EA	BO	PO	EA		5 MG		0.2	10/21/2005	12/31/2015						
59746-0172-06		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 5 MG	100 EA	BO	PO	EA		5 MG		1	08/03/2007	12/31/2015						
59746-0172-10		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 5 MG	1000 EA	BO	PO	EA		5 MG		1	08/03/2007	12/31/2015						
59746-0173-06		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 10 MG	100 EA	BO	PO	EA		5 MG		2	08/03/2007	12/31/2015						
59746-0173-09		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 10 MG	500 EA	BO	PO	EA		5 MG		2	08/03/2007	12/31/2015						
59746-0173-10		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 10 MG	1000 EA	BO	PO	EA		5 MG		2	08/03/2007	12/31/2015						
59746-0175-06		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 20 MG	100 EA	BO	PO	EA		5 MG		4	08/03/2007	12/31/2015						
59746-0175-09		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 20 MG	500 EA	BO	PO	EA		5 MG		4	08/03/2007	12/31/2015						
59746-0175-10		J7506		08/03/2007	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 20 MG	1000 EA	BO	PO	EA		5 MG		4	08/03/2007	12/31/2015						
59762-0100-01		J8515		01/01/2006	99/99/9999	CABERGOLINE, ORAL, 0.25 MG	CABERGOLINE 0.5 MG	8 EA	BO	PO	EA		0.25 MG		2	01/01/2006	99/99/9999						
59762-2576-01		J9211		08/27/2007	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (PF) 1 MG/ML	5 ML	VL	IV	ML		5 MG		0.2	08/27/2007	99/99/9999						
59762-2586-01		J9211		08/27/2007	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (PF) 1 MG/ML	10 ML	VL	IV	ML		5 MG		0.2	08/27/2007	99/99/9999						
59762-2596-01		J9211		08/27/2007	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (PF) 1 MG/ML	20 ML	VL	IV	ML		5 MG		0.2	08/27/2007	99/99/9999						
59762-3051-01		Q0144		07/07/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 1 GM/Packet	10 EA	BX	PO	EA		1 GM		1	07/07/2006	99/99/9999						
59762-3051-02		Q0144		07/07/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 1 GM/Packet	3 EA	BX	PO	EA		1 GM		1	07/07/2006	99/99/9999						
59762-3060-01		Q0144		11/14/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	6 EA	DP	PO	EA		1 GM		0.25	11/14/2005	99/99/9999						
59762-3060-02		Q0144		11/14/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	30 EA	BO	PO	EA		1 GM		0.25	11/14/2005	99/99/9999						
59762-3060-03		Q0144		11/14/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 250 MG	50 EA	BX	PO	EA		1 GM		0.25	11/14/2005	99/99/9999						
59762-3070-01		Q0144		11/14/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	3 EA	DP	PO	EA		1 GM		0.5	11/14/2005	99/99/9999						
59762-3070-02		Q0144		11/14/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	30 EA	BO	PO	EA		1 GM		0.5	11/14/2005	99/99/9999						
59762-3080-01		Q0144		11/14/2005	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 600 MG	30 EA	BO	PO	EA		1 GM		0.6	11/14/2005	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
59762-3110-01		Q0144		07/07/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY) 100 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.02	07/07/2006	99/99/9999						
59762-3120-01		Q0144		07/07/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY) 200 MG/5 ML	15 ML	BO	PO	ML		1 GM		0.04	07/07/2006	99/99/9999						
59762-3130-01		Q0144		07/07/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY) 200 MG/5 ML	22.5 ML	BO	PO	ML		1 GM		0.04	07/07/2006	99/99/9999						
59762-3140-01		Q0144		07/07/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY) 200 MG/5 ML	30 ML	BO	PO	ML		1 GM		0.04	07/07/2006	99/99/9999						
59762-4537-01		J1055		09/27/2004	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	MEDROXYPROGESTERONE ACETATE 150 MG/ML	1 ML	VL	IM	ML		150 MG		1	09/27/2004	12/31/2012						
59762-4537-02		J1055		09/27/2004	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	MEDROXYPROGESTERONE ACETATE 150 MG/ML	1 ML	VL	IM	ML		150 MG		1	09/27/2004	12/31/2012						
59762-5091-01		J9178		08/08/2007	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (SINGLE USE, PF) 2 MG/ML	25 ML	VL	IV	ML		2 MG		1	08/08/2007	99/99/9999						
59762-5093-01		J9178		08/08/2007	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (SINGLE USE, PF) 2 MG/ML	100 ML	VL	IV	ML		2 MG		1	08/08/2007	99/99/9999						
59762-7529-01		J9206		02/27/2008	01/01/2013	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML SDV) 20 MG/ML	2 ML	VL	IV	ML		20 MG		1	02/27/2008	01/01/2013						
59762-7529-02		J9206		02/27/2008	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML, SDV) 20 MG/ML	5 ML	VL	IV	ML		20 MG		1	02/27/2008	99/99/9999						
60242-0202-01		Q0163		07/06/2007	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 50 MG	100 EA	BO	PO	EA		50 MG		1	07/06/2007	02/03/2016						
60242-0202-10		Q0163		07/06/2007	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 50 MG	1000 EA	BO	PO	EA		50 MG		1	07/06/2007	02/03/2016						
67457-0533-16		J9171		09/05/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (USP;MULTI-USE VIAL) 10 MG/1 ML	16 ML	VL	IV	ML		1 MG		10	09/05/2018	99/99/9999						
67457-0705-75		J3370		08/31/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (LYOPHILIZED) 750 MG	10 EA	VL	IV	EA		500 MG		1.5	08/31/2018	99/99/9999						
67457-0813-50		J0878		09/04/2018	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (SDV,PF,LYOPHILIZED) 500 MG	1 EA	VL	IV	EA		1 MG		500	09/04/2018	99/99/9999						
67457-0822-99		J3370		08/31/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (LYOPHILIZED) 250 MG	10 EA	VL	IV	EA		500 MG		0.5	08/31/2018	99/99/9999						
60432-0126-08		J8999		11/17/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (LEMON-LIME) 40 MG/ML	240 ML	BO	PO	ML		1 EA		1	11/17/2004	99/99/9999						
60432-0126-16		J8999		12/01/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (LEMON-LIME) 40 MG/ML	480 ML	BO	PO	ML		1 EA		1	12/01/2006	99/99/9999						
60432-0140-50		J7502		09/28/2004	02/01/2015	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE 100 MG/ML	50 ML	BO	PO	ML		100 MG		1	09/28/2004	02/01/2015						
60432-0212-08		J7510		10/25/2004	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (DYE-FREE, GRAPE) 15 MG/5 ML	237 ML	BO	PO	ML		5 MG		0.6	10/25/2004	99/99/9999						
60432-0466-08		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (RASPERRY) 0.5 MG/5 ML	240 ML	BO	PO	ML		0.25 MG		0.4	01/01/2006	99/99/9999						
67457-0853-50		J1120		09/13/2018	99/99/9999	INJECTION, ACETAZOLAMIDE SODIUM, UP TO 500 MG	ACETAZOLAMIDE (USP,PF,LATEX-FREE) 500 MG	1 EA	VL	IV	EA		500 MG		1	09/13/2018	99/99/9999						
60432-0608-04		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (FRUIT,TROPICAL) 6.25 MG/5 ML	120 ML	BO	PO	ML		25 MG		0.05	01/01/2002	12/31/2013						
60432-0608-16		Q0170		01/01/2002	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (FRUIT,TROPICAL) 6.25 MG/5 ML	480 ML	BO	PO	ML		25 MG		0.05	01/01/2002	12/31/2013						
60492-0051-01		J1571		01/01/2008	04/17/2013	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAMUSCULAR, 0.5 ML	HEPAGAM B (SDV,PF)	5 ML	VL	IM	ML		0.5 ML		2	01/01/2008	04/17/2013						
60492-0051-02		J1573		01/01/2008	04/17/2013	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAVENOUS, 0.5 ML	NOVAPLUS HEPAGAM B (>1560/5ML,PF)	5 ML	VL	IJ	ML		0.5 ML		2	01/01/2008	04/17/2013						
60492-0052-01		J1571		01/01/2008	04/17/2013	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAMUSCULAR, 0.5 ML	HEPAGAM B (SDV,PF)	1 ML	VL	IM	ML		0.5 ML		2	01/01/2008	04/17/2013						
60492-0052-02		J1573		01/01/2008	04/17/2013	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAVENOUS, 0.5 ML	NOVAPLUS HEPAGAM B (>312IU/ML,PF)	1 ML	VL	IJ	ML		0.5 ML		2	01/01/2008	04/17/2013						
60505-0042-06		J8499		03/01/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (USP) 200 MG	100 EA	BO	PO	EA		1 EA		1	03/01/2006	99/99/9999						
60505-0133-00		J7515		05/17/2002	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE 25 MG	30 EA	BO	PO	EA		25 MG		1	05/17/2002	99/99/9999						
60505-0134-00		J7502		05/17/2002	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE 100 MG	30 EA	BO	PO	EA		100 MG		1	05/17/2002	99/99/9999						
60505-0354-01		J7502		08/01/2005	01/31/2014	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (U.S.P.) 100 MG/ML	50 ML	BO	PO	ML		100 MG		1	08/01/2005	01/31/2014						
60505-0368-01		J8999		06/23/2006	01/31/2014	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (USP,LEMON-LIME) 40 MG/ML	240 ML	BO	PO	ML		1 EA		1	06/23/2006	01/31/2014						
60505-0679-05		J0696		09/01/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (1X100ML,BULK PKG) 10 GM	1 EA	VL	IV	EA		250 MG		40	09/01/2005	99/99/9999						
60505-0679-08		J0696		09/01/2005	04/17/2013	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (1X100ML,PIGGYBACK) 1 GM	1 EA	VL	IJ	EA		250 MG		4	09/01/2005	04/17/2013						
60505-0679-09		J0696		09/01/2005	04/17/2013	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (1X100ML) 2 GM	1 EA	VL	IJ	EA		250 MG		8	09/01/2005	04/17/2013						
60505-0681-00		J0692		06/19/2007	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP) 2 GM	1 EA	VL	IJ	EA		500 MG		4	06/19/2007	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
60505-0702-01		J1631		01/01/2002	01/31/2014	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (M.D.V.) 50 MG/ML	5 ML	VL	IM	ML	ML	50 MG		1	01/01/2002	01/31/2014							
60505-0703-01		J1631		01/01/2002	01/31/2014	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (M.D.V.) 100 MG/ML	5 ML	VL	IM	ML	ML	50 MG		2	01/01/2002	01/31/2014							
60505-0705-00		J1885		02/28/2005	01/31/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV) 15 MG/ML	1 ML	VL	IJ	ML	ML	15 MG		1	02/28/2005	01/31/2014							
60505-0706-00		J1885		02/28/2005	01/31/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV) 30 MG/ML	1 ML	VL	IJ	ML	ML	15 MG		2	02/28/2005	01/31/2014							
60505-0706-01		J1885		02/28/2005	01/31/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 30 MG/ML	2 ML	VL	IM	ML	ML	15 MG		2	02/28/2005	01/31/2014							
60505-0715-00		J1245		08/01/2004	01/31/2014	INJECTION, DIPYRIDAMOLE, PER 10 MG	DIPYRIDAMOLE 5 MG/ML	2 ML	VL	IV	ML	ML	10 MG		0.5	08/01/2004	01/31/2014							
60505-0715-01		J1245		08/01/2004	01/31/2014	INJECTION, DIPYRIDAMOLE, PER 10 MG	DIPYRIDAMOLE (10X10) 5 MG/ML	10 ML	VL	IV	ML	ML	10 MG		0.5	08/01/2004	01/31/2014							
60505-0722-00		J0282		06/01/2003	01/31/2014	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL (SDV) 50 MG/ML	3 ML	VL	IV	ML	ML	30 MG		1.66666	06/01/2003	01/31/2014							
60505-0722-01		J0282		12/20/2005	01/31/2014	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL (SDS, 10X3ML) 50 MG/ML	3 ML	SR	IV	ML	ML	30 MG		1.66666	12/20/2005	01/31/2014							
60505-0725-01		J1885		11/01/2004	01/31/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 15 MG/ML	1 ML	SR	IJ	ML	ML	15 MG		1	11/01/2004	01/31/2014							
60505-0726-01		J1885		11/01/2004	01/31/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/ML	1 ML	SR	IJ	ML	ML	15 MG		2	11/01/2004	01/31/2014							
60505-0726-02		J1885		11/01/2004	01/31/2014	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/ML	2 ML	SR	IJ	ML	ML	15 MG		2	11/01/2004	01/31/2014							
60505-0727-03		J1630		01/24/2005	01/31/2014	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (PF) 5 MG/ML	1 ML	SR	IM	ML	ML	5 MG		1	01/24/2005	01/31/2014							
60505-0733-01		J1450		05/25/2005	08/08/2013	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (FLEXBAG) 200 MG/100 ML	100 ML	PC	IV	ML	ML	200 MG		0.01	05/25/2005	08/08/2013							
60505-0733-02		J1450		05/25/2005	08/08/2013	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (FLEXBAG) 400 MG/200 ML	200 ML	PC	IV	ML	ML	200 MG		0.01	05/25/2005	08/08/2013							
60505-0734-01		J1450		05/25/2005	08/08/2013	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (FLEXBAG, DEXTROSE) 200 MG/100 ML	100 ML	PC	IV	ML	ML	200 MG		0.01	05/25/2005	08/08/2013							
60505-0734-02		J1450		05/25/2005	01/31/2014	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (FLEXBAG, DEXTROSE) 400 MG/200 ML	200 ML	PC	IV	ML	ML	200 MG		0.01	05/25/2005	01/31/2014							
60505-0744-01		J2405		12/26/2006	01/31/2014	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (5X2ML,SDV,USP) 2 MG/ML	2 ML	VL	IJ	ML	ML	1 MG		2	12/26/2006	01/31/2014							
60505-0744-06		J2405		12/26/2006	01/31/2014	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV,USP) 2 MG/ML	20 ML	VL	IJ	ML	ML	1 MG		2	12/26/2006	01/31/2014							
60505-0748-04		J0690		09/19/2005	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM 500 MG	1 EA	VL	IJ	EA	EA	500 MG		1	09/19/2005	99/99/9999							
60505-0748-05		J0690		09/19/2005	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM 500 MG	1 EA	VL	IJ	EA	EA	500 MG		1	09/19/2005	99/99/9999							
60505-0749-04		J0690		09/19/2005	05/26/2016	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM 1 GM	1 EA	VL	IJ	EA	EA	500 MG		2	09/19/2005	05/26/2016							
60505-0749-05		J0690		09/16/2005	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM 1 GM	1 EA	VL	IJ	EA	EA	500 MG		2	09/16/2005	99/99/9999							
60505-0750-00		J0696		08/02/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (1X10ML) 250 MG	1 EA	VL	IJ	EA	EA	250 MG		1	08/02/2005	99/99/9999							
60505-0750-04		J0696		08/02/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (10X10ML) 250 MG	1 EA	VL	IJ	EA	EA	250 MG		1	08/02/2005	99/99/9999							
60505-0751-00		J0696		08/02/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (1X10ML) 500 MG	1 EA	VL	IJ	EA	EA	250 MG		2	08/02/2005	99/99/9999							
60505-0751-04		J0696		08/02/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (10X10ML) 500 MG	1 EA	VL	IJ	EA	EA	250 MG		2	08/02/2005	99/99/9999							
60505-0752-00		J0696		08/02/2005	04/17/2013	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (1X20ML) 1 GM	1 EA	VL	IJ	EA	EA	250 MG		4	08/02/2005	04/17/2013							
60505-0752-04		J0696		08/02/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (10X20ML) 1 GM	1 EA	VL	IJ	EA	EA	250 MG		4	08/02/2005	99/99/9999							
60505-0753-04		J0696		08/02/2005	99/99/9999	INJECTION, CEFTRIAZONE SODIUM, PER 250 MG	CEFTRIAZONE (10X20ML) 2 GM	1 EA	VL	IJ	EA	EA	250 MG		8	08/02/2005	99/99/9999							
60505-0759-05		J0694		01/23/2006	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN 1 GM	1 EA	VL	IV	EA	EA	1 GM		1	01/23/2006	99/99/9999							
60505-0760-05		J0694		01/23/2006	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN 2 GM	1 EA	VL	IV	EA	EA	1 GM		2	01/23/2006	99/99/9999							
60505-0761-04		J0694		02/13/2006	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN (BULK PACKAGE) 10 GM	1 EA	VL	IV	EA	EA	1 GM		10	02/13/2006	99/99/9999							
68001-0366-25		J3489		09/17/2018	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SDV) 4 MG/5 ML	5 ML	VL	IV	ML	ML	1 MG		0.8	09/17/2018	99/99/9999							
60505-0769-00		J0690		06/13/2006	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN 10 GM	1 EA	VL	IV	EA	EA	500 MG		20	06/13/2006	99/99/9999							
60505-0802-01		J7631		05/31/2002	01/31/2014	CROMOLYN SODIUM, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (AMP) 10 MG/ML	2 ML	PC	IH	ML	ML	10 MG		1	05/31/2002	01/31/2014							
60505-0802-01	KO	J7631	KO	05/31/2002	01/31/2014	CROMOLYN SODIUM, INHALATION SOLUTION ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (AMP) 10 MG/ML	2 ML	PC	IH	ML	ML	10 MG		1	05/31/2002	01/31/2014							
60505-0806-01		J7644		01/01/2002	01/31/2014	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (AMP) 0.02%	2.5 ML	PC	IH	ML	ML	1 MG		0.2	01/01/2002	01/31/2014							
60505-0806-01	KO	J7644	KO	01/01/2002	01/31/2014	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (AMP) 0.02%	2.5 ML	PC	IH	ML	ML	1 MG		0.2	01/01/2002	01/31/2014							
60505-0807-01		J7669		01/01/2002	01/31/2014	METAPROTERENOL SULFATE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (AMP) 0.4%	2.5 ML	PC	IH	ML	ML	10 MG		0.4	01/01/2002	01/31/2014							
60505-0807-01	KO	J7669	KO	01/01/2002	01/31/2014	METAPROTERENOL SULFATE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (AMP) 0.4%	2.5 ML	PC	IH	ML	ML	10 MG		0.4	01/01/2002	01/31/2014							
60505-0808-01		J7669		01/01/2002	01/31/2014	METAPROTERENOL SULFATE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (AMP) 0.6%	2.5 ML	PC	IH	ML	ML	10 MG		0.6	01/01/2002	01/31/2014							
60505-0808-01	KO	J7669	KO	01/01/2002	01/31/2014	METAPROTERENOL SULFATE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (AMP) 0.6%	2.5 ML	PC	IH	ML	ML	10 MG		0.6	01/01/2002	01/31/2014							
60505-0834-00		J0692		06/19/2007	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP) 1 GM	1 EA	VL	IJ	EA	EA	500 MG		2	06/19/2007	99/99/9999							
60505-0834-04		J0692		06/19/2007	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP) 1 GM	10 EA	VL	IJ	EA	EA	500 MG		2	06/19/2007	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
60505-5306-01		J8499		03/01/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (USP) 400 MG	100 EA	BO PO EA				1 EA		1	03/01/2006	99/99/9999						
60505-5306-08		J8499		05/21/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	1000 EA	BO PO EA				1 EA		1	05/21/2007	99/99/9999						
60505-5307-01		J8499		03/01/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (USP) 800 MG	100 EA	BO PO EA				1 EA		1	03/01/2006	99/99/9999						
60505-5307-05		J8499		05/21/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	500 EA	BO PO EA				1 EA		1	05/21/2007	99/99/9999						
60505-6020-02		J1631		01/30/2008	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	NOVAPLUS HALOPERIDOL DECAANOATE (1X5ML.MDV) 50 MG/ML	5 ML	VL IM ML				50 MG		1	01/30/2008	99/99/9999						
60505-6021-02		J1631		12/14/2007	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	NOVAPLUS HALOPERIDOL DECAANOATE (1X5ML.MDV) 100 MG/ML	5 ML	VL IM ML				50 MG		2	12/14/2007	99/99/9999						
60505-6025-05		J0694		02/27/2008	02/22/2018	INJECTION, CEFOXITIN SODIUM, 1 GM	NOVAPLUS CEFOXITIN (USP) 1 GM	1 EA	VL IV EA				1 GM		1	02/27/2008	02/22/2018						
60505-6026-05		J0694		02/27/2008	04/24/2018	INJECTION, CEFOXITIN SODIUM, 1 GM	NOVAPLUS CEFOXITIN (USP) 2 GM	1 EA	VL IV EA				1 GM		2	02/27/2008	04/24/2018						
60505-6030-04		J0692		04/11/2008	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP) 1 GM	1 EA	VL U EA				500 MG		2	04/11/2008	99/99/9999						
60505-6031-04		J0692		04/11/2008	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP) 2 GM	1 EA	VL U EA				500 MG		4	04/11/2008	99/99/9999						
60760-0002-21		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	21 EA	BO PO EA				5 MG		4	05/15/2009	12/31/2015	01/01/2002	09/26/2002	4	03/01/2006	09/01/2007	4
60760-0330-30		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30 EA	BO PO EA				50 MG		0.5	01/01/2002	02/03/2016						
60760-0830-20		Q0170		06/01/2005	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20 EA	BO PO EA				25 MG		1	06/01/2005	12/31/2013						
60793-0130-10		J2510		09/14/2007	99/99/9999	INJECTION, PENICILLIN G PROCAINE, AQUEOUS, UP TO 600,000 UNITS	PENICILLIN G PROCAINE (21GX1&1/2,1MLX10) 600000 U/ML	1 ML	SR IM ML				600000 U		1	09/14/2007	99/99/9999						
60793-0131-10		J2510		09/14/2007	99/99/9999	INJECTION, PENICILLIN G PROCAINE, AQUEOUS, UP TO 600,000 UNITS	PENICILLIN G PROCAINE (21GX1&1/4,2MLX10) 600000 U/ML	2 ML	SR IM ML				600000 U		1	09/14/2007	99/99/9999						
60977-0001-43		J2550		05/05/2007	10/17/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PHENERGAN 25 MG/ML	1 ML	AM U ML				50 MG		0.5	05/05/2007	10/17/2016						
60977-0001-44		J2550		05/05/2007	04/30/2013	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PHENERGAN 25 MG/ML	1 ML	VL U ML				50 MG		0.5	05/05/2007	04/30/2013						
60977-0001-44		J2550		05/05/2007	04/30/2013	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), NOT OTHERWISE SPECIFIED, 500 MG	PANZYGA (PF,LATEX-FREE) 100 MG/1 ML	10 ML	BO IV ML				500 MG		0.2	11/12/2018	99/99/9999						
60977-0002-43		J2550		05/05/2007	10/17/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PHENERGAN 50 MG/ML	1 ML	AM U ML				50 MG		1	05/05/2007	10/17/2016						
60977-0002-44		J2550		05/05/2007	10/17/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PHENERGAN 50 MG/ML	1 ML	VL U ML				50 MG		1	05/05/2007	10/17/2016						
60977-0016-02		J2275		01/15/2004	07/02/2012	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	DURAMORPH (AMP,DOSETTE,PF) 0.5 MG/ML	10 ML	AM U ML				10 MG		0.05	01/15/2004	07/02/2012						
60977-0016-73		J2275		05/05/2007	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	DURAMORPH (PF) 0.5 MG/ML	10 ML	AM U ML				10 MG		0.05	05/05/2007	12/31/2014						
60977-0017-01		J2275		01/15/2004	07/02/2012	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	DURAMORPH (AMP,DOSETTE,PF) 1 MG/ML	10 ML	AM U ML				10 MG		0.1	01/15/2004	07/02/2012						
60977-0112-01		J2060		02/13/2004	04/05/2012	INJECTION, LORAZEPAM, 2 MG	ATVAN (S.D.V.) 2 MG/ML	1 ML	VL U ML				2 MG		1	02/13/2004	04/05/2012						
60977-0112-81		J2060		05/05/2007	02/28/2014	INJECTION, LORAZEPAM, 2 MG	ATVAN (SDV) 2 MG/ML	1 ML	VL U ML				2 MG		1	05/05/2007	02/28/2014						
60977-0113-01		J2060		02/13/2004	04/05/2012	INJECTION, LORAZEPAM, 2 MG	ATVAN (S.D.V.) 4 MG/ML	1 ML	VL U ML				2 MG		2	02/13/2004	04/05/2012						
60977-0113-02		J2060		02/13/2004	04/05/2012	INJECTION, LORAZEPAM, 2 MG	ATVAN (M.D.V.) 4 MG/ML	10 ML	VL U ML				2 MG		2	02/13/2004	04/05/2012						
60977-0113-71		J2060		05/05/2007	12/31/2013	INJECTION, LORAZEPAM, 2 MG	ATVAN (MDV) 4 MG/ML	10 ML	VL U ML				2 MG		2	05/05/2007	12/31/2013						
60977-0113-81		J2060		05/05/2007	01/31/2014	INJECTION, LORAZEPAM, 2 MG	ATVAN 4 MG/ML	1 ML	VL U ML				2 MG		2	05/05/2007	01/31/2014						
60977-0114-01		J2275		01/01/2004	07/24/2012	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	INFUMORPH 200 (AMP, DOSETTE,PF) 10 MG/ML	20 ML	AM U ML				10 MG		1	01/01/2004	07/24/2012						
60977-0114-74		J2275		05/05/2007	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	INFUMORPH 200 (PF) 10 MG/ML	1 ML	NA U ML				10 MG		1	05/05/2007	12/31/2014						
60977-0115-01		J2275		01/01/2004	07/24/2012	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	INFUMORPH 500 (AMP, DOSETTE,PF) 25 MG/ML	20 ML	AM U ML				10 MG		2.5	01/01/2004	07/24/2012						
60977-0115-74		J2275		05/05/2007	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	INFUMORPH 500 (PF) 25 MG/ML	1 ML	NA U ML				10 MG		2.5	05/05/2007	12/31/2014						
60977-0141-01		J2730		12/20/2004	99/99/9999	INJECTION, PRALDOXIME CHLORIDE, UP TO 1 GM	PROTOPAM CHLORIDE (S.D.V.) 1 GM	1 EA	VL U EA				1 GM		1	12/20/2004	99/99/9999						
60977-0141-27		J2730		05/05/2007	99/99/9999	INJECTION, PRALDOXIME CHLORIDE, UP TO 1 GM	PROTOPAM CHLORIDE 1 GM	1 EA	VL U EA				1 GM		1	05/05/2007	99/99/9999						
60977-0150-01		J2800		01/01/2004	07/24/2012	INJECTION, METHOCARBAMOL, UP TO 10 ML	ROBAXIN (S.D.V.) 100 MG/ML	10 ML	VL U ML				10 ML		0.1	01/01/2004	07/24/2012						
60977-0150-71		J2800		05/05/2007	10/17/2016	INJECTION, METHOCARBAMOL, UP TO 10 ML	ROBAXIN (SDV) 100 MG/ML	10 ML	VL U ML				10 ML		0.1	05/05/2007	10/17/2016						
60977-0155-01		J7643		02/13/2004	10/18/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (S.D.V.) 0.2 MG/ML	1 ML	VL U ML				1 MG		0.2	02/13/2004	10/18/2012						
60977-0155-01	KO	J7643	KO	02/13/2004	10/18/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (S.D.V.) 0.2 MG/ML	1 ML	VL U ML				1 MG		0.2	02/13/2004	10/18/2012						
60977-0155-02		J7643		02/13/2004	10/18/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (S.D.V.) 0.2 MG/ML	2 ML	VL U ML				1 MG		0.2	02/13/2004	10/18/2012						
60977-0155-02	KO	J7643	KO	02/13/2004	10/18/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (S.D.V.) 0.2 MG/ML	2 ML	VL U ML				1 MG		0.2	02/13/2004	10/18/2012						
60977-0155-03		J7643		02/13/2004	07/24/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (M.D.V.) 0.2 MG/ML	5 ML	VL U ML				1 MG		0.2	02/13/2004	07/24/2012						
60977-0155-03	KO	J7643	KO	02/13/2004	07/24/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (M.D.V.) 0.2 MG/ML	5 ML	VL U ML				1 MG		0.2	02/13/2004	07/24/2012						
60977-0155-06		J7643		03/02/2006	07/24/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (10X20ML.MDV) 0.2 MG/ML	20 ML	VL U ML				1 MG		0.2	03/02/2006	07/24/2012						
60977-0155-06	KO	J7643	KO	03/02/2006	07/24/2012	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (10X20ML.MDV) 0.2 MG/ML	20 ML	VL U ML				1 MG		0.2	03/02/2006	07/24/2012						
60977-0155-17		J7643		05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL 0.2 MG/ML	2 ML	VL U ML				1 MG		0.2	05/05/2007	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
60977-0155-17	KO	J7643	KO	05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL 0.2 MG/ML	2 ML	VL	U	ML		1 MG		0.2	05/05/2007	02/03/2016						
60977-0155-54		J7643		05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL 0.2 MG/ML	5 ML	VL	U	ML		1 MG		0.2	05/05/2007	02/03/2016						
60977-0155-54	KO	J7643	KO	05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL 0.2 MG/ML	5 ML	VL	U	ML		1 MG		0.2	05/05/2007	02/03/2016						
60977-0155-63		J7643		05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (MDV) 0.2 MG/ML	20 ML	VL	U	ML		1 MG		0.2	05/05/2007	02/03/2016						
60977-0155-63	KO	J7643	KO	05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL (MDV) 0.2 MG/ML	20 ML	VL	U	ML		1 MG		0.2	05/05/2007	02/03/2016						
60977-0155-81		J7643		05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL 0.2 MG/ML	1 ML	VL	U	ML		1 MG		0.2	05/05/2007	02/03/2016						
60977-0155-81	KO	J7643	KO	05/05/2007	02/03/2016	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	ROBINUL 0.2 MG/ML	1 ML	VL	U	ML		1 MG		0.2	05/05/2007	02/03/2016						
12496-0100-01		J3490		01/01/2018	06/30/2018	UNCLASSIFIED DRUGS	SUBLOCADE 100 MG/0.5 ML	0.5 ML	SR	SC	ML		1 MG		1	01/01/2018	06/30/2018						
12496-0300-01		J3490		01/01/2018	06/30/2018	UNCLASSIFIED DRUGS	SUBLOCADE 100 MG/0.5 ML	1.5 ML	SR	SC	ML		1 MG		1	01/01/2018	06/30/2018						
60977-0451-17		J2765		05/05/2007	04/30/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	REGLAN (PF) 5 MG/ML	2 ML	VL	IV	ML		10 MG		0.5	05/05/2007	04/30/2013						
60977-0451-71		J2765		05/05/2007	04/30/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	REGLAN (PF) 5 MG/ML	10 ML	VL	IV	ML		10 MG		0.5	05/05/2007	04/30/2013						
60977-0451-82		J2765		05/05/2007	04/30/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	REGLAN (PF) 5 MG/ML	30 ML	VL	IV	ML		10 MG		0.5	05/05/2007	04/30/2013						
61553-0107-02		J3010		02/02/2004	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (INTRAVIA) 0.5 MG/100 ML-0.9%	250 ML	BG	IV	ML		0.1 MG		0.05	02/02/2004	99/99/9999						
61553-0109-72		J3010		02/02/2004	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (SRN,12 ML) 0.5 MG/100 ML-0.9%	10 ML	SR	IV	ML		0.1 MG		0.05	02/02/2004	99/99/9999						
61553-0111-48		J3010		02/02/2004	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (INTRAVIA) 1 MG/100 ML-0.9%	100 ML	BG	IV	ML		0.1 MG		0.1	02/02/2004	99/99/9999						
61553-0112-48		J3010		02/02/2004	06/30/2017	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (PUMP BAG) 1 MG/100 ML-0.9%	100 ML	BG	IV	ML		0.1 MG		0.1	02/02/2004	06/30/2017						
61553-0113-02		J3010		02/02/2004	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (INTRAVIA) 1 MG/100 ML-0.9%	250 ML	BG	IV	ML		0.1 MG		0.1	02/02/2004	99/99/9999						
61553-0114-02		J3010		02/02/2004	02/17/2015	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (PUMP BAG) 1 MG/100 ML-0.9%	250 ML	BG	IV	ML		0.1 MG		0.1	02/02/2004	02/17/2015						
61553-0116-48		J3010		02/02/2004	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (INTRAVIA) 2 MG/100 ML-0.9%	100 ML	BG	IV	ML		0.1 MG		0.2	02/02/2004	99/99/9999						
61553-0118-41		J3010		02/02/2004	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (INTRAVIA) 0.05 MG/ML	50 ML	NA	IV	ML		0.1 MG		0.5	02/02/2004	99/99/9999						
61553-0161-41		J1170		02/02/2004	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL/SODIUM CHLORIDE (INTRAVIA) 10 MG/50 ML-0.9%	50 ML	BG	IV	ML		4 MG		0.05	02/02/2004	99/99/9999						
61553-0162-67		J1170		02/02/2004	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL/SODIUM CHLORIDE (SRN,35 ML) 1 MG/5 ML-0.9%	25 ML	SR	IV	ML		4 MG		0.05	02/02/2004	99/99/9999						
61553-0163-75		J1170		02/02/2004	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL/SODIUM CHLORIDE (SRN,60 ML) 1 MG/5 ML-0.9%	50 ML	SR	IV	ML		4 MG		0.05	02/02/2004	99/99/9999						
61553-0165-41		J1170		02/02/2004	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL/SODIUM CHLORIDE (INTRAVIA) 50 MG/50 ML-0.9%	50 ML	BG	IV	ML		4 MG		0.25	02/02/2004	99/99/9999						
61553-0166-67		J1170		02/02/2004	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL/SODIUM CHLORIDE (SRN,35 ML) 1 MG/ML-0.9%	25 ML	SR	IV	ML		4 MG		0.25	02/02/2004	99/99/9999						
61553-0167-75		J1170		02/02/2004	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL/SODIUM CHLORIDE (SRN,50 ML) 1 MG/ML-0.9%	50 ML	SR	IV	ML		4 MG		0.25	02/02/2004	99/99/9999						
61553-0170-41		J2175		02/02/2004	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL/SODIUM CHLORIDE (INTRAVIA) 500 MG/50 ML-0.9%	50 ML	BG	IV	ML		100 MG		0.1	02/02/2004	99/99/9999						
61553-0172-48		J2175		02/02/2004	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL/SODIUM CHLORIDE (INTRAVIA) 1 GM/100 ML-0.9%	100 ML	BG	IV	ML		100 MG		0.1	02/02/2004	99/99/9999						
61553-0173-48		J2175		02/02/2004	06/30/2017	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL/SODIUM CHLORIDE (PUMP BAG) 1 GM/100 ML-0.9%	100 ML	BG	IV	ML		100 MG		0.1	02/02/2004	06/30/2017						
61553-0177-41		J2270		02/02/2004	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE/SODIUM CHLORIDE (INTRAVIA) 50 MG/50 ML-0.9%	50 ML	BG	IV	ML		10 MG		0.1	02/02/2004	99/99/9999						
61553-0178-48		J2270		02/02/2004	06/30/2017	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE/SODIUM CHLORIDE (PUMP BAG) 100 MG/100 ML-0.9%	100 ML	BG	IV	ML		10 MG		0.1	02/02/2004	06/30/2017						
61553-0179-48		J2270		02/02/2004	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE/SODIUM CHLORIDE (INTRAVIA) 100 MG/100 ML-0.9%	150 ML	BG	IV	ML		10 MG		0.1	02/02/2004	99/99/9999						
61553-0181-02		J2270		02/02/2004	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE/SODIUM CHLORIDE (INTRAVIA) 250 MG/250 ML-0.9%	250 ML	BG	IV	ML		10 MG		0.1	02/02/2004	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3		
61553-0183-48	J2270			02/02/2004	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	DEXTROSE/MORPHINE SULFATE (INTRAVIA) 5%-100 MG/100 ML	100	ML	NA	IV	ML	10	MG	0.1	02/02/2004	99/99/9999								
61553-0185-02	J2270			02/02/2004	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	DEXTROSE/MORPHINE SULFATE (INTRAVIA) 5%-100 MG/100 ML	250	ML	NA	IV	ML	10	MG	0.1	02/02/2004	99/99/9999								
61553-0186-67	J2270			02/02/2004	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	DEXTROSE/MORPHINE SULFATE (SRN,35 ML) 5%-2 MG/ML	25	ML	NA	IV	ML	10	MG	0.2	02/02/2004	99/99/9999								
61553-0187-75	J2270			02/02/2004	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	DEXTROSE/MORPHINE SULFATE (SRN,60 ML) 5%-2 MG/ML	50	ML	NA	IV	ML	10	MG	0.2	02/02/2004	99/99/9999								
61553-0189-48	J3490			02/02/2004	03/31/2017	UNCLASSIFIED DRUGS	BUPIVACAINE/SODIUM CHLORIDE (INTRAVIA) 0.0625%-0.9%	100	ML	BG	IV	ML	1	EA	1	02/02/2004	03/31/2017								
61553-0190-48	J3490			02/02/2004	06/30/2017	UNCLASSIFIED DRUGS	BUPIVACAINE/SODIUM CHLORIDE (PUMP BAG) 0.0625%-0.9%	100	ML	BG	IV	ML	1	EA	1	02/02/2004	06/30/2017								
61553-0191-48	J3490			02/02/2004	03/31/2017	UNCLASSIFIED DRUGS	BUPIVACAINE/SODIUM CHLORIDE (INTRAVIA) 0.125%-0.9%	100	ML	BG	IV	ML	1	EA	1	02/02/2004	03/31/2017								
61553-0192-02	J3490			02/02/2004	03/31/2017	UNCLASSIFIED DRUGS	BUPIVACAINE/SODIUM CHLORIDE (INTRAVIA) 0.125%-0.9%	250	ML	BG	IV	ML	1	EA	1	02/02/2004	03/31/2017								
61553-0193-41	J3490			02/02/2004	03/31/2017	UNCLASSIFIED DRUGS	BUPIVACAINE/SODIUM CHLORIDE (INTRAVIA) 0.25%-0.9%	50	ML	BG	IV	ML	1	EA	1	02/02/2004	03/31/2017								
61553-0194-48	J3490			02/02/2004	06/30/2017	UNCLASSIFIED DRUGS	BUPIVACAINE/SODIUM CHLORIDE (PUMP BAG) 0.125%-0.9%	100	ML	BG	IV	ML	1	EA	1	02/02/2004	06/30/2017								
61553-0228-02	J3490			11/21/2007	03/31/2017	UNCLASSIFIED DRUGS	ROPIVACAINE HYDROCHLORIDE-SODIUM CHLORIDE 0.2%-0.9%	250	ML	NA	EP	ML	1	EA	1	11/21/2007	03/31/2017								
61553-0421-04	J3475			02/01/2005	03/31/2017	INJECTION, MAGNESIUM SULFATE, PER 500 MG	DEXTROSE-MAGNESIUM SULFATE (6X1000ML, VIAFLEX BAG) 5%-20 GM	1000	ML	NA	IV	ML	500	MG	0.04	02/01/2005	03/31/2017								
61553-0423-02	J3475			07/11/2005	12/31/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE IN DEXTROSE (24X250ML) 5%-8 GM/100 ML	250	ML	NA	IV	ML	500	MG	0.16	07/11/2005	12/31/2016								
61553-0602-48	J3010			02/02/2004	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE/SODIUM CHLORIDE (INTRAVIA) 0.2 MG/100 ML-0.9%	100	ML	BG	IV	ML	0.1	MG	0.02	02/02/2004	99/99/9999								
61553-0624-48	J1170			02/02/2004	06/30/2017	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL/SODIUM CHLORIDE (PUMP BAG) 20 MG/100 ML-0.9%	100	ML	BG	IV	ML	4	MG	0.05	02/02/2004	06/30/2017								
61553-0681-76	J1170			11/21/2007	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (5X60ML, BD SYRINGES) 0.2 MG/ML-0.9%	60	ML	SR	IV	ML	4	MG	0.05	11/21/2007	99/99/9999								
61553-0701-68	J1170			12/01/2006	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 0.1 MG/ML-0.9%	30	ML	VL	IV	ML	4	MG	0.025	12/01/2006	99/99/9999								
61553-0702-68	J1170			12/01/2006	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 0.2 MG/ML-0.9%	30	ML	VL	IV	ML	4	MG	0.05	12/01/2006	99/99/9999								
61553-0704-68	J1170			12/01/2006	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 0.4 MG/ML-0.9%	30	ML	VL	IV	ML	4	MG	0.1	12/01/2006	99/99/9999								
61553-0705-68	J1170			12/01/2006	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 0.5 MG/ML-0.9%	30	ML	VL	IV	ML	4	MG	0.125	12/01/2006	99/99/9999								
61553-0706-68	J1170			12/01/2006	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 0.6 MG/ML-0.9%	30	ML	VL	IV	ML	4	MG	0.15	12/01/2006	99/99/9999								
61553-0710-68	J1170			12/01/2006	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 1 MG/ML-0.9%	30	ML	VL	IV	ML	4	MG	0.25	12/01/2006	99/99/9999								
61553-0712-68	J1170			12/01/2006	06/30/2017	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 1.2 MG/ML-0.9%	30	ML	VL	IV	ML	4	MG	0.3	12/01/2006	06/30/2017								
61553-0730-68	J3010			11/21/2007	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 25 MCG/ML-0.9%	30	ML	VL	IV	ML	0.1	MG	0.25	11/21/2007	99/99/9999								
61553-0732-03	J2590			02/06/2004	12/31/2016	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN-SODIUM CHLORIDE (12X500ML, VIAFLEX BAG) 10 U-0.9%	500	ML	NA	IV	ML	10	U	1	02/06/2004	12/31/2016								
61553-0780-68	J1170			12/01/2006	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (10X30ML, PCA VIAL) 2 MG/ML	30	ML	VL	IV	ML	4	MG	0.5	12/01/2006	99/99/9999								
61553-0791-68	J3010			12/01/2006	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 10 MCG/ML-0.9%	30	ML	VL	IV	ML	0.1	MG	100	12/01/2006	99/99/9999								
61553-0792-68	J3010			12/01/2006	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 20 MCG/ML-0.9%	30	ML	VL	IV	ML	0.1	MG	200	12/01/2006	99/99/9999								
61553-0793-68	J3010			12/01/2006	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 30 MCG/ML-0.9%	30	ML	VL	IV	ML	0.1	MG	300	12/01/2006	99/99/9999								
61553-0794-68	J3010			12/01/2006	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE-SODIUM CHLORIDE (10X30ML, PCA VIAL) 40 MCG/ML-0.9%	30	ML	VL	IV	ML	0.1	MG	400	12/01/2006	99/99/9999								
61553-0795-68	J3010			12/01/2006	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (10X30ML, PCA VIAL) 50 MCG/ML	30	ML	VL	IV	ML	0.1	MG	500	12/01/2006	99/99/9999								

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
						TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TIGAN 300 MG	100	EA	BO	PO	EA	250	MG	1.2	02/13/2002	99/99/9999							
61570-0079-01		Q0173		02/13/2002	99/99/9999																			
61570-0260-10		J2770		06/27/2003	99/99/9999	INJECTION, QUINUPRISTIN/DALFOPRISTIN, 500 MG (150/350)	SYNERCID (PF) 350 MG-150 MG	1	EA	VL	IV	EA	500	MG	1	06/27/2003	99/99/9999							
61703-0245-22		J2405		12/26/2006	10/17/2016	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (M.D.V.,USP) 2 MG/ML	20	ML	VL	IJ	ML	1	MG	2	12/26/2006	10/17/2016							
68982-0820-02		J1599		11/12/2018	99/99/9999	OTHERWISE SPECIFIED, 500 MG	PANZYGA (PF,LATEX-FREE) 100 MG/1 ML	25	ML	BO	IV	ML	500	MG	0.2	11/12/2018	99/99/9999							
61703-0305-38		J9100		05/01/2003	99/99/9999	INJECTION, CYTARABINE, 100 MG	CYTARABINE (S.D.V. X 5,PF) 20 MG/ML	5	ML	VL	IJ	ML	100	MG	0.2	05/01/2003	99/99/9999							
61703-0309-06		J9370		01/01/2002	99/99/9999	VINCRIStINE SULFATE, 1 MG	VINCRIStINE SULFATE (S.D.V.,PF) 1 MG/ML	1	ML	VL	IV	ML	1	MG	1	01/01/2002	99/99/9999							
61703-0309-16		J9370		01/01/2002	99/99/9999	VINCRIStINE SULFATE, 1 MG	VINCRIStINE SULFATE (S.D.V.,PF) 1 MG/ML	2	ML	VL	IV	ML	1	MG	1	01/01/2002	99/99/9999							
61703-0317-45		J0595		06/25/2004	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (S.D.V.) 1 MG/ML	1	ML	VL	IJ	ML	1	MG	1	06/25/2004	99/99/9999							
61703-0318-45		J0595		06/25/2004	99/99/9999	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (S.D.V.) 2 MG/ML	1	ML	VL	IJ	ML	1	MG	2	06/25/2004	99/99/9999							
61703-0323-22		J9040		01/01/2002	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN SULFATE 30 U	1	EA	VL	IJ	EA	15	U	2	01/01/2002	99/99/9999							
61703-0324-18		J2430		12/15/2006	99/99/9999	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (SDV) 3 MG/ML	10	ML	VL	IV	ML	30	MG	0.1	12/15/2006	99/99/9999							
61703-0325-18		J2430		01/27/2003	99/99/9999	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (PF) 6 MG/ML	10	ML	VL	IV	ML	30	MG	0.2	01/27/2003	99/99/9999							
61703-0326-18		J2430		09/15/2005	99/99/9999	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM 9 MG/ML	10	ML	VL	IV	ML	30	MG	0.3	09/15/2005	99/99/9999							
61703-0332-18		J9040		01/01/2002	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN SULFATE 15 U	1	EA	VL	IJ	EA	15	U	1	01/01/2002	99/99/9999							
68982-0820-03		J1599		11/12/2018	99/99/9999	OTHERWISE SPECIFIED, 500 MG	PANZYGA (PF,LATEX-FREE) 100 MG/1 ML	50	ML	BO	IV	ML	500	MG	0.2	11/12/2018	99/99/9999							
61703-0339-18		J9045		04/14/2004	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (MDV) 10 MG/ML	5	ML	VL	IV	ML	50	MG	0.2	04/14/2004	99/99/9999							
61703-0339-22		J9045		04/14/2004	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (MDV) 10 MG/ML	15	ML	VL	IV	ML	50	MG	0.2	04/14/2004	99/99/9999							
61703-0339-50		J9045		04/14/2004	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (MDV) 10 MG/ML	45	ML	VL	IV	ML	50	MG	0.2	04/14/2004	99/99/9999							
61703-0339-56		J9045		02/09/2005	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (MDV) 10 MG/ML	60	ML	VL	IV	ML	50	MG	0.2	02/09/2005	99/99/9999							
61703-0341-06		J9390		09/07/2005	10/31/2017	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (S.D.V.,PF) 10 MG/ML	1	ML	VL	IV	ML	10	MG	1	09/07/2005	10/31/2017							
61703-0341-09		J9390		11/07/2005	03/30/2018	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (S.D.V.,PF) 10 MG/ML	5	ML	VL	IV	ML	10	MG	1	11/07/2005	03/30/2018							
61703-0342-09		J9265		04/21/2004	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (M.D.V.) 6 MG/ML	5	ML	VL	IV	ML	30	MG	0.2	04/21/2004	12/31/2014							
61703-0342-22		J9265		04/21/2004	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (M.D.V.) 6 MG/ML	16.7	ML	VL	IV	ML	30	MG	0.2	04/21/2004	12/31/2014							
61703-0342-50		J9265		04/21/2004	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (M.D.V.) 6 MG/ML	50	ML	VL	IV	ML	30	MG	0.2	04/21/2004	12/31/2014							
61703-0343-18		J9293		04/11/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (USP,CONCENTRATE,MDV,PF) 2 MG/ML	10	ML	VL	IV	ML	5	MG	0.4	04/11/2006	99/99/9999							
61703-0343-65		J9293		04/11/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (USP,CONCENTRATE,MDV,PF) 2 MG/ML	12.5	ML	VL	IV	ML	5	MG	0.4	04/11/2006	99/99/9999							
61703-0343-66		J9293		04/11/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (USP,CONCENTRATE,MDV,PF) 2 MG/ML	15	ML	VL	IV	ML	5	MG	0.4	04/11/2006	99/99/9999							
61703-0347-35		J9178		11/06/2006	08/31/2014	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (S.D.V.) 50 MG	1	EA	VL	IV	EA	2	MG	25	11/06/2006	08/31/2014							
61703-0349-09		J9206		02/27/2008	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML) 20 MG/ML	5	ML	VL	IV	ML	20	MG	1	02/27/2008	99/99/9999							
61703-0349-16		J9206		02/27/2008	99/99/9999	IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML) 20 MG/ML	2	ML	VL	IV	ML	20	MG	1	02/27/2008	99/99/9999							
61703-0349-36		J9206		02/27/2008	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X25ML,SDV) 20 MG/ML	25	ML	VL	IV	ML	20	MG	1	02/27/2008	99/99/9999							
61703-0350-38		J9250		06/27/2005	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (MDV,5X2ML) 25 MG/ML	2	ML	VL	IJ	ML	5	MG	5	06/27/2005	99/99/9999							
61703-0356-18		J2430		12/15/2006	08/31/2015	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	NOVAPLUS PAMIDRONATE DISODIUM (SDV) 9 MG/ML	10	ML	VL	IV	ML	30	MG	0.3	12/15/2006	08/31/2015							
61703-0359-01		J9178		04/10/2008	11/30/2015	INJECTION, EPIRUBICIN HCL, 2 MG	NOVAPLUS EPIRUBICIN HYDROCHLORIDE (1X25ML,SINGLE USE,PF) 2 MG/ML	25	ML	VL	IV	ML	2	MG	1	04/10/2008	11/30/2015							
61703-0359-02		J9178		04/10/2008	01/31/2015	INJECTION, EPIRUBICIN HCL, 2 MG	NOVAPLUS EPIRUBICIN HYDROCHLORIDE (1X100ML,SINGLE USE,PF) 2 MG/ML	100	ML	VL	IV	ML	2	MG	1	04/10/2008	01/31/2015							
61703-0359-59		J9178		08/08/2007	06/05/2017	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (PF) 2 MG/ML	100	ML	VL	IV	ML	2	MG	1	08/08/2007	06/05/2017							
61703-0359-93		J9178		08/08/2007	06/05/2017	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (PF) 2 MG/ML	25	ML	VL	IV	ML	2	MG	1	08/08/2007	06/05/2017							
68982-0820-04		J1599		11/12/2018	99/99/9999	OTHERWISE SPECIFIED, 500 MG	PANZYGA (PF,LATEX-FREE) 100 MG/1 ML	100	ML	BO	IV	ML	500	MG	0.2	11/12/2018	99/99/9999							
61703-0360-18		J9045		06/28/2006	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	NOVAPLUS CARBOPLATIN (MDV) 10 MG/ML	5	ML	VL	IV	ML	50	MG	0.2	06/28/2006	99/99/9999							
61703-0360-22		J9045		06/28/2006	10/31/2015	INJECTION, CARBOPLATIN, 50 MG	NOVAPLUS CARBOPLATIN (MDV) 10 MG/ML	15	ML	VL	IV	ML	50	MG	0.2	06/28/2006	10/31/2015							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
61703-0360-50		J9045		06/28/2006	01/31/2016	INJECTION, CARBOPLATIN, 50 MG	NOVAPLUS CARBOPLATIN (MDV) 10 MG/ML	45	ML	VL	IV	ML	50	MG	0.2	06/28/2006	01/31/2016						
61703-0408-41		J9250		04/09/2004	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (SDV,PF) 25 MG/ML	40	ML	VL	IJ	ML	5	MG	5	06/27/2005	99/99/9999	04/09/2004	01/17/2005	5			
68982-0820-05		J1599		11/12/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), NOT OTHERWISE SPECIFIED, 500 MG	PANZYGA (PF,LATEX-FREE) 100 MG/1 ML	200	ML	BO	IV	ML	500	MG	0.2	11/12/2018	99/99/9999						
61953-0004-01		J1572		01/01/2008	99/99/9999	NON-LYOPHILIZED (E.G., LIQUID), 500 MG	FLEBOGAMMA (DIF,PF) 5 GM/100 ML	10	ML	VL	IV	ML	500	MG	0.1	01/01/2008	99/99/9999						
61953-0004-02		J1572		01/01/2008	99/99/9999	NON-LYOPHILIZED (E.G., LIQUID), 500 MG	FLEBOGAMMA (DIF,PF) 5 GM/100 ML	50	ML	VL	IV	ML	500	MG	0.1	01/01/2008	99/99/9999						
61953-0004-03		J1572		01/01/2008	99/99/9999	NON-LYOPHILIZED (E.G., LIQUID), 500 MG	FLEBOGAMMA (DIF,PF) 5 GM/100 ML	100	ML	VL	IV	ML	500	MG	0.1	01/01/2008	99/99/9999						
61953-0004-04		J1572		01/01/2008	99/99/9999	NON-LYOPHILIZED (E.G., LIQUID), 500 MG	FLEBOGAMMA (DIF,PF) 5 GM/100 ML	200	ML	VL	IV	ML	500	MG	0.1	01/01/2008	99/99/9999						
61953-0004-05		J1572		01/01/2008	99/99/9999	NON-LYOPHILIZED (E.G., LIQUID), 500 MG	FLEBOGAMMA (DIF,PF) 5 GM/100 ML	400	ML	VL	IV	ML	500	MG	0.1	01/01/2008	99/99/9999						
61958-0101-01		J0740		01/01/2002	12/01/2016	INJECTION, IMMUNE GLOBULIN, (FLEBOGAMMA/FLEBOGAMMA DIF), INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), 500 MG	VISTIDE (S.D.V.,PF) 75 MG/ML	5	ML	VL	IV	ML	375	MG	0.2	01/01/2002	12/01/2016						
62033-0204-10		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	100	EA	BO	PO	EA	1	EA	1	01/01/2002	02/03/2016						
62033-0204-14		J8499		01/01/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	400	EA	BO	PO	EA	1	EA	1	01/01/2002	02/03/2016						
68982-0820-06		J1599		11/12/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), NOT OTHERWISE SPECIFIED, 500 MG	PANZYGA (PF,LATEX-FREE) 100 MG/1 ML	300	ML	BO	IV	ML	500	MG	0.2	11/12/2018	99/99/9999						
69543-0371-10		J2469		09/20/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL 0.05 MG/1 ML	5	ML	VL	IV	ML	25	MCG	2	09/20/2018	99/99/9999						
70069-0030-03		J1631		10/04/2018	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (3X1ML) 50 MG/1 ML	1	ML	AM	IM	ML	50	MG	1	10/04/2018	99/99/9999						
70069-0031-05		J1631		10/04/2018	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (5X1ML) 100 MG/1 ML	1	ML	AM	IM	ML	50	MG	2	10/04/2018	99/99/9999						
70121-1577-05		J2370		10/04/2018	99/99/9999	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	PHENYLEPHRINE HCL (LATEX-FREE) 10 MG/1 ML	1	ML	VL	IV	ML	1	ML	1	10/04/2018	99/99/9999						
70655-0002-06		J1450		08/31/2018	99/99/9999	INJECTION, FLUCONAZOLE, 200 MG	FLUCONAZOLE (PF,LATEX-FREE) 200 MG/100 ML	100	ML	BX	IV	ML	200	MG	0.01	08/31/2018	99/99/9999						
62756-0181-01		J2405		12/27/2006	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (5X2ML,SDA,USP) 2 MG/ML	2	ML	AM	IJ	ML	1	MG	2	12/27/2006	99/99/9999						
62756-0581-40		J0207		03/26/2008	99/99/9999	INJECTION, AMIFOSTINE, 500 MG	AMIFOSTINE (USP) 500 MG	1	EA	VL	IV	EA	500	MG	1	03/26/2008	99/99/9999						
62756-0581-42		J0207		03/26/2008	99/99/9999	INJECTION, AMIFOSTINE, 500 MG	AMIFOSTINE (USP) 500 MG	1	EA	VL	IV	EA	500	MG	1	03/26/2008	99/99/9999						
62856-0101-10		J1645		11/20/2006	03/31/2015	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (27GX1/2"W/NDL,GUARD) 10000 IU/ML	1	ML	SR	SC	ML	2500	IU	4	11/20/2006	03/31/2015						
62856-0125-10		J1645		08/25/2007	03/31/2015	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (SINGLE DOSE,PF) 12500 IU/0.5 ML	0.5	ML	SR	SC	ML	2500	IU	10	08/25/2007	03/31/2015						
62856-0150-10		J1645		08/25/2007	03/31/2015	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (SINGLE DOSE,PF) 15000 IU/0.6 ML	0.6	ML	SR	SC	ML	2500	IU	10.66666	08/25/2007	03/31/2015						
62856-0180-10		J1645		08/25/2007	03/31/2015	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (SINGLE DOSE,PF) 18000 IU/0.72 ML	0.72	ML	SR	SC	ML	2500	IU	10	08/25/2007	03/31/2015						
62856-0250-10		J1645		06/26/2007	03/31/2015	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (10X0.2ML,PF) 2500 IU/0.2 ML	0.2	ML	SR	SC	ML	2500	IU	5	06/26/2007	03/31/2015						
62856-0251-01		J1645		11/20/2006	12/01/2014	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (MDV) 25000 IU/ML	3.8	ML	VL	SC	ML	2500	IU	10	11/20/2006	12/01/2014						
62856-0500-10		J1645		10/10/2006	03/31/2015	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (27GX1/2",10X0.2ML,PF) 5000 IU/0.2 ML	0.2	ML	SR	SC	ML	2500	IU	10	10/10/2006	03/31/2015						
62856-0750-10		J1645		02/06/2007	02/02/2015	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED) 7500 IU/0.3 ML	0.3	ML	SR	SC	ML	2500	IU	10	02/06/2007	02/02/2015						
62927-0621-04		Q0177		01/01/2002	12/17/2015	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (BANANA) 25 MG/5 ML	120	ML	EA	PO	ML	25	MG	0.2	01/01/2002	12/17/2015						
62927-0621-16		Q0177		01/01/2002	12/17/2015	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (BANANA) 25 MG/5 ML	480	ML	EA	PO	ML	25	MG	0.2	01/01/2002	12/17/2015						
62991-1003-02		J7604		01/01/2008	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
62991-1003-02	KO	J7604	KO	01/01/2008	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
62991-1003-03		J7604		01/01/2008	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
62991-1003-03	KO	J7604	KO	01/01/2008	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
62991-1003-04		J7604		01/01/2008	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
62991-1003-04	KO	J7604	KO	01/01/2008	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1	EA	BO	NA	GM	1	GM	1	01/01/2008	99/99/9999						
62991-1004-01		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	01/01/2006	99/99/9999						
62991-1004-02		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	01/01/2006	99/99/9999						
66794-0160-02		J2274		07/23/2018	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10 MG	MITIGO (SINGLE USE,PF) 10 MG/1 ML	20	ML	VL	IJ	ML	10	MG	1	07/23/2018	99/99/9999						
66794-0162-02		J2274		07/27/2018	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10 MG	MITIGO (SINGLE USE,PF) 25 MG/1 ML	20	ML	VL	IJ	ML	10	MG	2.5	07/27/2018	99/99/9999						
62991-1013-01		J0475		01/01/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
66993-0039-01		J1729		08/09/2018	99/99/9999	INJECTION, HYDROXYPROGESTERONE CAPROATE, NOT OTHERWISE SPECIFIED, 10 MG	HYDROXYPROGESTERONE CAPROATE (MDV) 250 MG/1 ML	5	ML	VL	IM	ML	10	MG	25	08/09/2018	99/99/9999							
67457-0397-99		J2780		08/17/2018	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE (10X2ML,SDV,USP) 25 MG/1 ML	2	ML	VL	IJ	ML	25	MG	1	08/17/2018	99/99/9999							
67457-0398-62		J2780		08/17/2018	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE (SDV,USP) 25 MG/1 ML	6	ML	VL	IJ	ML	25	MG	1	08/17/2018	99/99/9999							
67457-0843-30		J2020		07/31/2018	99/99/9999	INJECTION, LINEZOLID, 200 MG	LINEZOLID (10X300ML BAGS,PF) 600 MG/300 ML	300	ML	BG	IV	ML	200	MG	0.01	07/31/2018	99/99/9999							
70710-1377-01		J0330		07/18/2018	99/99/9999	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	SUCCINYLCHOLINE CHLORIDE (MDV, INNER PACK,STERILE) 20 MG/1 ML	10	ML	VL	IJ	ML	20	MG	1	07/18/2018	99/99/9999							
62991-1013-02		J0475		01/01/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2002	99/99/9999							
62991-1013-03		J0475		01/01/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2002	99/99/9999							
62991-1013-04		J0475		09/15/2003	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	09/15/2003	99/99/9999							
62991-1021-02		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	BENZOCAINE (U.S.P./N.F.)	1	EA	BO	NA	GM	1	EA	1	01/01/2002	99/99/9999							
62991-1021-04		J3490		09/15/2003	99/99/9999	UNCLASSIFIED DRUGS	BENZOCAINE (U.S.P.)	1	EA	BO	NA	GM	1	EA	1	09/15/2003	99/99/9999							
62991-1023-02		J7624		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1023-02	KO	J7624	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1023-03		J7624		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1023-03	KO	J7624	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1024-01		J7624		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1024-01	KO	J7624	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1024-02		J7624		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1024-02	KO	J7624	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1024-04		J7624		09/15/2003	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999							
62991-1024-04	KO	J7624	KO	09/15/2003	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999							
62991-1024-05		J7624		09/15/2003	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999							
62991-1024-05	KO	J7624	KO	09/15/2003	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	BETAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999							
62991-1038-01		J7632		01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1038-01	KO	J7632	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1038-02		J7632		01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1038-02	KO	J7632	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1038-03		J7632		01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1038-03	KO	J7632	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1038-04		J7632		01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1038-04	KO	J7632	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM (U.S.P.)	1	EA	BO	NA	GM	10	MG	100	01/01/2008	99/99/9999							
62991-1039-02		J3420		01/01/2002	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	1	EA	BO	NA	GM	1000	MCG	1000	01/01/2002	99/99/9999							
62991-1039-03		J3420		01/01/2002	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	1	EA	BO	NA	GM	1000	MCG	1000	01/01/2002	99/99/9999							
62991-1041-02		J7638		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1041-02	KO	J7638	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1041-03		J7638		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1041-03	KO	J7638	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1041-04		J7638		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1041-04	KO	J7638	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1047-02		J1200		01/01/2002	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (U.S.P.)	1	EA	VL	NA	GM	50	MG	20	01/01/2002	99/99/9999							
70860-0603-82		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM-SODIUM CHLORIDE (PF,LATEX-FREE) 1000 MG/100 ML-0.75%	100	ML	BG	IV	ML	10	MG	1	06/13/2018	99/99/9999							
62991-1051-02		J1435		01/01/2002	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999							
62991-1051-03		J1435		09/15/2003	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999							
62991-1051-04		J1435		09/15/2003	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
62991-1072-01		J7699		09/01/2002	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1 EA	BO NA GM				1 EA		1	09/01/2002	99/99/9999						
62991-1072-02		J7699		09/01/2002	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1 EA	BO NA GM				1 EA		1	09/01/2002	99/99/9999						
70860-0604-82		J1953		06/13/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM-SODIUM CHLORIDE (PF,LATEX-FREE) 1500 MG/100 ML-0.54%	100 ML	BG IV ML				10 MG		1.5	06/13/2018	99/99/9999						
43975-0307-10		None		04/05/2018	99/99/9999	CYCLOPHOSPHAMIDE, 25 MG, ORAL	CYCLOPHOSPHAMIDE 25 MG	100 EA	BO PO EA				25 MG		1	04/05/2018	99/99/9999						
47335-0890-72		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE (3X5,HARD GELATIN) 5 MG	15 EA	ST PO EA				5 MG		1	07/11/2018	99/99/9999						
70655-0002-10		J1450		08/31/2018	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (PF,LATEX-FREE) 200 MG/100 ML	100 ML	BX IV ML				200 MG		0.01	08/31/2018	99/99/9999						
60505-6144-04		J0692		03/15/2018	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME NOVAPLUS 1 GM	10 EA	VL UJ EA				500 MG		2	03/15/2018	99/99/9999						
47335-0890-74		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE (1X5,HARD GELATIN) 5 MG	5 EA	ST PO EA				5 MG		1	07/11/2018	99/99/9999						
47335-0891-72		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (3X5,HARD GELATIN) 20 MG	15 EA	ST PO EA				20 MG		1	07/11/2018	99/99/9999						
70655-0088-06		J1450		08/31/2018	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (PF,LATEX-FREE) 400 MG/200 ML	200 ML	BG IV ML				200 MG		0.01	08/31/2018	99/99/9999						
70655-0088-10		J1450		08/31/2018	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (PF,LATEX-FREE) 400 MG/200 ML	200 ML	BG IV ML				200 MG		0.01	08/31/2018	99/99/9999						
70710-1525-09		J9050		09/14/2018	99/99/9999	INJECTION, CARMUSTINE, 100 MG	CARMUSTINE (LYOPHILIZED) 100 MG	1 EA	VL IV EA				100 MG		1	09/14/2018	99/99/9999						
71288-0106-10		J9040		10/01/2018	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN (SDV,PF,LATEX-FREE) 15 U	1 EA	VL UJ EA				15 U		1	10/01/2018	99/99/9999						
62991-1095-01		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P., B.P.)	1 EA	BO NA GM				10 MG		100	01/01/2004	99/99/9999						
62991-1095-02		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P., B.P.)	1 EA	BO NA GM				10 MG		100	01/01/2004	99/99/9999						
62991-1095-03		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P., B.P.)	1 EA	BO NA GM				10 MG		100	01/01/2004	99/99/9999						
62991-1095-04		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P., B.P.)	1 EA	BO NA GM				10 MG		100	01/01/2004	99/99/9999						
71288-0107-20		J9040		10/01/2018	99/99/9999	INJECTION, BLEOMYCIN SULFATE, 15 UNITS	BLEOMYCIN (SDV,PF,LATEX-FREE) 30 U	1 EA	VL UJ EA				15 U		2	10/01/2018	99/99/9999						
71288-0109-20		J9100		11/05/2018	99/99/9999	INJECTION, CYTARABINE, 100 MG	CYTARABINE (SDV,PF,LATEX-FREE) 100 MG/1 ML	20 ML	VL UJ ML				100 MG		1	11/05/2018	99/99/9999						
62991-1095-06		J2001		04/01/2008	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (USP)	1 EA	BO NA GM				10 MG		100	04/01/2008	99/99/9999						
62991-1108-01		J2760		01/01/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM				5 MG		200	01/01/2002	99/99/9999						
62991-1108-02		J2760		01/01/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM				5 MG		200	01/01/2002	99/99/9999						
62991-1108-03		J2760		09/15/2003	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM				5 MG		200	09/15/2003	99/99/9999						
62991-1108-04		J2760		09/15/2003	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO NA GM				5 MG		200	09/15/2003	99/99/9999						
62991-1122-02		Q0165		01/01/2002	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	1 EA	BO NA GM				10 MG		100	01/01/2002	12/31/2013						
62991-1124-02		J2675		01/01/2002	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,MICRONIZED)	1 EA	BO NA GM				50 MG		20	01/01/2002	99/99/9999						
62991-1124-03		J2675		10/01/2007	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO NA GM				50 MG		20	10/01/2007	99/99/9999						
62991-1124-05		J2675		10/01/2007	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO NA GM				50 MG		20	10/01/2007	99/99/9999						
62991-1125-01		J2550		01/01/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	BO NA GM				50 MG		20	01/01/2002	99/99/9999						
62991-1125-02		J2550		01/01/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	BO NA GM				50 MG		20	01/01/2002	99/99/9999						
62991-1125-04		J2550		01/01/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	BO NA GM				50 MG		20	01/01/2002	99/99/9999						
62991-1128-02		J0270		09/15/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO NA GM				1.25 MCG		800000	09/15/2003	99/99/9999						
62991-1128-06		J0270		09/15/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO NA GM				1.25 MCG		800000	09/15/2003	99/99/9999						
62991-1128-07		J0270		09/15/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO NA GM				1.25 MCG		800000	09/15/2003	99/99/9999						
62991-1128-08		J0270		09/15/2003	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO NA GM				1.25 MCG		800000	09/15/2003	99/99/9999						
62991-1130-02		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1 EA	BO NA GM				100 MG		10	01/01/2004	99/99/9999						
62991-1130-03		J3415		01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1 EA	BO NA GM				100 MG		10	01/01/2004	99/99/9999						
62991-1132-01		J2780		09/15/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO NA GM				25 MG		40	09/15/2003	99/99/9999						
62991-1132-02		J2780		09/15/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO NA GM				25 MG		40	09/15/2003	99/99/9999						
62991-1132-03		J2780		09/15/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO NA GM				25 MG		40	09/15/2003	99/99/9999						
62991-1132-04		J2780		09/15/2003	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO NA GM				25 MG		40	09/15/2003	99/99/9999						
62991-1133-01		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	RIFAMPIN (U.S.P.)	1 EA	BO NA GM				1 EA		1	01/01/2002	99/99/9999						
62991-1133-02		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	RIFAMPIN (U.S.P.)	1 EA	BO NA GM				1 EA		1	01/01/2002	99/99/9999						
62991-1133-04		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	RIFAMPIN (U.S.P.)	1 EA	BO NA GM				1 EA		1	01/01/2002	99/99/9999						
62991-1152-01		J7681		01/01/2002	99/99/9999	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO NA GM				1 MG		1000	01/01/2002	99/99/9999						
62991-1152-01	KO	J7681	KO	01/01/2002	99/99/9999	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO NA GM				1 MG		1000	01/01/2002	99/99/9999						
62991-1152-02		J7681		01/01/2002	99/99/9999	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO NA GM				1 MG		1000	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
62991-1152-02	KO	J7681	KO	01/01/2002	99/99/9999	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
62991-1156-01		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., BP EP, MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
62991-1156-01	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., BP EP, MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
62991-1156-02		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., BP EP, MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
62991-1156-02	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., BP EP, MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
62991-1156-03		J7684		01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., BP EP, MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
62991-1156-03	KO	J7684	KO	01/01/2002	99/99/9999	TRIAMCINOLONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TRIAMCINOLONE ACETONIDE (U.S.P., BP EP, MICRONIZED)	1	EA	BO	NA	GM	1	MG	1000	01/01/2002	99/99/9999						
62991-1173-02		J0285		01/01/2002	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P., ORAL GRADE)	1	EA	BO	NA	GM	50	MG	20	01/01/2008	99/99/9999	01/01/2002	09/01/2004	20			
62991-1173-04		J0285		01/01/2002	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P., ORAL GRADE)	1	EA	BO	NA	GM	50	MG	20	01/01/2008	99/99/9999						
62991-1173-05		J0285		01/01/2008	99/99/9999	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (USP)	1	EA	BO	NA	GM	50	MG	20	01/01/2008	99/99/9999	01/01/2002	09/01/2004	20			
62991-1179-03		J7627		01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE MICRONIZED (EP)	1	EA	JR	NA	GM	0.5	MG	2000	01/01/2006	99/99/9999						
62991-1179-03	KO	J7627	KO	01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE MICRONIZED (EP)	1	EA	JR	NA	GM	0.5	MG	2000	01/01/2006	99/99/9999						
62991-1179-05		J7627		01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE MICRONIZED (EP)	1	EA	JR	NA	GM	0.5	MG	2000	01/01/2006	99/99/9999						
62991-1179-05	KO	J7627	KO	01/01/2006	99/99/9999	BUDESONIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE MICRONIZED (EP)	1	EA	JR	NA	GM	0.5	MG	2000	01/01/2006	99/99/9999						
62991-1206-01		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (U.S.P., MICRONIZED)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	12/31/2015						
62991-1206-02		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (U.S.P., MICRONIZED)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	12/31/2015						
62991-1257-01		J7510		01/01/2002	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	01/01/2002	99/99/9999						
62991-1257-02		J7510		09/15/2003	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE ANHYDROUS (U.S.P., MICRO)	1	EA	NA	NA	GM	5	MG	200	09/15/2003	99/99/9999						
62991-1351-02		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE	1	EA	BO	NA	GM	300	MG	3.33333	01/01/2007	99/99/9999						
62991-1351-02	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE	1	EA	BO	NA	GM	300	MG	3.33333	01/01/2007	99/99/9999						
62991-1351-03		J7685		01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE	1	EA	BO	NA	GM	300	MG	3.33333	01/01/2007	99/99/9999						
62991-1351-03	KO	J7685	KO	01/01/2007	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCIN SULFATE	1	EA	BO	NA	GM	300	MG	3.33333	01/01/2007	99/99/9999						
62991-1352-01		J3490		01/01/2007	99/99/9999	UNCLASSIFIED DRUGS	HYALURONIC ACID	1	EA	BO	NA	GM	1	EA	1	01/01/2007	99/99/9999						
62991-1352-02		J3490		01/01/2007	99/99/9999	UNCLASSIFIED DRUGS	HYALURONIC ACID	1	EA	NA	NA	GM	1	EA	1	01/01/2007	99/99/9999						
62991-1352-04		J3490		01/01/2007	99/99/9999	UNCLASSIFIED DRUGS	HYALURONIC ACID	1	EA	BO	NA	GM	1	EA	1	01/01/2007	99/99/9999						
62991-1382-01		J3350		01/01/2002	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P./N.F.)	1	EA	BO	NA	GM	40	GM	0.025	01/01/2002	99/99/9999						
62991-1412-01		J3150		09/01/2002	01/09/2013	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE MICRONIZED (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	09/01/2002	01/09/2013						
62991-1412-02		J3150		09/01/2002	11/01/2012	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE MICRONIZED (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	09/01/2002	11/01/2012						
62991-1412-03		J3150		09/01/2002	12/21/2012	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE MICRONIZED (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	09/01/2002	12/21/2012						
62991-1422-01		J0735		09/15/2003	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999						
62991-1422-02		J0735		09/15/2003	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	09/15/2003	99/99/9999						
62991-1486-02		J9190		09/15/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (U.S.P.)	1	EA	BO	NA	GM	500	MG	2	09/15/2003	99/99/9999						
62991-1486-03		J9190		09/15/2003	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (U.S.P.)	1	EA	BO	NA	GM	500	MG	2	09/15/2003	99/99/9999						
62991-1513-01		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	99/99/9999						
62991-1513-02		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	99/99/9999						
62991-1513-03		J0364		01/01/2007	99/99/9999	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1	EA	BO	NA	GM	1	MG	1000	01/01/2007	99/99/9999						
62991-1530-02		J0520		09/15/2003	99/99/9999	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/15/2003	99/99/9999						
62991-1530-03		J0520		09/15/2003	99/99/9999	INJECTION, BETHANECHOL CHLORIDE, MYOTONACHOL OR URECHOLINE, UP TO 5 MG	BETHANECHOL CHLORIDE (U.S.P.)	1	EA	BO	NA	GM	5	MG	200	09/15/2003	99/99/9999						
62991-1533-01		J7516		09/15/2003	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORINE (U.S.P., A)	1	EA	BO	NA	GM	250	MG	4	09/15/2003	99/99/9999						
62991-1533-02		J7516		09/15/2003	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORINE (U.S.P., A)	1	EA	BO	NA	GM	250	MG	4	09/15/2003	99/99/9999						
62991-1533-05		J7516		01/01/2008	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORINE (U.S.P., A)	1	EA	NA	NA	GM	250	MG	4	01/01/2008	99/99/9999						
62991-1568-01		J2150		09/15/2003	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	1	EA	BO	NA	GM	50	ML	0.08	01/01/2008	99/99/9999	09/15/2003	10/01/2007	0.08			
62991-1583-01		J0592		09/15/2003	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE	1	EA	BO	NA	GM	0.1	MG	10000	09/15/2003	99/99/9999						
62991-1583-02		J0592		09/15/2003	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE	1	EA	BO	NA	GM	0.1	MG	10000	09/15/2003	99/99/9999						
62991-1583-03		J0592		09/15/2003	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE	1	EA	BO	NA	GM	0.1	MG	10000	09/15/2003	99/99/9999						
62991-1635-02		J1030		09/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1	EA	BO	NA	GM	40	MG	25	09/01/2002	99/99/9999						
62991-1635-03		J1030		09/01/2002	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1	EA	BO	NA	GM	40	MG	25	09/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
62991-1635-04		J1030		09/15/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	40 MG	25	09/15/2003	99/99/9999									
62991-1635-05		J1030		09/15/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	40 MG	25	09/15/2003	99/99/9999									
62991-1635-06		J1030		09/15/2003	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	40 MG	25	09/15/2003	99/99/9999									
62991-1685-01		J3490		09/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM	1 EA	1	09/01/2002	99/99/9999									
62991-1685-02		J3490		09/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM	1 EA	1	09/01/2002	99/99/9999									
62991-1685-03		J3490		09/01/2002	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM	1 EA	1	09/01/2002	99/99/9999									
62991-1692-01		J2650		09/01/2002	99/99/9999	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED	1 EA	BO	NA	GM	1 ML	20	09/01/2002	99/99/9999									
62991-1692-02		J2650		09/01/2002	99/99/9999	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED	1 EA	BO	NA	GM	1 ML	20	09/01/2002	99/99/9999									
62991-1692-03		J2650		09/01/2002	99/99/9999	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED	1 EA	BO	NA	GM	1 ML	20	09/01/2002	99/99/9999									
62991-1707-01		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM	100 MG	10	01/01/2002	12/31/2014									
62991-1707-02		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM	100 MG	10	01/01/2002	12/31/2014									
62991-1707-03		J1070		01/01/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM	100 MG	10	01/01/2002	12/31/2014									
62991-2003-02		J0280		01/01/2002	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1 EA	BO	NA	GM	250 MG	4	01/01/2002	99/99/9999									
62991-2003-03		J0280		01/01/2002	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1 EA	BO	NA	GM	250 MG	4	01/01/2002	99/99/9999									
62991-2004-02		J1320		01/01/2002	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	1 EA	BO	NA	GM	20 MG	50	01/01/2002	99/99/9999									
62991-2004-03		J1320		01/01/2002	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	1 EA	BO	NA	GM	20 MG	50	01/01/2002	99/99/9999									
62991-2022-02		J7638		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000	01/01/2002	99/99/9999									
62991-2022-02	KO	J7638	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000	01/01/2002	99/99/9999									
62991-2022-04		J7638		01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000	01/01/2002	99/99/9999									
62991-2022-04	KO	J7638	KO	01/01/2002	99/99/9999	THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE (U.S.P.,MICRONIZED)	1 EA	BO	NA	GM	1 MG	1000	01/01/2002	99/99/9999									
62991-2026-02		J3520		01/01/2002	99/99/9999	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM (U.S.P.N.F.)	1 EA	BO	NA	GM	150 MG	6.66666	01/01/2002	99/99/9999									
62991-2026-03		J3520		01/01/2002	99/99/9999	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM (U.S.P.N.F.)	1 EA	BO	NA	GM	150 MG	6.66666	01/01/2002	99/99/9999									
62991-2026-04		J3520		09/15/2003	99/99/9999	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM (DIHYDRATE)	1 EA	BO	NA	GM	150 MG	6.66666	09/15/2003	99/99/9999									
62991-2031-02		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200	01/01/2002	99/99/9999									
62991-2031-03		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200	01/01/2002	99/99/9999									
62991-2031-04		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	1 EA	BO	NA	GM	5 MG	200	01/01/2002	99/99/9999									
4735-0891-74	None			07/11/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (1X5,HARD GELATIN) 20 MG	5 EA	ST	PO	EA	20 MG	1	07/11/2018	99/99/9999									
62991-2042-02		J2765		01/01/2002	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100	01/01/2002	99/99/9999									
62991-2042-03		J2765		01/01/2002	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100	01/01/2002	99/99/9999									
62991-2068-02		J3411		01/01/2004	99/99/9999	INJECTION, THIAMINE HCL, 100 MG	THIAMINE HYDROCHLORIDE (1X100GM, USP)	1 EA	BO	NA	GM	100 MG	10	10/01/2007	99/99/9999			01/01/2004	09/01/2004	10				
62991-2068-03		J3411		01/01/2004	99/99/9999	INJECTION, THIAMINE HCL, 100 MG	THIAMINE HYDROCHLORIDE (1X500GM, USP)	1 EA	BO	NA	GM	100 MG	10	10/01/2007	99/99/9999			01/01/2004	09/01/2004	10				
62991-2068-04		J3411		10/01/2007	99/99/9999	INJECTION, THIAMINE HCL, 100 MG	THIAMINE HYDROCHLORIDE (1X100GM,USP)	1 EA	NA	NA	GM	100 MG	10	10/01/2007	99/99/9999									
47781-0605-94		J9045		04/02/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	45 ML	VL	IV	ML	50 MG	0.2	04/02/2018	99/99/9999									
62991-2150-01		J3140		09/01/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	50 MG	20	09/01/2002	12/31/2014									
62991-2150-02		J3140		09/01/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	50 MG	20	09/01/2002	12/31/2014									
62991-2150-03		J3140		09/01/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	50 MG	20	09/01/2002	12/31/2014									
62991-2150-04		J3140		09/01/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	50 MG	20	09/01/2002	12/31/2014									
62991-2184-02		J2675		09/01/2002	03/05/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO	NA	GM	50 MG	20	09/01/2002	03/05/2013									
62991-2184-03		J2675		09/01/2002	03/05/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO	NA	GM	50 MG	20	09/01/2002	03/05/2013									
62991-2184-04		J2675		09/01/2002	02/06/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO	NA	GM	50 MG	20	09/01/2002	02/06/2013									
62991-2501-01		J3490		09/15/2003	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P., 24)	1 EA	BO	NA	GM	1 EA	1	09/15/2003	99/99/9999									
62991-2501-02		J3490		09/15/2003	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P., 24)	1 EA	BO	NA	GM	1 EA	1	09/15/2003	99/99/9999									
62991-2516-01		J7640		01/01/2006	99/99/9999	FORMOTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 12 MICROGRAMS	FORMOTEROL FUMARATE	1 EA	BO	NA	GM	12 MCG	83333.33	01/01/2006	99/99/9999									
62991-2516-01	KO	J7640	KO	01/01/2006	99/99/9999	FORMOTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 12 MICROGRAMS	FORMOTEROL FUMARATE	1 EA	BO	NA	GM	12 MCG	83333.33	01/01/2006	99/99/9999									
62991-2516-03		J7640		01/01/2006	99/99/9999	FORMOTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 12 MICROGRAMS	FORMOTEROL FUMARATE	1 EA	BO	NA	GM	12 MCG	83333.33	01/01/2006	99/99/9999									
62991-2516-03	KO	J7640	KO	01/01/2006	99/99/9999	FORMOTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 12 MICROGRAMS	FORMOTEROL FUMARATE	1 EA	BO	NA	GM	12 MCG	83333.33	01/01/2006	99/99/9999									
62991-2562-01		J1835		11/01/2005	99/99/9999	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE	1 EA	NA	NA	GM	50 MG	20	11/01/2005	99/99/9999									
62991-2562-02		J1835		11/01/2005	99/99/9999	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE	1 EA	NA	NA	GM	50 MG	20	11/01/2005	99/99/9999									
62991-2562-03		J1835		11/01/2005	99/99/9999	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE	1 EA	NA	NA	GM	50 MG	20	11/01/2005	99/99/9999									
62991-2577-02		J0456		10/01/2007	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN DIHYDRATE (1X100																	

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
62991-2577-03		J0456		10/01/2007	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN DIHYDRATE (1X500GM, USP)	1 EA	NA	NA	GM		500 MG		2	10/01/2007	99/99/9999						
62991-2599-01		J2405		01/01/2006	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HYDROCHLORIDE (1X100GM)	1 EA	BO	NA	GM		1 MG		1000	01/01/2006	99/99/9999						
62991-2599-02		J2405		01/01/2006	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HYDROCHLORIDE (1X1000GM)	1 EA	BO	NA	GM		1 MG		1000	01/01/2006	99/99/9999						
62991-2664-01		J7507		10/01/2007	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (1X100MG)	0.1 GM	NA	NA	GM		1 MG		1000	10/01/2007	99/99/9999						
62991-2664-02		J7507		10/01/2007	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (1X500MG)	0.5 GM	NA	NA	GM		1 MG		1000	10/01/2007	99/99/9999						
62991-2664-03		J7507		10/01/2007	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (1X1GM)	1 EA	NA	NA	GM		1 MG		1000	10/01/2007	99/99/9999						
62991-2664-04		J7507		10/01/2007	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (1X5GM)	5 GM	NA	NA	GM		1 MG		1000	10/01/2007	99/99/9999						
62991-2707-02		J1956		01/01/2008	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN	1 EA	BO	NA	GM		250 MG		4	01/01/2008	99/99/9999						
62991-2707-03		J1956		01/01/2008	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN	1 EA	BO	NA	GM		250 MG		4	01/01/2008	99/99/9999						
63004-7731-01		J0800		01/01/2002	01/06/2013	INJECTION, CORTICOTROPIN, UP TO 40 UNITS	H.P. ACTHAR (M.D.V.) 80 U/ML VELCADE (10ML SDV.LYOPHILIZED) 3.5 MG	5 ML	VL	IJ	ML		40 U		2	01/01/2002	01/06/2013						
63020-0049-01		J9041		01/01/2005	99/99/9999	INJECTION, BORTEZOMIB (VELCADE), 0.1 MG	TEMOZOLOMIDE (3X5,HARD GELATIN) 100 MG	1 EA	VL	IV	EA		0.1 MG		35	01/01/2005	99/99/9999						
47335-0892-72		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE (3X5,HARD GELATIN) 100 MG	15 EA	ST	PO	EA		100 MG		1	07/11/2018	99/99/9999						
63275-1025-04		J2271		12/03/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	12/03/2002	12/31/2014						
63275-1100-05		J2271		12/03/2002	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	12/03/2002	12/31/2014						
63275-1200-01		J1960		12/03/2002	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	1 EA	BO	NA	GM		2 MG		500	12/03/2002	99/99/9999						
63275-1200-02		J1960		12/03/2002	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	1 EA	BO	NA	GM		2 MG		500	12/03/2002	99/99/9999						
63275-1200-04		J1960		12/03/2002	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	1 EA	BO	NA	GM		2 MG		500	12/03/2002	99/99/9999						
63275-1200-07		J1960		12/03/2002	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	1 EA	BO	NA	GM		2 MG		500	12/03/2002	99/99/9999						
63275-2001-01		J1170		12/03/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	JR	NA	GM		4 MG		250	12/03/2002	99/99/9999						
63275-2005-02		J1170		12/03/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	12/03/2002	99/99/9999						
63275-2010-03		J1170		12/03/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	12/03/2002	99/99/9999						
63275-2100-05		J1170		12/03/2002	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	12/03/2002	99/99/9999						
63275-2100-09		J1170		09/01/2003	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 EA	BO	NA	GM		4 MG		250	09/01/2003	99/99/9999						
63275-5100-01		J3010		12/03/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 EA	BO	NA	GM		0.1 MG		10000	12/03/2002	99/99/9999						
63275-5100-02		J3010		09/01/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 EA	BO	NA	GM		0.1 MG		10000	09/01/2002	99/99/9999						
63275-5100-06		J3010		12/03/2002	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 EA	BO	NA	GM		0.1 MG		10000	12/03/2002	99/99/9999						
63275-6200-01		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	01/01/2002	99/99/9999						
63275-6200-06		J3490		12/03/2002	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	12/03/2002	99/99/9999						
63275-6200-07		J3490		12/03/2002	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	12/03/2002	99/99/9999						
63275-6200-09		J3490		12/03/2002	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P.)	1 EA	BO	NA	GM		1 EA		1	12/03/2002	99/99/9999						
63275-7100-04		J2175		12/03/2002	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	12/03/2002	99/99/9999						
63275-7100-05		J2175		12/03/2002	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	1 EA	BO	NA	GM		100 MG		10	12/03/2002	99/99/9999						
63275-8100-03		J0745		12/03/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		30 MG		33.33333	12/03/2002	99/99/9999						
63275-8100-04		J0745		12/03/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		30 MG		33.33333	12/03/2002	99/99/9999						
63275-8100-05		J0745		12/03/2002	99/99/9999	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	1 EA	BO	NA	GM		30 MG		33.33333	12/03/2002	99/99/9999						
63275-9100-04		J1230		12/03/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	BO	NA	GM		10 MG		100	12/03/2002	99/99/9999						
63275-9100-05		J1230		12/03/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	BO	NA	GM		10 MG		100	12/03/2002	99/99/9999						
63275-9936-02		J1320		01/01/2007	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HYDROCHLORIDE (1X5GM, USP)	1 EA	BO	NA	GM		20 MG		50	01/01/2007	99/99/9999						
63275-9936-04		J1320		01/01/2007	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HYDROCHLORIDE (1X25GM, USP)	1 EA	BO	NA	GM		20 MG		50	01/01/2007	99/99/9999						
63275-9936-05		J1320		01/01/2007	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HYDROCHLORIDE (1X100GM, USP)	1 EA	BO	NA	GM		20 MG		50	01/01/2007	99/99/9999						
63275-9936-08		J1320		01/01/2007	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HYDROCHLORIDE (1X500GM, USP)	1 EA	BO	NA	GM		20 MG		50	01/01/2007	99/99/9999						
63275-9955-01		J2405		01/27/2005	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HCL	1 EA	BO	NA	GM		1 MG		1000	01/27/2005	99/99/9999						
63275-9955-06		J2405		01/27/2005	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HCL	1 EA	BO	NA	GM		1 MG		1000	01/27/2005	99/99/9999						
63275-9955-07		J2405		01/27/2005	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HCL	1 EA	BO	NA	GM		1 MG		1000	01/27/2005	99/99/9999						
63275-9958-01		J7507		09/01/2004	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS	1 EA	BO	NA	GM		1 MG		1000	09/01/2004	99/99/9999						
63275-9958-02		J7507		09/01/2004	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS	5 EA	BO	NA	GM		1 MG		1000	09/01/2004	99/99/9999						
63275-9958-06		J7507		09/01/2004	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS	0.1 GM	BO	NA	GM		1 MG		1000	09/01/2004	99/99/9999						
63275-9958-07		J7507		09/01/2004	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS	0.5 GM	BO	NA	GM		1 MG		1000	09/01/2004	99/99/9999						
63275-9960-01		J1450		05/01/2004	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE	1 EA	NA	NA	GM		200 MG		5	05/01/2004	99/99/9999						
63275-9960-02		J1450		05/01/2004	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE	1 EA	BO	NA	GM		200 MG		5	05/01/2004	99/99/9999						
63275-9960-04		J1450		05/01/2004	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE	1 EA	BO	NA	GM		200 MG		5	05/01/2004	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63275-9965-03		J0456		01/01/2007	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN DIHYDRATE (1X10GM, USP)	1 EA	BO	NA	GM	500 MG			2	01/01/2007	99/99/9999						
63275-9965-04		J0456		01/01/2007	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN DIHYDRATE (1X25GM, USP)	1 EA	BO	NA	GM	500 MG			2	01/01/2007	99/99/9999						
63275-9965-05		J0456		01/01/2007	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN DIHYDRATE (1X100GM, USP)	1 EA	BO	NA	GM	500 MG			2	01/01/2007	99/99/9999						
63275-9974-01		J0735		01/01/2003	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (BULK COMPOUND)	1 EA	JR	NA	GM	1 MG	1000		1000	01/01/2003	99/99/9999						
63275-9974-02		J0735		01/01/2003	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (BULK COMPOUND)	1 EA	JR	NA	GM	1 MG	1000		1000	01/01/2003	99/99/9999						
63275-9974-03		J0735		01/01/2003	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (BULK COMPOUND)	1 EA	JR	NA	GM	1 MG	1000		1000	01/01/2003	99/99/9999						
63275-9979-02		J2060		12/04/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO	NA	GM	2 MG	500		500	12/04/2002	99/99/9999						
63275-9979-04		J2060		12/04/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO	NA	GM	2 MG	500		500	12/04/2002	99/99/9999						
63275-9979-05		J2060		12/04/2002	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	1 EA	BO	NA	GM	2 MG	500		500	12/04/2002	99/99/9999						
63275-9981-05		J2675		12/04/2002	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO	NA	GM	50 MG	20		20	12/04/2002	99/99/9999						
63275-9981-08		J2675		12/04/2002	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO	NA	GM	50 MG	20		20	12/04/2002	99/99/9999						
63275-9981-09		J2675		12/04/2002	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED	1 EA	BO	NA	GM	50 MG	20		20	12/04/2002	99/99/9999						
63275-9982-04		J1070		12/04/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM	100 MG	10		10	12/04/2002	12/31/2014						
63275-9982-05		J1070		12/04/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM	100 MG	10		10	12/04/2002	12/31/2014						
63275-9982-09		J1070		12/04/2002	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO	NA	GM	100 MG	10		10	12/04/2002	12/31/2014						
63275-9983-04		J3140		12/04/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED	1 EA	JR	NA	GM	50 MG	20		20	12/04/2002	12/31/2014						
63275-9983-05		J3140		12/04/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED	1 EA	JR	NA	GM	50 MG	20		20	12/04/2002	12/31/2014						
63275-9983-08		J3140		12/04/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED	1 EA	JR	NA	GM	50 MG	20		20	12/04/2002	12/31/2014						
63275-9983-09		J3140		12/04/2002	12/31/2014	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED	1 EA	JR	NA	GM	50 MG	20		20	12/04/2002	12/31/2014						
63275-9986-01		J1435		12/04/2002	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	12/04/2002	99/99/9999						
63275-9986-02		J1435		12/04/2002	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	12/04/2002	99/99/9999						
63275-9986-04		J1435		12/04/2002	99/99/9999	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	12/04/2002	99/99/9999						
63275-9988-09		J0270		12/04/2002	99/99/9999	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	PROSTAGLANDIN E1 (U.S.P.)	1 EA	BO	NA	GM	1.25 MCG	800000		800000	12/04/2002	99/99/9999						
63275-9989-01		J2760		12/04/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		200	12/04/2002	99/99/9999						
63275-9989-06		J2760		12/04/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		200	12/04/2002	99/99/9999						
63275-9989-07		J2760		12/04/2002	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		200	12/04/2002	99/99/9999						
63275-9990-02		J2440		12/04/2002	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		16.66666	12/04/2002	99/99/9999						
63275-9990-04		J2440		12/04/2002	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		16.66666	12/04/2002	99/99/9999						
63275-9990-05		J2440		12/04/2002	99/99/9999	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		16.66666	12/04/2002	99/99/9999						
63275-9991-04		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL	1 EA	BO	NA	GM	10 MG	100		100	01/01/2004	99/99/9999						
63275-9991-05		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL	1 EA	BO	NA	GM	10 MG	100		100	01/01/2004	99/99/9999						
63275-9991-08		J2001		01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL	1 EA	BO	NA	GM	10 MG	100		100	01/01/2004	99/99/9999						
63275-9992-02		J0475		12/04/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		100	12/04/2002	99/99/9999						
63275-9992-04		J0475		12/04/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		100	12/04/2002	99/99/9999						
63275-9992-05		J0475		12/04/2002	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		100	12/04/2002	99/99/9999						
63275-9998-01		J7645		01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9998-01	KO	J7645	KO	01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9998-02		J7645		01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9998-02	KO	J7645	KO	01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9998-04		J7645		01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9998-04	KO	J7645	KO	01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9998-05		J7645		01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9998-05	KO	J7645	KO	01/01/2007	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9999-04		J7609		01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9999-04	KO	J7609	KO	01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9999-05		J7609		01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63275-9999-05	KO	J7609	KO	01/01/2007	99/99/9999	ALBUTEROL, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM	1 MG	1000		1000	01/01/2007	99/99/9999						
63304-0504-01		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100 EA	BO	PO	EA	1 EA	1		1	01/01/2002	99/99/9999						
63304-0505-01		J8499		01/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100 EA	BO	PO	EA	1 EA	1		1	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
63323-0010-02		J1580		01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (M.D.V.) 40 MG/ML	2	ML	VL	U	ML	80	MG	0.5	01/01/2002	99/99/9999							
63323-0010-20		J1580		01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE (M.D.V.) 40 MG/ML	20	ML	VL	U	ML	80	MG	0.5	01/01/2002	99/99/9999							
63323-0011-15		J0720		01/01/2002	99/99/9999	INJECTION, CHLORAMPHENICOL SODIUM SUCCINATE, UP TO 1 GM	CHLORAMPHENICOL SODIUM SUCCINATE (VIAL,PF) 1 GM	1	EA	VL	IV	GM	1	GM	1	01/01/2002	99/99/9999							
63323-0012-01		J2590		01/01/2002	99/99/9999	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN (VIAL,P.C.) 10 U/ML	1	ML	VL	IV	ML	10	U	1	01/01/2002	99/99/9999							
63323-0012-10		J2590		01/01/2002	99/99/9999	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN (M.D.V.) 10 U/ML	10	ML	VL	IV	ML	10	U	1	01/01/2002	99/99/9999							
63323-0012-12		J2590		01/28/2008	99/99/9999	INJECTION, OXYTOCIN, UP TO 10 UNITS	NOVAPLUS OXYTOCIN (25X1ML,USP) 10 U/ML	1	ML	VL	U	ML	10	U	1	01/28/2008	99/99/9999							
63323-0012-30		J2590		09/24/2007	99/99/9999	INJECTION, OXYTOCIN, UP TO 10 UNITS	OXYTOCIN (M.D.V.) 10 U/ML	30	ML	VL	IV	ML	10	U	1	09/24/2007	99/99/9999							
63323-0013-02		J3411		01/01/2004	99/99/9999	INJECTION, THIAMINE HCL, 100 MG	THIAMINE HCL (M.D.V.) 100 MG/ML	2	ML	VL	U	ML	100	MG	1	01/01/2004	99/99/9999							
63323-0017-10		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPFLUSH-10 (S.D.V.,PF) 10 U/ML	10	ML	VL	IV	ML	10	U	1	01/01/2002	99/99/9999							
63323-0024-25		J2150		01/01/2002	99/99/9999	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (FLIPOFF TOP,PF) 25% CHORIONIC GONADOTROPIN (M.D.V. W/DILUENT) 10000 U	50	ML	VL	IV	ML	50	ML	0.02	01/01/2002	99/99/9999							
63323-0025-10		J0725		01/01/2002	99/99/9999	INJECTION, CHORIONIC GONADOTROPIN, PER 1,000 USP UNITS	CHORIONIC GONADOTROPIN (M.D.V. W/DILUENT) 10000 U	1	EA	VL	IM	EA	1000	Units	10	01/01/2002	99/99/9999							
63323-0044-01		J3420		01/01/2002	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (M.D.V.) 1000 MCG/ML	1	ML	VL	IM	ML	1000	MCG	1	01/01/2002	99/99/9999							
63323-0047-10		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.) 5000 U/ML	10	ML	VL	U	ML	1000	U	5	01/01/2002	99/99/9999							
63323-0064-02		J3475		01/01/2002	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.,P.C.) 500 MG/ML	2	ML	VL	U	ML	500	MG	1	01/01/2002	99/99/9999							
63323-0064-10		J3475		01/01/2002	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.,P.C.,PF) 500 MG/ML	10	ML	VL	U	ML	500	MG	1	01/01/2002	99/99/9999							
63323-0064-20		J3475		01/01/2002	05/17/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.) 500 MG/ML	20	ML	VL	U	ML	500	MG	1	01/01/2002	05/17/2016							
63323-0064-50		J3475		01/01/2002	05/17/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.) 500 MG/ML	50	ML	VL	U	ML	500	MG	1	01/01/2002	05/17/2016							
63323-0088-61		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE CONCENTRATE (MAXIVAL,BULK PACK,PF) 23.4%	100	ML	VL	IV	ML	1	EA	1	01/01/2002	99/99/9999							
63323-0088-63		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE CONCENTRATE (MAXIVAL,BULK PACK,PF) 23.4%	200	ML	VL	IV	ML	1	EA	1	01/01/2002	99/99/9999							
63323-0101-61		J9000		08/06/2007	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HYDROCHLORIDE (USP,STERILE,MDV,PF) 2 MG/ML	100	ML	VL	IV	ML	10	MG	0.2	08/06/2007	99/99/9999							
63323-0104-05		J9181		01/01/2002	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V.) 20 MG/ML	5	ML	VL	IV	ML	10	MG	2	01/01/2002	99/99/9999							
63323-0104-25		J9181		01/01/2002	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V.) 20 MG/ML	25	ML	VL	IV	ML	10	MG	2	01/01/2002	99/99/9999							
63323-0104-50		J9181		01/01/2002	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (M.D.V.) 20 MG/ML	50	ML	VL	IV	ML	10	MG	2	01/01/2002	99/99/9999							
63323-0105-10		J0133		01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR SODIUM (VIAL,PF) 500 MG	1	EA	VL	IV	EA	5	MG	100	01/01/2006	99/99/9999							
63323-0113-10		J7676		01/01/2008	99/99/9999	PENTAMIDINE ISETHIONATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MG	PENTAM (S.D.V.,PF) 300 MG	1	EA	VL	U	EA	300	MG	1	01/01/2008	99/99/9999							
63323-0113-10	KO	J7676	KO	01/01/2008	99/99/9999	ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MG	PENTAM (S.D.V.,PF) 300 MG	1	EA	VL	U	EA	300	MG	1	01/01/2008	99/99/9999							
63323-0117-10		J9190		01/01/2002	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (S.D.V.,PF) 50 MG/ML	10	ML	VL	IV	ML	500	MG	0.1	01/01/2002	99/99/9999							
63323-0117-20		J9190		01/01/2002	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (S.D.V.,PF) 50 MG/ML	20	ML	VL	IV	ML	500	MG	0.1	01/01/2002	99/99/9999							
63323-0117-51		J9190		01/01/2002	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (BULK PACKAGE PF) 50 MG/ML	50	ML	VL	IV	ML	500	MG	0.1	01/01/2002	99/99/9999							
63323-0117-61		J9190		01/01/2002	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (BULK PACKAGE,PF) 50 MG/ML	100	ML	VL	IV	ML	500	MG	0.1	01/01/2002	99/99/9999							
63323-0119-08		J9150		01/01/2002	99/99/9999	INJECTION, DAUNORUBICIN, 10 MG	DAUNORUBICIN HCL (S.D.V.,PF) 20 MG	1	EA	VL	IV	EA	10	MG	2	01/01/2002	99/99/9999							
63323-0121-02		J9250		01/01/2002	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	2	ML	VL	U	ML	5	MG	5	01/01/2002	99/99/9999							
63323-0121-04		J9250		01/01/2002	02/03/2016	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	4	ML	VL	U	ML	5	MG	5	01/01/2002	02/03/2016							
63323-0121-08		J9250		01/01/2002	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	8	ML	VL	U	ML	5	MG	5	01/01/2002	99/99/9999							
63323-0121-10		J9250		01/01/2002	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (S.D.V.,PF) 25 MG/ML	10	ML	VL	U	ML	5	MG	5	01/01/2002	99/99/9999							
63323-0121-40		J9250		03/08/2002	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (VIAL,PF) 25 MG/ML	40	ML	VL	U	ML	5	MG	5	03/08/2002	99/99/9999							
63323-0122-50		J9260		01/01/2002	99/99/9999	METHOTREXATE SODIUM, 50 MG	METHOTREXATE SODIUM (S.D.V.,PF) 1 GM	1	EA	VL	U	EA	50	MG	20	01/01/2002	99/99/9999							
63323-0123-02		J9250		01/01/2002	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (VIAL) 25 MG/ML	2	ML	VL	U	ML	5	MG	5	01/01/2002	99/99/9999							
63323-0123-10		J9250		01/01/2002	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (VIAL) 25 MG/ML	10	ML	VL	U	ML	5	MG	5	01/01/2002	99/99/9999							
63323-0127-10		J9130		01/01/2002	99/99/9999	DACARBAZINE, 100 MG	DACARBAZINE (S.D.V.) 100 MG	1	EA	VL	IV	EA	100	MG	1	01/01/2002	99/99/9999							
63323-0132-10		J9293		03/17/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (USP,PF,LATEX-FREE) 2 MG/ML	10	ML	VL	IV	ML	5	MG	0.4	03/17/2006	99/99/9999							
63323-0132-12		J9293		03/17/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (USP,PF,LATEX-FREE) 2 MG/ML	12.5	ML	VL	IV	ML	5	MG	0.4	03/17/2006	99/99/9999							
63323-0132-15		J9293		03/17/2006	99/99/9999	INJECTION, MITOXANTRONE HYDROCHLORIDE, PER 5 MG	MITOXANTRONE (USP,PF,LATEX-FREE) 2 MG/ML	15	ML	VL	IV	ML	5	MG	0.4	03/17/2006	99/99/9999							
63323-0139-20		J7799		01/01/2002	02/15/2013	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (S.D.V.) 14.6%	20	ML	VL	IV	ML	1	EA	1	01/01/2002	02/15/2013							
63323-0139-40		J7799		01/01/2002	99/99/9999	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE (S.D.V.) 14.6%	40	ML	VL	IV	ML	1	EA	1	01/01/2002	99/99/9999							
63323-0140-10		J9065		09/13/2004	99/99/9999	INJECTION, CLADRIBINE, PER 1 MG	CLADRIBINE (S.D.V.,PF) 1 MG/ML	10	ML	VL	IV	ML	1	MG	1	09/13/2004	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63323-0142-10	J9208			07/25/2002	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (S.D.V.) 1 GM	1 EA	VL	IV	EA		1 GM		1	07/25/2002	99/99/9999						
63323-0142-12	J9208			11/18/2002	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (SDV) 1 GM	1 EA	VL	IV	EA		1 GM		1	11/18/2002	99/99/9999						
63323-0145-07	J9200			01/01/2002	99/99/9999	INJECTION, FLOXURIDINE, 500 MG	FLOXURIDINE 0.5 GM	1 EA	VL	IJ	EA		500 MG		1	01/01/2002	99/99/9999						
63323-0148-01	J9390			06/22/2005	99/99/9999	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (USP,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	06/22/2005	99/99/9999						
63323-0148-05	J9390			06/22/2005	99/99/9999	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE TARTRATE (USP,PF) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	06/22/2005	99/99/9999						
63323-0151-00	J9178			12/07/2007	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (1X100ML,PF) 2 MG/ML	100 ML	VL	IV	ML		2 MG		1	12/07/2007	99/99/9999						
63323-0151-25	J9178			12/07/2007	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HYDROCHLORIDE (1X25ML,PF) 2 MG/ML	25 ML	VL	IV	ML		2 MG		1	12/07/2007	99/99/9999						
63323-0161-01	J1885			01/01/2002	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 15 MG/ML	1 ML	VL	IJ	ML		15 MG		1	01/01/2002	99/99/9999						
63323-0162-01	J1885			01/01/2002	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 30 MG/ML	1 ML	VL	IJ	ML		15 MG		2	01/01/2002	99/99/9999						
63323-0162-02	J1885			01/01/2002	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (S.D.V.) 30 MG/ML	2 ML	VL	IM	ML		15 MG		2	01/01/2002	99/99/9999						
63323-0165-01	J1100			01/01/2002	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (VIAL) 4 MG/ML	1 ML	VL	IJ	ML		1 MG		4	01/01/2002	99/99/9999						
63323-0165-05	J1100			01/01/2002	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (M.D.V.) 4 MG/ML	5 ML	VL	IJ	ML		1 MG		4	01/01/2002	99/99/9999						
63323-0165-30	J1100			01/01/2002	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (M.D.V.) 4 MG/ML	30 ML	VL	IJ	ML		1 MG		4	01/01/2002	99/99/9999						
63323-0167-21	J9045			04/01/2004	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN 150 MG	1 EA	VL	IV	EA		50 MG		3	04/01/2004	99/99/9999						
63323-0172-45	J9045			04/28/2006	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (MDV,LATEX-FREE) 10 MG/ML	50 ML	VL	IV	ML		50 MG		0.2	04/28/2006	99/99/9999						
63323-0172-60	J9045			04/07/2006	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (600MG/60ML,LATEX-FREE) 10 MG/ML	60 ML	VL	IV	ML		50 MG		0.2	04/07/2006	99/99/9999						
63323-0173-02	J1580			01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE PEDIATRIC (PEDIATRIC S.D.V.,PF) 10 MG/ML	2 ML	VL	IJ	ML		80 MG		0.125	01/01/2002	99/99/9999						
63323-0180-01	J3415			01/01/2004	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (M.D.V.,AMBER) 100 MG/ML	1 ML	VL	IJ	ML		100 MG		1	01/01/2004	99/99/9999						
63323-0185-00	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.,TEAR TOP)	100 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0185-05	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.)	5 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0185-10	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.,P.C.)	10 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0185-20	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.,P.C.)	20 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0185-50	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	WATER FOR INJECTION (S.D.V.,P.C.,PF) SODIUM CHLORIDE (S.D.V.,TEAR TOP) 0.9%	50 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0186-00	J7050			01/01/2002	99/99/9999	INFUSION, NORMAL SALINE SOLUTION , 250 CC	SODIUM CHLORIDE (S.D.V.,P.C.) 0.9%	100 ML	VL	IV	ML		250 ML		0.004	01/01/2002	99/99/9999						
63323-0186-02	A4216			01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (S.D.V.,P.C.) 0.9%	2 ML	VL	IV	ML		10 ML		0.1	01/01/2007	99/99/9999						
63323-0186-10	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (S.D.V.,P.C.) 0.9%	10 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0186-20	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (S.D.V.,P.C.) 0.9%	20 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0187-30	J7799			01/01/2002	01/15/2013	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	SODIUM CHLORIDE CONCENTRATE (S.D.V.,PF) 23.4%	30 ML	VL	IV	ML		1 EA		1	01/01/2002	01/15/2013						
63323-0193-02	J9206			02/05/2008	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML,SINGLE DOSE) 20 MG/ML	2 ML	VL	IV	ML		20 MG		1	02/05/2008	99/99/9999						
63323-0193-05	J9206			02/05/2008	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML,SINGLE DOSE) 20 MG/ML	5 ML	VL	IV	ML		20 MG		1	02/05/2008	99/99/9999						
63323-0196-06	J9185			12/07/2007	99/99/9999	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE (USP) 50 MG	1 EA	VL	IV	EA		50 MG		1	12/07/2007	99/99/9999						
63323-0201-02	J2001			01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (S.D.V.,P.C.) 1%	2 ML	VL	EP	ML		10 MG		1	01/01/2004	99/99/9999						
63323-0201-10	J2001			01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (M.D.V.) 1%	10 ML	VL	EP	ML		10 MG		1	01/01/2004	99/99/9999						
63323-0202-02	J2001			01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (S.D.V.) 2%	2 ML	VL	IJ	ML		10 MG		2	01/01/2004	99/99/9999						
63323-0208-05	J2001			01/01/2004	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (S.D.V.,PF) 2%	5 ML	VL	IV	ML		10 MG		2	01/01/2004	99/99/9999						
63323-0221-10	J3370			01/01/2002	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (VIAL,PF) 500 MG	1 EA	VL	IV	EA		500 MG		1	01/01/2002	99/99/9999						
63323-0229-05	J2720			01/01/2002	99/99/9999	INJECTION, PROTAMINE SULFATE, PER 10 MG	PROTAMINE SULFATE (S.D.V.) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	01/01/2002	99/99/9999						
63323-0229-15	J2720			01/07/2008	99/99/9999	INJECTION, PROTAMINE SULFATE, PER 10 MG	NOVAPLUS PROTAMINE SULFATE (25X5ML,SDV,FLIPTOP,USP) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	01/07/2008	99/99/9999						
63323-0229-30	J2720			01/01/2002	99/99/9999	INJECTION, PROTAMINE SULFATE, PER 10 MG	PROTAMINE SULFATE (S.D.V.) 10 MG/ML	25 ML	VL	IV	ML		10 MG		1	01/01/2002	99/99/9999						
63323-0229-35	J2720			01/07/2008	99/99/9999	INJECTION, PROTAMINE SULFATE, PER 10 MG	NOVAPLUS PROTAMINE SULFATE (1X25ML,SDV,FLIPTOP,USP) 10 MG/ML	25 ML	VL	IV	ML		10 MG		1	01/07/2008	99/99/9999						
63323-0236-10	J0690			01/01/2002	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (VIAL,PF) 500 MG	1 EA	VL	IJ	EA		500 MG		1	01/01/2002	99/99/9999						
63323-0237-10	J0690			01/01/2002	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (VIAL) 1 GM	1 EA	VL	IJ	EA		500 MG		2	01/01/2002	99/99/9999						
63323-0237-65	J0690			01/01/2002	10/17/2016	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (P.B.,PF) 1 GM	1 EA	VL	IJ	EA		500 MG		2	01/01/2002	10/17/2016						
63323-0238-61	J0690			01/01/2002	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM (BULK PACKAGE,PF) 10 GM	1 EA	VL	IJ	EA		500 MG		20	01/01/2002	99/99/9999						
63323-0249-30	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	STERILE WATER BACTERIOSTATIC (M.D.V.)	30 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0255-03	J2920			09/22/2004	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 40 MG	METHYLPREDNISOLONE SODIUM SUCCINATE 40 MG	1 EA	VL	IJ	EA		40 MG		1	09/22/2004	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63323-0258-03	J2930			08/23/2004	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	METHYLPREDNISOLONE SODIUM SUCCINATE 125 MG	1 EA	VL	IJ	EA		125 MG		1	08/23/2004	99/99/9999						
63323-0259-30	A4216			01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (M.D.V.) 0.9% PROGESTERONE IN SESAME OIL (M.D.V.) 50 MG/ML	30 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0261-10	J2675			01/01/2002	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE IN SESAME OIL (M.D.V.) 50 MG/ML	10 ML	VL	IM	ML		50 MG		1	01/01/2002	99/99/9999						
63323-0262-01	J1644			01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.,P.C.) 5000 U/ML	1 ML	VL	IJ	ML		1000 U		5	01/01/2002	99/99/9999						
63323-0265-30	J2930			10/27/2004	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	METHYLPREDNISOLONE SODIUM SUCCINATE (PF) 1 GM	1 EA	VL	IJ	EA		125 MG		8	10/27/2004	99/99/9999						
63323-0269-20	J3490			02/21/2008	03/06/2013	UNCLASSIFIED DRUGS	DIPRIVAN (20X25ML) 10 MG/ML NOVAPLUS DIPRIVAN (25X20ML) 10 MG/ML	20 ML	VL	IV	ML		1 EA		1	02/21/2008	03/06/2013						
63323-0269-27	J3490			01/15/2008	09/07/2016	UNCLASSIFIED DRUGS	DIPRIVAN (20X50ML) 10 MG/ML	20 ML	VL	IV	ML		1 EA		1	01/15/2008	09/07/2016						
63323-0269-50	J3490			04/28/2008	99/99/9999	UNCLASSIFIED DRUGS	DIPRIVAN (20X50ML) 10 MG/ML NOVAPLUS DIPRIVAN (20X50ML) 10 MG/ML	50 ML	VL	IV	ML		1 EA		1	04/28/2008	99/99/9999						
63323-0269-57	J3490			03/05/2008	99/99/9999	UNCLASSIFIED DRUGS	DIPRIVAN (10X100ML) 10 MG/ML	100 ML	VL	IV	ML		1 EA		1	03/05/2008	99/99/9999						
63323-0269-65	J3490			03/06/2008	99/99/9999	UNCLASSIFIED DRUGS	NOVAPLUS DIPRIVAN (10X100ML, INFUSION) 10 MG/ML	100 ML	VL	IV	ML		1 EA		1	03/06/2008	99/99/9999						
63323-0269-67	J3490			02/01/2008	99/99/9999	UNCLASSIFIED DRUGS	FLUPHENAZINE DECANOATE (M.D.V.) 25 MG/ML	5 ML	VL	IJ	ML		25 MG		1	01/01/2002	99/99/9999						
63323-0272-05	J2680			01/01/2002	99/99/9999	INJECTION, FLUPHENAZINE DECANOATE, UP TO 25 MG	HEPARIN SODIUM (S.D.V.) 1000 U/ML	2 ML	VL	IJ	ML		1000 U		1	01/01/2002	99/99/9999						
63323-0276-02	J1644			01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (S.D.V.) 1000 U/ML	2 ML	VL	IJ	ML		1000 U		1	01/01/2002	99/99/9999						
63323-0278-10	J9360			01/01/2002	99/99/9999	INJECTION, VINBLASTINE SULFATE, 1 MG	VINBLASTINE SULFATE (M.D.V.) 1 MG/ML	10 ML	VL	IV	ML		1 MG		1	01/01/2002	99/99/9999						
63323-0280-02	J1940			01/01/2002	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (S.D.V.,AMBER) 10 MG/ML	2 ML	VL	IJ	ML		20 MG		0.5	01/01/2002	99/99/9999						
63323-0280-04	J1940			01/01/2002	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (S.D.V.,AMBER) 10 MG/ML	4 ML	VL	IJ	ML		20 MG		0.5	01/01/2002	99/99/9999						
63323-0280-10	J1940			01/01/2002	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (S.D.V.,AMBER) 10 MG/ML	10 ML	VL	IJ	ML		20 MG		0.5	01/01/2002	99/99/9999						
63323-0282-02	J3490			05/11/2007	99/99/9999	UNCLASSIFIED DRUGS	FUROSEMIDE (S.D.V.,AMBER) 10 MG/ML CLINDAMYCIN (SDV,USP,2MLX25) 150 MG/ML	2 ML	VL	IJ	ML		1 EA		1	05/11/2007	99/99/9999						
63323-0282-04	J3490			05/11/2007	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN (SDV,USP,4MLX25) 150 MG/ML	4 ML	VL	IJ	ML		1 EA		1	05/11/2007	99/99/9999						
63323-0282-06	J3490			05/11/2007	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN (SDV,USP,6MLX25) 150 MG/ML	6 ML	VL	IJ	ML		1 EA		1	05/11/2007	99/99/9999						
63323-0282-60	J3490			05/11/2007	99/99/9999	UNCLASSIFIED DRUGS	CLINDAMYCIN (USP) 150 MG/ML	60 ML	VL	IV	ML		1 EA		1	05/11/2007	99/99/9999						
63323-0284-20	J3370			01/01/2002	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (VIAL,PF) 1 GM VANCOMYCIN HCL (BULK PACKAGE,PF) 5 GM	1 EA	VL	IV	EA		500 MG		2	01/01/2002	99/99/9999						
63323-0295-61	J3370			01/01/2002	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (BULK PACKAGE,PF) 5 GM	1 EA	VL	IV	GM		500 MG		2	01/01/2002	99/99/9999						
63323-0303-51	J3260			01/01/2007	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (BULK VIAL,PF LATEX-FREE) 1.2 GM	6 EA	VL	IV	EA		80 MG		15	01/01/2007	99/99/9999						
63323-0303-55	J3260			01/01/2007	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE NOVAPLUS (BULK PKG,50ML VIAL X 6) 1.2 GM	6 EA	VL	IV	EA		80 MG		15	01/01/2007	99/99/9999						
63323-0305-02	J3260			04/05/2004	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (PEDIATRIC M.D.V.) 10 MG/ML	2 ML	VL	IJ	ML		80 MG		0.125	04/05/2004	99/99/9999						
63323-0306-02	J3260			04/05/2004	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (M.D.V.,LATEX-FREE) 40 MG/ML	2 ML	VL	IJ	ML		80 MG		0.5	04/05/2004	99/99/9999						
63323-0306-30	J3260			04/05/2004	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (M.D.V.,LATEX-FREE) 40 MG/ML	30 ML	VL	IJ	ML		80 MG		0.5	04/05/2004	99/99/9999						
63323-0307-51	J3260			04/05/2004	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (PHARMACY BULK PACKAGE) 40 MG/ML	50 ML	VL	IJ	ML		80 MG		0.5	04/05/2004	99/99/9999						
63323-0308-61	J1450			07/08/2004	11/14/2012	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (GLASS BOTTLE) 200 MG/100 ML	100 ML	VL	IV	ML		200 MG		0.01	07/08/2004	11/14/2012						
63323-0308-63	J1450			07/08/2004	11/14/2012	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (GLASS BOTTLE) 400 MG/200 ML	200 ML	VL	IV	ML		200 MG		0.01	07/08/2004	11/14/2012						
63323-0311-10	J0610			01/01/2002	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (S.D.V.) 100 MG/ML	10 ML	VL	IV	ML		10 ML		0.1	01/01/2002	99/99/9999						
63323-0311-50	J0610			01/01/2002	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (S.D.V.) 100 MG/ML	50 ML	VL	IV	ML		10 ML		0.1	01/01/2002	99/99/9999						
63323-0311-61	J0610			01/01/2002	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (MAXIVIAL,BULK PACK,PF) 100 MG/ML	100 ML	VL	IV	ML		10 ML		0.1	01/01/2002	99/99/9999						
63323-0311-63	J0610			01/01/2002	02/15/2013	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (MAXIVIAL,BULK PACK) 100 MG/ML	200 ML	VL	IV	ML		10 ML		0.1	01/01/2002	02/15/2013						
63323-0314-61	J3370			01/01/2002	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (BULK PACKAGE,PF) 10 GM	1 EA	VL	IV	GM		500 MG		2	01/01/2002	99/99/9999						
63323-0317-01	J1626			12/14/2007	99/99/9999	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (10X1ML,S.D.V,PF) 0.1 MG/ML	1 ML	VL	IV	ML		100 MCG		1	12/14/2007	99/99/9999						
63323-0325-10	J0133			01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR SODIUM (S.D.V.,PF) 50 MG/ML	10 ML	VL	IV	ML		5 MG		10	01/01/2006	99/99/9999						
63323-0325-20	J0133			01/01/2006	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR SODIUM (S.D.V.,PF) 50 MG/ML	20 ML	VL	IV	ML		5 MG		10	01/01/2006	99/99/9999						
63323-0326-20	J0692			03/17/2008	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP,10X1GM) 1 GM	1 EA	VL	IJ	EA		500 MG		2	03/17/2008	99/99/9999						
63323-0329-30	J3490			04/23/2004	99/99/9999	UNCLASSIFIED DRUGS	BACITRACIN (LATEX-FREE) 50000 U	1 EA	VL	IM	EA		1 EA		1	04/23/2004	99/99/9999						
47335-0892-74	None			07/11/2018	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE (1X5,HARD GELATIN) 100 MG	5 EA	ST	PO	EA		100 MG		1	07/11/2018	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
47335-0893-74		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE (1X5,HARD GELATIN) 250 MG	5 EA	ST	PO	EA		250 MG		1	07/11/2018	99/99/9999						
47335-0929-72		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (3X5,HARD GELATIN) 140 MG	15 EA	ST	PO	EA		20 MG		7	07/11/2018	99/99/9999						
47335-0929-74		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (1X5,HARD GELATIN) 140 MG	5 EA	ST	PO	EA		20 MG		7	07/11/2018	99/99/9999						
47335-0930-72		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (3X5,HARD GELATIN) 180 MG	15 EA	ST	PO	EA		20 MG		9	07/11/2018	99/99/9999						
70710-1377-02	J0330			07/18/2018	99/99/9999	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	(MDV,STERILE) 20 MG/1 ML	10 ML	VL	IJ	ML		20 MG		1	07/18/2018	99/99/9999						
63323-0344-10	J0696			02/16/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (S.D.V.) 250 MG	1 EA	VL	IJ	EA		250 MG		1	02/16/2006	99/99/9999						
63323-0345-10	J0696			02/16/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (S.D.V.) 500 MG	1 EA	VL	IJ	EA		250 MG		2	02/16/2006	99/99/9999						
63323-0346-10	J0696			02/16/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (S.D.V.) 1 GM	1 EA	VL	IJ	EA		250 MG		4	02/16/2006	99/99/9999						
63323-0347-20	J0696			02/16/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (S.D.V.) 2 GM	1 EA	VL	IJ	EA		250 MG		8	02/16/2006	99/99/9999						
63323-0348-61	J0696			02/16/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (BULK PACKAGE,1X100ML) 10 GM	1 EA	VL	IV	EA		250 MG		40	02/16/2006	99/99/9999						
70860-0112-15	J0290			08/01/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,PF,LATEX-FREE) 250 MG	10 EA	VL	IJ	EA		500 MG		0.5	08/01/2018	99/99/9999						
63323-0359-03	J1840			01/03/2003	01/31/2013	INJECTION, KANAMYCIN SULFATE, UP TO 500 MG	KANAMYCIN SULFATE 1 GM/3 ML	3 ML	VL	IJ	ML		500 MG		0.666	01/03/2003	01/31/2013						
63323-0365-01	J2354			04/13/2006	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (SDV,1MLX10,PF) 50 MCG/ML	1 ML	VL	IJ	ML		25 MCG		2	04/13/2006	99/99/9999						
63323-0366-01	J1240			07/01/2004	99/99/9999	INJECTION, DIMENHYDRINATE, UP TO 50 MG	DIMENHYDRINATE (VIAL) 50 MG/ML	1 ML	VL	IJ	ML		50 MG		1	07/01/2004	99/99/9999						
63323-0368-20	J0295			11/30/2005	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN/SULBACTAM 1 GM-0.5 GM	1 EA	VL	IJ	EA		1.5 GM		1	11/30/2005	99/99/9999						
63323-0369-20	J0295			11/30/2005	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN/SULBACTAM 2 GM-1 GM	1 EA	VL	IJ	EA		1.5 GM		2	11/30/2005	99/99/9999						
63323-0370-62	J0295			11/08/2006	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN AND SULBACTAM (USP,PHARMACY BULK PKG) 10 GM-5 GM	1 EA	VL	IV	EA		1.5 GM		10	11/08/2006	99/99/9999						
63323-0373-02	J2405			12/27/2006	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (SDV,25X2ML,PF) 2 MG/ML	2 ML	VL	IJ	ML		1 MG		2	12/27/2006	99/99/9999						
63323-0374-20	J2405			12/27/2006	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV) 2 MG/ML	20 ML	VL	IJ	ML		1 MG		2	12/27/2006	99/99/9999						
63323-0376-01	J2354			04/13/2006	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (SDV,1MLX10,PF) 100 MCG/ML	1 ML	VL	IJ	ML		25 MCG		4	04/13/2006	99/99/9999						
63323-0377-01	J2354			04/13/2006	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (SDV,1MLX10,PF) 500 MCG/ML	1 ML	VL	IJ	ML		25 MCG		20	04/13/2006	99/99/9999						
63323-0378-05	J2354			05/12/2006	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (MDV) 200 MCG/ML	5 ML	VL	IJ	ML		25 MCG		8	05/12/2006	99/99/9999						
63323-0379-05	J2354			05/12/2006	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS	OCTREOTIDE ACETATE (MDV) 1000 MCG/ML	5 ML	VL	IJ	ML		25 MCG		40	05/12/2006	99/99/9999						
63323-0382-10	J2710			01/01/2002	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (M.D.V.,AMBER) 0.5 MG/ML	10 ML	VL	IJ	ML		0.5 MG		1	01/01/2002	99/99/9999						
63323-0383-10	J2710			01/01/2002	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (M.D.V.,AMBER) 1 MG/ML	10 ML	VL	IJ	ML		0.5 MG		2	01/01/2002	99/99/9999						
63323-0385-10	J3490			08/13/2007	99/99/9999	UNCLASSIFIED DRUGS	CEFOTETAN 1 GM	1 EA	VL	IJ	EA		1 EA		1	08/13/2007	99/99/9999						
63323-0386-20	J3490			08/13/2007	99/99/9999	UNCLASSIFIED DRUGS	CEFOTETAN 2 GM	1 EA	VL	IJ	EA		1 EA		1	08/13/2007	99/99/9999						
63323-0387-10	J0290			01/01/2002	01/04/2017	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (VIAL) 250 MG	1 EA	VL	IJ	EA		500 MG		0.5	01/01/2002	01/04/2017						
63323-0388-10	J0290			01/01/2002	11/30/2017	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (VIAL) 500 MG	1 EA	VL	IJ	EA		500 MG		1	01/01/2002	11/30/2017						
63323-0389-10	J0290			01/01/2002	06/22/2017	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (VIAL) 1 GM	1 EA	VL	IJ	EA		500 MG		2	01/01/2002	06/22/2017						
63323-0393-06	J0770			03/10/2008	99/99/9999	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE (USP,LYOPHILIZED CAKE) 150 MG	1 EA	VL	IJ	EA		150 MG		1	03/10/2008	99/99/9999						
63323-0398-10	J0456			02/27/2006	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN (10X10ML,LATEX-FREE) 500 MG	1 EA	VL	IV	EA		500 MG		1	02/27/2006	99/99/9999						
63323-0398-12	J0456			02/27/2006	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	NOVAPLUS AZITHROMYCIN (10X10ML) 500 MG	1 EA	VL	IV	EA		500 MG		1	02/27/2006	99/99/9999						
63323-0399-23	J0290			01/01/2002	01/04/2017	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (VIAL) 2 GM	1 EA	VL	IJ	EA		500 MG		4	01/01/2002	01/04/2017						
63323-0407-03	J0706			08/03/2007	99/99/9999	INJECTION, CAFFEINE CITRATE, 5MG	CAFFEINE CITRATE (USP,SDV,PF) 20 MG/ML	3 ML	VL	IV	ML		5 MG		4	08/03/2007	99/99/9999						
63323-0411-10	J2250			01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (M.D.V.) 1 MG/ML	10 ML	VL	IJ	ML		1 MG		1	01/01/2002	99/99/9999						
63323-0411-12	J2250			01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (M.D.V.) 1 MG/ML	2 ML	VL	IJ	ML		1 MG		1	01/01/2002	99/99/9999						
63323-0411-25	J2250			12/08/2003	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (M.D.V.) 1 MG/ML	5 ML	VL	IJ	ML		1 MG		1	12/08/2003	99/99/9999						
63323-0412-02	J2250			01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (M.D.V.) 5 MG/ML	2 ML	VL	IJ	ML		1 MG		5	01/01/2002	99/99/9999						
63323-0412-05	J2250			01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (M.D.V.) 5 MG/ML	5 ML	VL	IJ	ML		1 MG		5	01/01/2002	99/99/9999						
63323-0412-10	J2250			01/01/2002	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (M.D.V.) 5 MG/ML	10 ML	VL	IJ	ML		1 MG		5	01/01/2002	99/99/9999						
63323-0412-25	J2250			01/07/2004	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (M.D.V.) 5 MG/ML	1 ML	VL	IJ	ML		1 MG		5	01/07/2004	99/99/9999						
63323-0469-01	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (VIAL) 50 MG/ML	1 ML	VL	IM	ML		50 MG		1	01/01/2002	99/99/9999						
63323-0469-05	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (M.D.V.) 50 MG/ML	5 ML	VL	IM	ML		50 MG		1	01/01/2002	99/99/9999						
47335-0930-74		None		07/11/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (1X5,HARD GELATIN) 180 MG	5 EA	ST	PO	EA		20 MG		9	07/11/2018	99/99/9999						
63323-0469-51	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL AMERINET CHOICE (VIAL,FLIP-TOP) 50 MG/ML	1 ML	VL	IM	ML		50 MG		1	01/01/2002	99/99/9999						
63323-0471-01	J1631			01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (VIAL) 100 MG/ML	1 ML	VL	IM	ML		50 MG		2	01/01/2002	99/99/9999						

NDC	NDC Mod	HPCS	HPCS Mod	Relationship Start Date	Relationship End Date	HPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HPCS Amount #1	HPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63323-0471-05		J1631		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (M.D.V.) 100 MG/ML	5 ML	VL	IM	ML		50 MG		2	01/01/2002	99/99/9999						
63323-0471-51		J1631		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL AMERINET CHOICE (VIAL,FLIP-TOP) 100 MG/ML	1 ML	VL	IM	ML		50 MG		2	01/01/2002	99/99/9999						
63323-0471-55		J1631		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL AMERINET CHOICE (M.D.V.,FLIP-TOP) 100 MG/ML	5 ML	VL	IM	ML		50 MG		2	01/01/2002	99/99/9999						
63323-0474-01		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL LACTATE (VIAL) 5 MG/ML	1 ML	VL	IM	ML		5 MG		1	01/01/2002	99/99/9999						
63323-0474-10		J1630		01/01/2002	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL LACTATE (M.D.V.) 5 MG/ML	10 ML	VL	IM	ML		5 MG		1	01/01/2002	99/99/9999						
63323-0506-01		J1100		05/30/2003	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (LATEX-FREE) 10 MG/ML	1 ML	VL	IJ	ML		1 MG		10	05/30/2003	99/99/9999						
63323-0513-02		J1580		01/01/2002	99/99/9999	INJECTION, GARAMYCIN, GENTAMICIN, UP TO 80 MG	GENTAMICIN SULFATE PEDIATRIC (PEDIATRIC M.D.V.,PF) 10 MG/ML	2 ML	VL	IJ	ML		80 MG	0.125		01/01/2002	99/99/9999						
63323-0516-10		J1100		08/23/2005	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE 10 MG/ML	10 ML	VL	IJ	ML		1 MG		10	08/23/2005	99/99/9999						
63323-0540-01		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.,P.C.) 1000 U/ML	1 ML	VL	IJ	ML		1000 U		1	01/01/2002	99/99/9999						
63323-0540-11		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.) 1000 U/ML	10 ML	VL	IJ	ML		1000 U		1	01/01/2002	99/99/9999						
63323-0540-31		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.) 1000 U/ML	30 ML	VL	IJ	ML		1000 U		1	01/01/2002	99/99/9999						
63323-0542-01		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.,P.C.) 10000 U/ML	1 ML	VL	IJ	ML		1000 U		10	01/01/2002	99/99/9999						
63323-0542-07		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.) 10000 U/ML	5 ML	VL	IJ	ML		1000 U		10	01/01/2002	99/99/9999						
63323-0544-01		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (M.D.V.,P.C.) 10 U/ML	1 ML	VL	IV	ML		10 U		1	01/01/2002	99/99/9999						
63323-0544-11		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (M.D.V.) 10 U/ML	10 ML	VL	IV	ML		10 U		1	01/01/2002	99/99/9999						
63323-0545-01		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (M.D.V.,P.C.) 100 U/ML	1 ML	VL	IV	ML		10 U		10	01/01/2002	99/99/9999						
63323-0545-05		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (M.D.V.) 100 U/ML	5 ML	VL	IV	ML		10 U		10	01/01/2002	99/99/9999						
63323-0640-01		J1800		01/01/2002	99/99/9999	INJECTION, PROPRANLOLOL HCL, UP TO 1 MG	PROPRANLOLOL HCL (S.D.V.) 1 MG/ML	1 ML	VL	IV	ML		1 MG		1	01/01/2002	99/99/9999						
63323-0614-01		J0360		01/01/2002	99/99/9999	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HCL (S.D.V.) 20 MG/ML	1 ML	VL	IJ	ML		20 MG		1	01/01/2002	99/99/9999						
63323-0614-55		J0360		03/26/2007	99/99/9999	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	NOVAPLUS HYDRALAZINE HYDROCHLORIDE (USP,SDV,LATEX-FREE) 20 MG/ML	1 ML	VL	IJ	ML		20 MG		1	03/26/2007	99/99/9999						
63323-0616-03		J0282		08/02/2002	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL (S.D.V.) 50 MG/ML	3 ML	VL	IV	ML		30 MG	1.66666		08/02/2002	99/99/9999						
63323-0616-09		J0282		12/16/2003	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL (S.D.V.) 50 MG/ML	9 ML	VL	IV	ML		30 MG	1.66666		12/16/2003	99/99/9999						
63323-0617-10		J2260		05/14/2002	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (S.D.V.) 1 MG/ML	10 ML	VL	IV	ML		5 MG	0.2		05/14/2002	99/99/9999						
63323-0617-20		J2260		05/14/2002	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (S.D.V.) 1 MG/ML	20 ML	VL	IV	ML		5 MG	0.2		05/14/2002	99/99/9999						
63323-0617-50		J2260		05/14/2002	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (S.D.V.) 1 MG/ML	50 ML	VL	IV	ML		5 MG	0.2		05/14/2002	99/99/9999						
63323-0651-02		J0150		06/27/2005	12/31/2014	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS, INSTEAD USE A9270)	ADENOSINE (PF) 3 MG/ML	2 ML	VL	IV	ML		6 MG		0.5	06/27/2005	12/31/2014						
63323-0651-04		J0150		06/27/2005	12/31/2014	INJECTION, ADENOSINE PHOSPHATE COMPOUNDS, INSTEAD USE A9270)	ADENOSINE (PF) 3 MG/ML	4 ML	VL	IV	ML		6 MG		0.5	06/27/2005	12/31/2014						
63323-0664-01		J1200		06/12/2002	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL 50 MG/ML	1 ML	VL	IJ	ML		50 MG		1	06/12/2002	99/99/9999						
63323-0665-01		J3105		06/21/2004	99/99/9999	INJECTION, TERBUTALINE SULFATE, UP TO 1 MG	TERBUTALINE SULFATE 1 MG/ML	1 ML	VL	SC	ML		1 MG		1	06/21/2004	99/99/9999						
63323-0731-01		J0636		03/17/2003	99/99/9999	INJECTION, CALCITRIOL, 0.1 MCG	CALCITRIOL 1 MCG/ML	1 ML	AM	IV	ML		0.1 MCG		10	03/17/2003	99/99/9999						
63323-0733-10		J9209		01/01/2002	99/99/9999	INJECTION, MESNA, 200 MG	MESNA (M.D.V.) 100 MG/ML	10 ML	VL	IV	ML		200 MG		0.5	01/01/2002	99/99/9999						
63323-0733-11		J9209		01/01/2002	99/99/9999	INJECTION, MESNA, 200 MG	MESNA (M.D.V.) 100 MG/ML	10 ML	VL	IV	ML		200 MG		0.5	01/01/2002	99/99/9999						
63323-0734-10		J2430		04/25/2002	99/99/9999	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (S.D.V.) 3 MG/ML	10 ML	VL	IV	ML		30 MG		0.1	04/25/2002	99/99/9999						
63323-0734-35		J2430		07/20/2004	02/03/2016	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM OTN (S.D.V.,LATEX-FREE) 3 MG/ML	10 ML	VL	IV	ML		30 MG		0.1	07/20/2004	02/03/2016						
63323-0735-10		J2430		04/25/2002	99/99/9999	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM (S.D.V.) 9 MG/ML	10 ML	VL	IV	ML		30 MG		0.3	04/25/2002	99/99/9999						
63323-0735-35		J2430		09/11/2003	02/03/2016	INJECTION, PAMIDRONATE DISODIUM, PER 30 MG	PAMIDRONATE DISODIUM OTN (S.D.V.) 9 MG/ML	10 ML	VL	IV	ML		30 MG		0.3	09/11/2003	02/03/2016						
63323-0738-04		J3490		01/01/2002	11/12/2012	UNCLASSIFIED DRUGS	FAMOTIDINE (M.D.V.) 10 MG/ML	4 ML	VL	IV	ML		1 EA		1	01/01/2002	11/12/2012						
63323-0738-20		J3490		01/01/2002	99/99/9999	UNCLASSIFIED DRUGS	FAMOTIDINE (M.D.V.) 10 MG/ML	20 ML	VL	IV	ML		1 EA		1	01/01/2002	99/99/9999						
63323-0739-12		J3490		05/14/2002	99/99/9999	UNCLASSIFIED DRUGS	FAMOTIDINE (S.D.V.) 10 MG/ML	2 ML	VL	IV	ML		1 EA		1	05/14/2002	99/99/9999						
63323-0877-15		J2545		01/01/2007	99/99/9999	PENTAMIDINE ISETHIONATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MG	NEBUPENT (S.D.V.,PF) 300 MG	1 EA	VL	IH	EA		300 MG		1	01/01/2007	99/99/9999						
63323-0883-05		J9000		08/06/2007	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HYDROCHLORIDE (USP,STERILE,SDV,PF) 2 MG/ML	5 ML	VL	IV	ML		10 MG		0.2	08/06/2007	99/99/9999						
63323-0883-10		J9000		08/06/2007	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HYDROCHLORIDE (USP,STERILE,SDV,PF) 2 MG/ML	10 ML	VL	IV	ML		10 MG		0.2	08/06/2007	99/99/9999						
63323-0883-30		J9000		08/06/2007	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HYDROCHLORIDE (USP,STERILE,SDV,PF) 2 MG/ML	25 ML	VL	IV	ML		10 MG		0.2	08/06/2007	99/99/9999						
63323-0915-01		J1644		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (M.D.V.,P.C.) 20000 U/ML	1 ML	VL	IJ	ML		1000 U		20	01/01/2002	99/99/9999						
63323-0924-10		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (M.D.V.,P.C.) 0.9%	10 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						
63323-0924-30		A4216		01/01/2004	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (M.D.V.,P.C.) 0.9%	30 ML	VL	IV	ML		10 ML		0.1	01/01/2004	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63323-0965-05		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE CONCENTRATE (S.D.V.,P.C.) 2 MEQ/ML	5 ML	VL	IV	ML		2 MEQ		1	01/01/2002	99/99/9999						
63323-0965-10		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE CONCENTRATE (S.D.V.,P.C.) 2 MEQ/ML	10 ML	VL	IV	ML		2 MEQ		1	01/01/2002	99/99/9999						
63323-0965-15		J3480		01/01/2002	06/04/2012	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE CONCENTRATE (S.D.V.,P.C.) 2 MEQ/ML	15 ML	VL	IV	ML		2 MEQ		1	01/01/2002	06/04/2012						
63323-0965-20		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE CONCENTRATE (S.D.V.,P.C.) 2 MEQ/ML	20 ML	VL	IV	ML		2 MEQ		1	01/01/2002	99/99/9999						
63323-0967-30		J3480		01/01/2002	99/99/9999	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE CONCENTRATE (M.D.V.,P.C.) 2 MEQ/ML	30 ML	VL	IV	ML		2 MEQ		1	01/01/2002	99/99/9999						
63370-0005-25		J7604		01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-25	KO	J7604	KO	01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-35		J7604		01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-35	KO	J7604	KO	01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-45		J7604		01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-45	KO	J7604	KO	01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-50		J7604		01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-50	KO	J7604	KO	01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-55		J7604		01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-55	KO	J7604	KO	01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-62		J7604		01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0005-62	KO	J7604	KO	01/01/2008	05/31/2013	THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYL CYSTEINE (U.S.P.)	1 EA	BO	NA	GM		1 GM		1	01/01/2008	05/31/2013						
63370-0007-25		J0133		01/01/2006	05/31/2013	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM		5 MG	200	01/01/2006	05/31/2013							
63370-0007-35		J0133		01/01/2006	05/31/2013	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM		5 MG	200	01/01/2006	05/31/2013							
63370-0007-50		J0133		01/01/2006	05/31/2013	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	1 EA	BO	NA	GM		5 MG	200	01/01/2006	05/31/2013							
63370-0010-25		J7609		01/01/2007	05/31/2013	THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0010-35		J7609		01/01/2007	05/31/2013	THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0010-35	KO	J7609	KO	01/01/2007	05/31/2013	THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0010-45		J7609		01/01/2007	05/31/2013	THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0010-45	KO	J7609	KO	01/01/2007	05/31/2013	THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0010-50		J7609		01/01/2007	05/31/2013	THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0010-50	KO	J7609	KO	01/01/2007	05/31/2013	THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0016-15		J0278		01/01/2006	05/31/2013	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG	10	01/01/2006	05/31/2013							
63370-0016-25		J0278		01/01/2006	05/31/2013	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG	10	01/01/2006	05/31/2013							
63370-0016-35		J0278		01/01/2006	05/31/2013	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG	10	01/01/2006	05/31/2013							
63370-0016-50		J0278		01/01/2006	05/31/2013	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (U.S.P.)	1 EA	BO	NA	GM		100 MG	10	01/01/2006	05/31/2013							
63370-0018-15		J1320		07/08/2003	05/31/2013	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	1 EA	BO	NA	GM		20 MG	50	07/08/2003	05/31/2013							
63370-0018-25		J1320		07/08/2003	05/31/2013	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	1 EA	BO	NA	GM		20 MG	50	07/08/2003	05/31/2013							
63370-0018-35		J1320		07/08/2003	05/31/2013	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	1 EA	BO	NA	GM		20 MG	50	07/08/2003	05/31/2013							
63370-0020-10		J0285		07/08/2003	05/31/2013	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO	NA	GM		50 MG	20	07/08/2003	05/31/2013							
63370-0020-15		J0285		09/04/2002	05/31/2013	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	JR	NA	GM		50 MG	20	09/04/2002	05/31/2013							
63370-0020-25		J0285		09/04/2002	05/31/2013	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO	NA	GM		50 MG	20	09/04/2002	05/31/2013							
63370-0020-35		J0285		09/04/2002	05/31/2013	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO	NA	GM		50 MG	20	09/04/2002	05/31/2013							
63370-0020-50		J0285		09/04/2002	05/31/2013	INJECTION, AMPHOTERICIN B, 50 MG	AMPHOTERICIN B (U.S.P.)	1 EA	BO	NA	GM		50 MG	20	09/04/2002	05/31/2013							
63370-0022-06		J0364		01/01/2007	05/31/2013	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0022-09		J0364		01/01/2007	05/31/2013	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0022-15		J0364		01/01/2007	05/31/2013	INJECTION, APOMORPHINE HYDROCHLORIDE, 1 MG	APOMORPHINE HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG	1000	01/01/2007	05/31/2013							
63370-0025-10		J7501		07/08/2003	05/31/2013	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (U.S.P.)	1 EA	BO	NA	GM		100 MG	10	07/08/2003	05/31/2013							
63370-0025-15		J7501		07/08/2003	05/31/2013	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (U.S.P.)	1 EA	BO	NA	GM		100 MG	10	07/08/2003	05/31/2013							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63370-0090-35		J1435		07/08/2003	05/31/2013	INJECTION, ESTRONE, PER 1 MG	ESTRONE (USP,1X100GM)	1 EA	JR NA GM	1 MG			1000			07/08/2003	05/31/2013						
63370-0091-25		J3490		07/12/2004	05/31/2013	UNCLASSIFIED DRUGS	FAMOTIDINE (USP)	1 EA	BO NA GM	1 EA						07/12/2004	05/31/2013						
63370-0091-35		J3490		07/12/2004	05/31/2013	UNCLASSIFIED DRUGS	FAMOTIDINE (USP)	1 EA	BO NA GM	1 EA						07/12/2004	05/31/2013						
63370-0091-45		J3490		07/12/2004	05/31/2013	UNCLASSIFIED DRUGS	FAMOTIDINE (USP)	1 EA	BO NA GM	1 EA						07/12/2004	05/31/2013						
63370-0095-15		J9190		07/08/2003	05/31/2013	INJECTION, FLUOROURACIL, 500 MG	5-FLUOROURACIL (U.S.P.)	1 EA	BO NA GM	500 MG			2			07/08/2003	05/31/2013						
63370-0095-25		J9190		07/08/2003	05/31/2013	INJECTION, FLUOROURACIL, 500 MG	5-FLUOROURACIL (U.S.P.)	1 EA	BO NA GM	500 MG			2			07/08/2003	05/31/2013						
63370-0095-35		J9190		07/08/2003	05/31/2013	INJECTION, FLUOROURACIL, 500 MG	5-FLUOROURACIL (U.S.P.)	1 EA	BO NA GM	500 MG			2			07/08/2003	05/31/2013						
63370-0098-15		J7699		07/08/2003	05/31/2013	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1 EA	JR NA GM	1 EA			1			07/08/2003	05/31/2013						
63370-0098-25		J7699		07/08/2003	05/31/2013	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1 EA	BO NA GM	1 EA			1			07/08/2003	05/31/2013						
63370-0098-35		J7699		07/08/2003	05/31/2013	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1 EA	BO NA GM	1 EA			1			07/08/2003	05/31/2013						
63370-0098-45		J7699		07/08/2003	05/31/2013	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1 EA	BO NA GM	1 EA			1			07/08/2003	05/31/2013						
63370-0098-55		J7699		07/08/2003	05/31/2013	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1 EA	BO NA GM	1 EA			1			07/08/2003	05/31/2013						
63370-0102-15		J1630		07/08/2003	05/31/2013	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.,BASE)	1 EA	BO NA GM	5 MG			200			07/08/2003	05/31/2013						
63370-0102-25		J1630		07/08/2003	05/31/2013	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.,BASE)	1 EA	BO NA GM	5 MG			200			07/08/2003	05/31/2013						
63370-0102-35		J1630		07/08/2003	05/31/2013	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.,BASE)	1 EA	BO NA GM	5 MG			200			07/08/2003	05/31/2013						
63370-0107-25		J3410		07/08/2003	05/31/2013	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (U.S.P.)	1 EA	BO NA GM	25 MG			40			07/08/2003	05/31/2013						
63370-0107-35		J3410		07/08/2003	05/31/2013	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (U.S.P.)	1 EA	BO NA GM	25 MG			40			07/08/2003	05/31/2013						
63370-0107-50		J3410		07/08/2003	05/31/2013	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (U.S.P.)	1 EA	BO NA GM	25 MG			40			07/08/2003	05/31/2013						
63370-0107-55		J3410		07/08/2003	05/31/2013	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (U.S.P.)	1 EA	BO NA GM	25 MG			40			07/08/2003	05/31/2013						
63370-0108-15		J1700		07/12/2004	05/31/2013	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE MICRONIZED (USP)	1 EA	BO NA GM	25 MG			40			07/12/2004	05/31/2013						
63370-0108-25		J1700		07/12/2004	05/31/2013	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE MICRONIZED (USP)	1 EA	BO NA GM	25 MG			40			07/12/2004	05/31/2013						
63370-0108-35		J1700		07/12/2004	05/31/2013	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE MICRONIZED (USP)	1 EA	BO NA GM	25 MG			40			07/12/2004	05/31/2013						
63370-0108-45		J1700		07/12/2004	05/31/2013	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE MICRONIZED (USP)	1 EA	BO NA GM	25 MG			40			07/12/2004	05/31/2013						
63370-0108-50		J1700		07/12/2004	05/31/2013	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE MICRONIZED (USP)	1 EA	BO NA GM	25 MG			40			07/12/2004	05/31/2013						
63370-0109-10		J3490		01/01/2007	05/31/2013	UNCLASSIFIED DRUGS	SODIUM HYALURONATE (1X1GM)	1 EA	NA NA GM	1 EA			1			01/01/2007	05/31/2013						
63370-0109-16		J3490		01/01/2007	05/31/2013	UNCLASSIFIED DRUGS	SODIUM HYALURONATE (1X0.2GM)	1 EA	NA NA GM	1 EA			1			01/01/2007	05/31/2013						
63370-0120-10		J7645		01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-10	KO	J7645	KO	01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-15		J7645		01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-15	KO	J7645	KO	01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-25		J7645		01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-25	KO	J7645	KO	01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-35		J7645		01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-35	KO	J7645	KO	01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-50		J7645		01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0120-50	KO	J7645	KO	01/01/2007	05/31/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (EP)	1 EA	BO NA GM	1 MG			1000			01/01/2007	05/31/2013						
63370-0122-15		J1835		07/08/2003	05/31/2013	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE MICRONIZED	1 EA	JR NA GM	50 MG			20			07/08/2003	05/31/2013						
63370-0122-25		J1835		07/08/2003	05/31/2013	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE MICRONIZED	1 EA	BO NA GM	50 MG			20			07/08/2003	05/31/2013						
63370-0122-35		J1835		07/08/2003	05/31/2013	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE MICRONIZED	1 EA	BO NA GM	50 MG			20			07/08/2003	05/31/2013						
63370-0124-20		J1840		07/08/2003	05/31/2013	INJECTION, KANAMYCIN SULFATE, UP TO 500 MG	KANAMYCIN SULFATE (U.S.P.)	1 EA	BO NA GM	500 MG			2			07/08/2003	05/31/2013						
63370-0124-25		J1840		07/08/2003	05/31/2013	INJECTION, KANAMYCIN SULFATE, UP TO 500 MG	KANAMYCIN SULFATE (U.S.P.)	1 EA	BO NA GM	500 MG			2			07/08/2003	05/31/2013						
63370-0124-35		J1840		07/08/2003	05/31/2013	INJECTION, KANAMYCIN SULFATE, UP TO 500 MG	KANAMYCIN SULFATE (U.S.P.)	1 EA	BO NA GM	500 MG			2			07/08/2003	05/31/2013						
63370-0138-10		J1030		10/25/2006	05/31/2013	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRO (1X1GM,USP)	1 EA	NA NA GM	40 MG			25			10/25/2006	05/31/2013						
63370-0138-15		J1030		10/25/2006	05/31/2013	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRO (1X5GM,USP)	1 EA	NA NA GM	40 MG			25			10/25/2006	05/31/2013						
63370-0138-25		J1030		10/25/2006	05/31/2013	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRO (1X25GM,USP)	1 EA	NA NA GM	40 MG			25			10/25/2006	05/31/2013						
63370-0138-35		J1030		10/25/2006	05/31/2013	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRO (1X100GM,USP)	1 EA	NA NA GM	40 MG			25			10/25/2006	05/31/2013						
63370-0138-50		J1030		10/25/2006	05/31/2013	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE MICRO (1X100GM,USP)	1 EA	NA NA GM	40 MG			25			10/25/2006	05/31/2013						
63370-0141-15		J2765		07/08/2003	05/31/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P., MONOHYDRATE)	1 EA	BO NA GM	10 MG			100			07/08/2003	05/31/2013						
63370-0141-25		J2765		07/08/2003	05/31/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P., MONOHYDRATE)	1 EA	BO NA GM	10 MG			100			07/08/2003	05/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63370-0141-35		J2765		07/08/2003	05/31/2013	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL (U.S.P., MONOHYDRATE)	1 EA	BO	NA	GM	10 MG	100		07/08/2003	05/31/2013							
63370-0143-45		J2800		07/08/2003	05/31/2013	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (U.S.P.)	1 EA	BO	NA	GM	10 ML	1		07/08/2003	05/31/2013							
63370-0143-45		J2800		07/08/2003	05/31/2013	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (U.S.P.)	1 EA	BO	NA	GM	10 ML	1		07/08/2003	05/31/2013							
63370-0143-50		J2800		07/08/2003	05/31/2013	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (U.S.P.)	1 EA	BO	NA	GM	10 ML	1		07/08/2003	05/31/2013							
63370-0145-14		J2001		01/01/2004	05/31/2013	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2004	05/31/2013							
63370-0145-25		J2001		01/01/2004	05/31/2013	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2004	05/31/2013							
63370-0145-35		J2001		01/01/2004	05/31/2013	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2004	05/31/2013							
63370-0145-50		J2001		01/01/2004	05/31/2013	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2004	05/31/2013							
63370-0145-55		J2001		01/01/2004	05/31/2013	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2004	05/31/2013							
63370-0152-25		J3490		07/08/2003	05/31/2013	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM	1 EA	1		07/08/2003	05/31/2013							
63370-0152-35		J3490		07/08/2003	05/31/2013	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM	1 EA	1		07/08/2003	05/31/2013							
63370-0152-45		J3490		07/08/2003	05/31/2013	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	1 EA	BO	NA	GM	1 EA	1		07/08/2003	05/31/2013							
63370-0153-20		J7670		01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0153-20	KO	J7670	KO	01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0153-25		J7670		01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0153-25	KO	J7670	KO	01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0153-35		J7670		01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0153-35	KO	J7670	KO	01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0153-45		J7670		01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0153-45	KO	J7670	KO	01/01/2007	05/31/2013	METAPROTERENOL SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	METAPROTERENOL SULFATE (U.S.P.)	1 EA	BO	NA	GM	10 MG	100		01/01/2007	05/31/2013							
63370-0154-10		J8610		07/08/2003	05/31/2013	METHOTREXATE; ORAL, 2.5 MG	METHOTREXATE (U.S.P.)	1 EA	BO	NA	GM	2.5 MG	400		07/08/2003	05/31/2013							
63370-0154-15		J8610		07/08/2003	05/31/2013	METHOTREXATE; ORAL, 2.5 MG	METHOTREXATE (U.S.P.)	1 EA	BO	NA	GM	2.5 MG	400		07/08/2003	05/31/2013							
63370-0154-25		J8610		07/08/2003	05/31/2013	METHOTREXATE; ORAL, 2.5 MG	METHOTREXATE (U.S.P.)	1 EA	BO	NA	GM	2.5 MG	400		07/08/2003	05/31/2013							
63370-0165-15		J2440		07/08/2003	05/31/2013	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		07/08/2003	05/31/2013							
63370-0165-25		J2440		07/08/2003	05/31/2013	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		07/08/2003	05/31/2013							
63370-0165-35		J2440		07/08/2003	05/31/2013	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HYDROCHLORIDE (U.S.P.)	1 EA	BO	NA	GM	60 MG	16.66666		07/08/2003	05/31/2013							
63370-0170-06		J2760		07/08/2003	05/31/2013	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/08/2003	05/31/2013							
63370-0170-09		J2760		07/08/2003	05/31/2013	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/08/2003	05/31/2013							
63370-0170-10		J2760		07/08/2003	05/31/2013	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/08/2003	05/31/2013							
63370-0170-15		J2760		07/08/2003	05/31/2013	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/08/2003	05/31/2013							
63370-0176-25		J1165		07/08/2003	05/31/2013	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	BO	NA	GM	50 MG	20		07/08/2003	05/31/2013							
63370-0176-35		J1165		07/08/2003	05/31/2013	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	BO	NA	GM	50 MG	20		07/08/2003	05/31/2013							
63370-0176-45		J1165		07/08/2003	05/31/2013	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	BO	NA	GM	50 MG	20		07/08/2003	05/31/2013							
63370-0176-53		J1165		07/08/2003	05/31/2013	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	1 EA	BO	NA	GM	50 MG	20		07/08/2003	05/31/2013							
63370-0194-15		J7506		07/12/2004	05/31/2013	PREDNISONE, ORAL, PER SMG	PREDNISONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/12/2004	05/31/2013							
63370-0194-25		J7506		07/12/2004	05/31/2013	PREDNISONE, ORAL, PER SMG	PREDNISONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/12/2004	05/31/2013							
63370-0194-35		J7506		07/12/2004	05/31/2013	PREDNISONE, ORAL, PER SMG	PREDNISONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/12/2004	05/31/2013							
63370-0194-45		J7506		07/12/2004	05/31/2013	PREDNISONE, ORAL, PER SMG	PREDNISONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/12/2004	05/31/2013							
63370-0194-50		J7506		07/12/2004	05/31/2013	PREDNISONE, ORAL, PER SMG	PREDNISONE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	5 MG	200		07/12/2004	05/31/2013							
63370-0195-15		J2650		07/08/2003	05/31/2013	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	1 ML	20		07/08/2003	05/31/2013							
63370-0195-25		J2650		07/08/2003	05/31/2013	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	1 ML	20		07/08/2003	05/31/2013							
63370-0195-35		J2650		07/08/2003	05/31/2013	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	1 ML	20		07/08/2003	05/31/2013							
63370-0195-50		J2650		07/08/2003	05/31/2013	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	1 ML	20		07/08/2003	05/31/2013							
63370-0195-55		J2650		07/08/2003	05/31/2013	INJECTION, PREDNISOLONE ACETATE, UP TO 1 ML	PREDNISOLONE ACETATE MICRONIZED (U.S.P.)	1 EA	BO	NA	GM	1 ML	20		07/08/2003	05/31/2013							
63370-0198-25		Q0165		12/19/2003	05/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (USP)	1 EA	BO	NA	GM	10 MG	100		12/19/2003	05/31/2013							
63370-0198-35		Q0165		12/19/2003	05/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (USP)	1 EA	BO	NA	GM	10 MG	100		12/19/2003	05/31/2013							
63370-0198-45		Q0165		12/19/2003	05/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (USP)	1 EA	BO	NA	GM	10 MG	100		12/19/2003	05/31/2013							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63370-0199-35		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (USP,SOY)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0199-45		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (USP,SOY)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0199-50		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (USP,SOY)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0199-55		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (USP,SOY)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0199-62		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (USP,SOY)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0200-35		J2675		12/19/2003	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (USP,YAM)	1 EA	BO	NA	GM		50 MG		20	12/19/2003	05/31/2013						
63370-0200-45		J2675		12/19/2003	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (USP,YAM)	1 EA	BO	NA	GM		50 MG		20	12/19/2003	05/31/2013						
63370-0200-50		J2675		07/08/2003	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (YAM)	1 EA	BO	NA	GM		50 MG		20	07/08/2003	05/31/2013						
63370-0200-55		J2675		12/19/2003	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (USP,YAM)	1 EA	BO	NA	GM		50 MG		20	12/19/2003	05/31/2013						
63370-0202-35		J2675		07/12/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE WETTABLE (U.S.P.,YAM)	1 EA	BO	NA	GM		50 MG		20	07/12/2004	05/31/2013						
63370-0202-45		J2675		07/12/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE WETTABLE (U.S.P.,YAM)	1 EA	BO	NA	GM		50 MG		20	07/12/2004	05/31/2013						
63370-0202-50		J2675		07/12/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE WETTABLE (U.S.P.,YAM)	1 EA	BO	NA	GM		50 MG		20	07/12/2004	05/31/2013						
63370-0203-25		J2550		07/08/2003	05/31/2013	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	BO	NA	GM		50 MG		20	07/08/2003	05/31/2013						
63370-0203-35		J2550		07/08/2003	05/31/2013	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	BO	NA	GM		50 MG		20	07/08/2003	05/31/2013						
63370-0203-45		J2550		07/08/2003	05/31/2013	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	BO	NA	GM		50 MG		20	07/08/2003	05/31/2013						
63370-0203-50		J2550		07/08/2003	05/31/2013	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	1 EA	BO	NA	GM		50 MG		20	07/08/2003	05/31/2013						
63370-0204-35		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (YAM)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0204-45		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (YAM)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0204-50		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (YAM)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0204-55		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (YAM)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0204-62		J2675		02/25/2004	05/31/2013	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE MICRONIZED (YAM)	1 EA	BO	NA	GM		50 MG		20	02/25/2004	05/31/2013						
63370-0205-25		J1800		07/08/2003	05/31/2013	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0205-35		J1800		07/08/2003	05/31/2013	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0205-45		J1800		07/08/2003	05/31/2013	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0210-04		J0270		07/08/2003	05/31/2013	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO	NA	GM		1.25 MCG		800000	07/08/2003	05/31/2013						
63370-0210-06		J0270		07/08/2003	05/31/2013	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO	NA	GM		1.25 MCG		800000	07/08/2003	05/31/2013						
63370-0210-10		J0270		07/08/2003	05/31/2013	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	1 EA	BO	NA	GM		1.25 MCG		800000	07/08/2003	05/31/2013						
63370-0218-25		J2780		07/08/2003	05/31/2013	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	JR	NA	GM		25 MG		40	07/08/2003	05/31/2013						
63370-0218-35		J2780		07/08/2003	05/31/2013	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	JR	NA	GM		25 MG		40	07/08/2003	05/31/2013						
63370-0218-45		J2780		07/08/2003	05/31/2013	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO	NA	GM		25 MG		40	07/08/2003	05/31/2013						
63370-0218-50		J2780		07/08/2003	05/31/2013	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	RANITIDINE HCL (U.S.P.)	1 EA	BO	NA	GM		25 MG		40	07/08/2003	05/31/2013						
63370-0233-35		J3000		07/08/2003	05/31/2013	INJECTION, STREPTOMYCIN, UP TO 1 GM	STREPTOMYCIN SULFATE (U.S.P., NON-STERILE)	1 EA	BO	NA	GM		1 GM		1	07/08/2003	05/31/2013						
63370-0233-50		J3000		07/08/2003	05/31/2013	INJECTION, STREPTOMYCIN, UP TO 1 GM	STREPTOMYCIN SULFATE (U.S.P., NON-STERILE)	1 EA	BO	NA	GM		1 GM		1	07/08/2003	05/31/2013						
63370-0250-15		J7681		07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0250-15	KO	J7681	KO	07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0250-20		J7681		07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0250-20	KO	J7681	KO	07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0250-25		J7681		07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0250-25	KO	J7681	KO	07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0250-35		J7681		07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0250-35	KO	J7681	KO	07/08/2003	05/31/2013	TERBUTALINE SULFATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	TERBUTALINE SULFATE (U.S.P.)	1 EA	BO	NA	GM		1 MG		1000	07/08/2003	05/31/2013						
63370-0275-10		J7685		01/01/2007	05/31/2013	TOBRAMYCN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCN SULFATE (U.S.P.)	1 EA	BO	NA	GM		300 MG		3.33333	01/01/2007	05/31/2013						
63370-0275-10	KO	J7685	KO	01/01/2007	05/31/2013	TOBRAMYCN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCN SULFATE (U.S.P.)	1 EA	BO	NA	GM		300 MG		3.33333	01/01/2007	05/31/2013						
63370-0275-15		J7685		01/01/2007	05/31/2013	TOBRAMYCN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCN SULFATE (U.S.P.)	1 EA	BO	NA	GM		300 MG		3.33333	01/01/2007	05/31/2013						
63370-0275-15	KO	J7685	KO	01/01/2007	05/31/2013	TOBRAMYCN, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 300 MILLIGRAMS	TOBRAMYCN SULFATE (U.S.P.)	1 EA	BO	NA	GM		300 MG		3.33333	01/01/2007	05/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63370-0939-15		J1230		07/08/2003	05/31/2013	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	BO NA GM	10 MG			100		07/08/2003	05/31/2013							
63370-0939-25		J1230		07/08/2003	05/31/2013	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	BO NA GM	10 MG			100		07/08/2003	05/31/2013							
63370-0939-35		J1230		07/08/2003	05/31/2013	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	1 EA	BO NA GM	10 MG			100		07/08/2003	05/31/2013							
63370-0950-25		J2271		07/08/2003	05/31/2013	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0950-35		J2271		07/08/2003	05/31/2013	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0950-45		J2271		07/08/2003	05/31/2013	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0950-50		J2271		07/08/2003	05/31/2013	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0968-04		J3490		07/08/2003	05/31/2013	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P.)	1 EA	NA NA GM	1 EA			1		07/08/2003	05/31/2013							
63370-0968-06		J3490		07/08/2003	05/31/2013	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P.)	1 EA	BO NA GM	1 EA			1		07/08/2003	05/31/2013							
63370-0970-25		J3140		01/31/2002	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	BO NA GM	50 MG			20		01/31/2002	05/31/2013							
63370-0970-35		J3140		01/31/2002	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	BO NA GM	50 MG			20		01/31/2002	05/31/2013							
63370-0970-45		J3140		01/31/2002	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	JR NA GM	50 MG			20		01/31/2002	05/31/2013							
63370-0970-50		J3140		01/31/2002	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (U.S.P.)	1 EA	JR NA GM	50 MG			20		01/31/2002	05/31/2013							
63370-0971-25		J3140		12/19/2003	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (USP,YAM)	1 EA	BO NA GM	50 MG			20		12/19/2003	05/31/2013							
63370-0971-35		J3140		12/19/2003	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (USP,YAM)	1 EA	BO NA GM	50 MG			20		12/19/2003	05/31/2013							
63370-0971-45		J3140		12/19/2003	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (USP,YAM)	1 EA	BO NA GM	50 MG			20		12/19/2003	05/31/2013							
63370-0971-50		J3140		12/19/2003	05/31/2013	INJECTION, TESTOSTERONE SUSPENSION, UP TO 50 MG	TESTOSTERONE MICRONIZED (USP,YAM)	1 EA	BO NA GM	50 MG			20		12/19/2003	05/31/2013							
63370-0980-25		J1070		07/08/2003	05/31/2013	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0980-35		J1070		07/08/2003	05/31/2013	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	JR NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0980-50		J1070		07/08/2003	05/31/2013	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1 EA	JR NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0983-15		J3130		01/19/2004	05/31/2013	INJECTION, TESTOSTERONE ENANTHATE, UP TO 200 MG	TESTOSTERONE ENANTHATE (U.S.P.)	1 EA	BO NA GM	200 MG			5		01/19/2004	05/31/2013							
63370-0983-25		J3130		01/19/2004	05/31/2013	INJECTION, TESTOSTERONE ENANTHATE, UP TO 200 MG	TESTOSTERONE ENANTHATE (U.S.P.)	1 EA	BO NA GM	200 MG			5		01/19/2004	05/31/2013							
63370-0983-35		J3130		01/19/2004	05/31/2013	INJECTION, TESTOSTERONE ENANTHATE, UP TO 200 MG	TESTOSTERONE ENANTHATE (U.S.P.)	1 EA	BO NA GM	200 MG			5		01/19/2004	05/31/2013							
63370-0983-50		J3130		01/19/2004	05/31/2013	INJECTION, TESTOSTERONE ENANTHATE, UP TO 200 MG	TESTOSTERONE ENANTHATE (U.S.P.)	1 EA	BO NA GM	200 MG			5		01/19/2004	05/31/2013							
63370-0985-25		J3150		07/08/2003	05/31/2013	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0985-35		J3150		07/08/2003	05/31/2013	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0985-45		J3150		07/08/2003	05/31/2013	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63370-0985-50		J3150		07/08/2003	05/31/2013	INJECTION, TESTOSTERONE PROPIONATE, UP TO 100 MG	TESTOSTERONE PROPIONATE (U.S.P.)	1 EA	BO NA GM	100 MG			10		07/08/2003	05/31/2013							
63402-0511-24		J7614		04/01/2008	04/20/2016	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX PEDIATRIC 0.31 MG/3 ML	3 ML	PC IH ML	0.5 MG			0.20666		04/01/2008	04/20/2016							
63402-0511-24	KO	J7614	KO	04/01/2008	04/20/2016	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX PEDIATRIC 0.31 MG/3 ML	3 ML	PC IH ML	0.5 MG			0.20666		04/01/2008	04/20/2016							
63402-0512-24		J7614		04/01/2008	12/14/2015	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.63 MG/3 ML	3 ML	PC IH ML	0.5 MG			0.42		04/01/2008	12/14/2015							
63402-0512-24	KO	J7614	KO	04/01/2008	12/14/2015	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.63 MG/3 ML	3 ML	PC IH ML	0.5 MG			0.42		04/01/2008	12/14/2015							
63402-0513-24		J7614		04/01/2008	10/21/2015	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 1.25 MG/3 ML	3 ML	PC IH ML	0.5 MG			0.83333		04/01/2008	10/21/2015							
63402-0513-24	KO	J7614	KO	04/01/2008	10/21/2015	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 1.25 MG/3 ML	3 ML	PC IH ML	0.5 MG			0.83333		04/01/2008	10/21/2015							
63402-0515-30		J7612		04/01/2008	06/21/2015	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 0.5 MG	XOPENEX (PF) 1.25 MG/0.5 ML	0.5 ML	PC IH ML	0.5 MG			5		04/01/2008	06/21/2015							
63402-0911-30	KO	J7605	KO	01/01/2008	99/99/9999	ARFORMOTEROL, INHALATION SOLUTION, FDA APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 15 MICROGRAMS	BROVANA 15 MCG/2 ML	2 ML	PC IH ML	15 MCG			0.5		01/01/2008	99/99/9999							
63402-0911-64	KO	J7605	KO	01/01/2008	99/99/9999	ARFORMOTEROL, INHALATION SOLUTION, FDA APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 15 MICROGRAMS	BROVANA (60X2ML) 15 MCG/2 ML	2 ML	VL IH ML	15 MCG			0.5		01/01/2008	99/99/9999							
63459-0391-20		J3490		03/31/2008	99/99/9999	UNCLASSIFIED DRUGS	TREANDA	1 EA	VL IV EA	1 EA			1		03/31/2008	99/99/9999							
63459-0600-10		J9017		07/15/2006	12/15/2017	INJECTION, ARSENIC TRIOXIDE, 1 MG	TRISENOX (10X10 AMP PF) 1 MG/ML OPANA (1MLX10,PABEN-FREE) 1 MG/ML	10 ML	AM IV ML	1 MG			1		07/15/2006	12/15/2017							
63481-0624-10		J2410		05/07/2007	04/11/2018	INJECTION, OXYMORPHONE HCL, UP TO 1 MG	OPANA (1MLX10,PABEN-FREE) 1 MG/ML	1 ML	AM IV ML	1 MG			1		05/07/2007	04/11/2018							
63629-1262-01		J8999		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	AROMASIN 25 MG	30 EA	NA PO EA	1 EA			1		11/01/2004	99/99/9999							
63629-1335-01		Q0165		11/01/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10 EA	NA PO EA	10 MG			1		11/01/2004	12/31/2013							
63629-1335-02		Q0165		11/01/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30 EA	NA PO EA	10 MG			1		11/01/2004	12/31/2013							
63629-1335-03		Q0165		11/01/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20 EA	NA PO EA	10 MG			1		11/01/2004	12/31/2013							
63629-1343-01		Q0163		11/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	30 EA	BO PO EA	50 MG			0.5		11/01/2004	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63629-1343-02		Q0163		11/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	20	EA	BO	PO	EA	50	MG	0.5	11/01/2004	99/99/9999						
63629-1343-03		Q0163		11/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	42	EA	BO	PO	EA	50	MG	0.5	11/01/2004	99/99/9999						
63629-1343-04		Q0163		11/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 25 MG	24	EA	BO	PO	EA	50	MG	0.5	11/01/2004	99/99/9999						
63629-1349-01		Q0163		11/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 50 MG	15	EA	BO	PO	EA	50	MG	1	11/01/2004	99/99/9999						
63629-1349-02		Q0163		11/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 50 MG	20	EA	BO	PO	EA	50	MG	1	11/01/2004	99/99/9999						
63629-1349-03		Q0163		11/01/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE 50 MG	30	EA	BO	PO	EA	50	MG	1	11/01/2004	99/99/9999						
63629-1472-01		None		11/01/2004	99/99/9999	METHOTREXATE, 2.5 MG, ORAL HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	METHOTREXATE 2.5 MG	30	EA	NA	PO	EA	2.5	MG	1	11/01/2004	99/99/9999						
63629-1533-01		Q0177		11/01/2004	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20	EA	NA	PO	EA	25	MG	1	11/01/2004	99/99/9999						
63629-1533-02		Q0177		11/01/2004	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	NA	PO	EA	25	MG	1	11/01/2004	99/99/9999						
63629-1579-01		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	21	EA	NA	PO	EA	5	MG	2	11/01/2004	12/31/2015						
63629-1579-02		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	40	EA	NA	PO	EA	5	MG	2	11/01/2004	12/31/2015						
63629-1579-03		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 10 MG	30	EA	NA	PO	EA	5	MG	2	11/01/2004	12/31/2015						
63629-1587-01		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	20	EA	NA	PO	EA	5	MG	4	11/01/2004	12/31/2015						
63629-1587-02		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	NA	PO	EA	5	MG	4	11/01/2004	12/31/2015						
63629-1587-03		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	40	EA	NA	PO	EA	5	MG	4	11/01/2004	12/31/2015						
63629-1587-04		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	15	EA	NA	PO	EA	5	MG	4	11/01/2004	12/31/2015						
63629-1591-01		Q0169		11/01/2004	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 12.5 MG	12	EA	NA	PO	EA	12.5	MG	1	11/01/2004	99/99/9999						
63629-1591-02		Q0169		11/01/2004	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 12.5 MG	4	EA	NA	PO	EA	12.5	MG	1	11/01/2004	99/99/9999						
63629-1591-03		Q0169		11/01/2004	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 12.5 MG	2	EA	NA	PO	EA	12.5	MG	1	11/01/2004	99/99/9999						
63629-1591-04		Q0169		11/01/2004	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 12.5 MG	30	EA	NA	PO	EA	12.5	MG	1	11/01/2004	99/99/9999						
63629-1605-01		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	30	EA	NA	PO	EA	5	MG	1	11/01/2004	12/31/2015						
63629-1605-02		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	78	EA	NA	PO	EA	5	MG	1	11/01/2004	12/31/2015						
63629-1605-03		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	38	EA	NA	PO	EA	5	MG	1	11/01/2004	12/31/2015						
63629-1605-04		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	21	EA	NA	PO	EA	5	MG	1	11/01/2004	12/31/2015						
63629-1605-05		J7506		11/01/2004	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	15	EA	NA	PO	EA	5	MG	1	11/01/2004	12/31/2015						
63629-1676-01		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1676-02		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1676-03		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	35	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1677-01		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	20	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1677-02		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	28	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1677-03		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1678-01		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	25	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1678-02		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	35	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1678-03		J8499		11/01/2004	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	11/01/2004	99/99/9999						
63629-1742-01		Q0170		11/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	15	EA	NA	PO	EA	25	MG	1	11/01/2004	12/31/2013						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63629-1742-02		Q0170		11/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	30	EA	NA	PO	EA	25	MG	1	11/01/2004	12/31/2013						
63629-1742-03		Q0170		11/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10	EA	NA	PO	EA	25	MG	1	11/01/2004	12/31/2013						
63629-1742-04		Q0170		11/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	20	EA	NA	PO	EA	25	MG	1	11/01/2004	12/31/2013						
63629-1841-01		Q0164		11/01/2004	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	20	EA	NA	PO	EA	5	MG	1	11/01/2004	99/99/9999						
63629-1856-01		Q0177		11/01/2004	99/99/9999	HYDROXYZYNE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZYNE PAMOATE 25 MG	30	EA	NA	PO	EA	25	MG	1	11/01/2004	99/99/9999						
63629-1856-02		Q0177		11/01/2004	99/99/9999	HYDROXYZYNE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZYNE PAMOATE 25 MG	60	EA	NA	PO	EA	25	MG	1	11/01/2004	99/99/9999						
63629-1862-01		J7510		11/01/2004	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE 15 MG/5 ML	60	ML	NA	PO	ML	5	MG	0.6	11/01/2004	99/99/9999						
63629-1870-01		Q0170		11/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	120	ML	NA	PO	ML	25	MG	0.05	11/01/2004	12/31/2013						
63629-1870-02		Q0170		11/01/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	240	ML	NA	PO	ML	25	MG	0.05	11/01/2004	12/31/2013						
63739-0161-10		J7509		02/27/2007	06/30/2012	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (USP) 4 MG	100	EA	BX	PO	EA	4	MG	1	02/27/2007	06/30/2012						
63739-0165-10		J8999		02/27/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (USP) 40 MG	100	EA	BX	PO	EA	1	EA	1	02/27/2007	99/99/9999						
63739-0207-10		J7506		03/01/2007	04/30/2012	PREDNISONE, ORAL, PER 5MG	PREDNISONE (USP) 5 MG	100	EA	BX	PO	EA	5	MG	1	03/01/2007	04/30/2012						
63739-0213-10		Q0170		02/27/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	100	EA	BX	PO	EA	25	MG	1	02/27/2007	12/31/2013						
63739-0269-10		J8999		02/27/2007	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	TAMOXIFEN CITRATE (USP) 10 MG	100	EA	BX	PO	EA	1	EA	1	02/27/2007	99/99/9999						
63807-0100-11		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (PF,LATEX-FREE) 0.9%	10	ML	BX	IJ	ML	10	ML	0.1	01/01/2007	99/99/9999						
63807-0100-20		A4216		04/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (30X10ML,PF) 0.9%	10	ML	SR	IJ	ML	10	ML	0.1	04/01/2007	99/99/9999						
63807-0100-30		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (SRN,PF) 0.9%	2.5	ML	SR	IJ	ML	10	ML	0.1	01/01/2007	99/99/9999						
63807-0100-33		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (PF,LATEX-FREE) 0.9%	2.5	ML	BX	IJ	ML	10	ML	0.1	01/01/2007	99/99/9999						
63807-0100-35		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (USP,10X3ML,SYRINGE,PF) 0.9%	3	ML	SR	IJ	ML	10	ML	0.1	01/01/2007	99/99/9999						
63807-0100-50		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (SRN,PF) 0.9%	5	ML	SR	IJ	ML	10	ML	0.1	01/01/2007	99/99/9999						
63807-0100-51		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (PF,LATEX-FREE) 0.9%	5	ML	BX	IJ	ML	10	ML	0.1	01/01/2007	99/99/9999						
63807-0100-55		A4216		04/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (100X5ML,PF) 0.9%	5	ML	SR	IJ	ML	10	ML	0.1	04/01/2007	99/99/9999						
63807-0100-75		A4216		01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (SRN,PF) 0.9%	10	ML	SR	IJ	ML	10	ML	0.1	01/01/2007	99/99/9999						
63807-0100-92		A4216		01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (2X10ML,PF,LATEX-FREE) 0.9%	10	ML	SR	IJ	ML	10	ML	0.1	01/01/2007	02/03/2016						
63807-0102-11		A4216		01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SYREX (PF,LATEX-FREE) 0.9%	10	ML	SR	IJ	ML	10	ML	0.1	01/01/2007	02/03/2016						
63807-0300-35		J1642		04/12/2007	11/25/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (USP,3MLX100,PF) 1 U/ML	3	ML	SR	IV	ML	10	U	0.1	04/12/2007	11/25/2016						
63807-0400-31		J1642		01/01/2007	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LATEX-FREE) 2 U/ML	5	ML	SR	IV	ML	10	U	0.2	01/01/2007	99/99/9999						
63807-0400-35		J1642		04/12/2007	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (USP,3MLX100,PF) 2 U/ML	3	ML	SR	IV	ML	10	U	0.2	04/12/2007	99/99/9999						
63807-0500-31		J1642		01/01/2007	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LATEX-FREE) 10 U/ML	3	ML	SR	IV	ML	10	U	1	01/01/2007	99/99/9999						
63807-0500-51		J1642		01/01/2007	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LATEX-FREE) 10 U/ML	5	ML	SR	IV	ML	10	U	1	01/01/2007	99/99/9999						
63807-0600-31		J1642		01/01/2007	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LATEX-FREE) 100 U/ML	3	ML	SR	IV	ML	10	U	10	01/01/2007	99/99/9999						
63807-0600-51		J1642		01/01/2007	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LATEX-FREE) 100 U/ML	5	ML	SR	IV	ML	10	U	10	01/01/2007	99/99/9999						
63807-0600-55		J1642		05/10/2005	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH 100 U/ML	5	ML	SR	IV	ML	10	U	10	05/10/2005	99/99/9999						
63868-0087-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MEDIPHEDRYL 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63868-0087-24		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MEDIPHEDRYL 25 MG	24	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63868-0500-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MEDIPHEDRYL (MINITAB) 25 MG	100	EA	BO	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63868-0611-32		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	NIGHT TIME SLEEP AID 25 MG	32	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63868-0612-32		Q0163		04/01/2006	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	QUALITY CHOICE SLEEP AID (SOFTGEL) 50 MG	32	EA	BO	PO	EA	50	MG	1	04/01/2006	99/99/9999						
63868-0789-24		Q0163		11/01/2003	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	QUALITY CHOICE REST SIMPLY (CAPLET) 25 MG	24	EA	BX	PO	EA	50	MG	0.5	11/01/2003	99/99/9999						
63868-0823-54		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALLERGY CHILDREN'S (AF,CHERRY) 12.5 MG/5 ML	118	ML	BO	PO	ML	50	MG	0.05	01/01/2002	99/99/9999						
63874-0005-01		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	100	EA	NA	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63874-0005-02		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	1000	EA	NA	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63874-0005-06		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	6	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-09		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	9	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-10		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	10	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-12		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	12	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-14		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	14	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-15		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15	EA	NA	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63874-0005-20		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20	EA	NA	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63874-0005-21		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	21	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-24		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	24	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-25		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	25	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-28		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	28	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63874-0005-30		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30	EA	BX	PO	EA	50	MG	0.5	01/01/2002	99/99/9999						
63874-0005-40		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	40	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-45		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	45	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-60		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0005-90		Q0163		05/10/2004	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	90	EA	BO	PO	EA	50	MG	0.5	05/10/2004	99/99/9999						
63874-0006-01		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2002	02/03/2016						
63874-0006-02		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	1000	EA	BO	PO	EA	50	MG	1	01/01/2002	02/03/2016						
63874-0006-07		Q0163		05/10/2004	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	7	EA	BO	PO	EA	50	MG	1	05/10/2004	02/03/2016						
63874-0006-10		Q0163		05/10/2004	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	10	EA	BO	PO	EA	50	MG	1	05/10/2004	02/03/2016						
63874-0006-12		Q0163		05/10/2004	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	12	EA	BO	PO	EA	50	MG	1	05/10/2004	02/03/2016						
63874-0006-14		Q0163		05/10/2004	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	14	EA	BO	PO	EA	50	MG	1	05/10/2004	02/03/2016						
63874-0006-15		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15	EA	BO	PO	EA	50	MG	1	01/01/2002	02/03/2016						
63874-0006-20		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20	EA	BO	PO	EA	50	MG	1	01/01/2002	02/03/2016						
63874-0006-25		Q0163		05/10/2004	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	25	EA	BO	PO	EA	50	MG	1	05/10/2004	02/03/2016						
63874-0006-28		Q0163		05/10/2004	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	28	EA	BO	PO	EA	50	MG	1	05/10/2004	02/03/2016						
63874-0006-30		Q0163		01/01/2002	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30	EA	BO	PO	EA	50	MG	1	01/01/2002	02/03/2016						
63874-0006-60		Q0163		05/10/2004	02/03/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	60	EA	BO	PO	EA	50	MG	1	05/10/2004	02/03/2016						
63874-0246-00		Q0144		03/15/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX (Z-PACK) 250 MG	6	EA	NA	PO	EA	1	GM	0.25	03/15/2006	99/99/9999						
63874-0246-04		Q0144		03/15/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	4	EA	BO	PO	EA	1	GM	0.25	03/15/2006	99/99/9999						
63874-0246-06		Q0144		03/15/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6	EA	BO	PO	EA	1	GM	0.25	03/15/2006	99/99/9999						
63874-0246-10		Q0144		03/15/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	10	EA	BO	PO	EA	1	GM	0.25	03/15/2006	99/99/9999						
63874-0246-15		Q0144		03/15/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	15	EA	BO	PO	EA	1	GM	0.25	03/15/2006	99/99/9999						
63874-0327-01		J7506		05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	100	EA	BO	PO	EA	5	MG	2	05/10/2004	12/31/2015						
63874-0327-02		J7506		05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	1000	EA	BO	PO	EA	5	MG	2	05/10/2004	12/31/2015						
63874-0327-10		J7506		05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	10	EA	BO	PO	EA	5	MG	2	05/10/2004	12/31/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63874-0327-12	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	12 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-14	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	14 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-15	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	15 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-18	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	18 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-19	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	19 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-20	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	20 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-21	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	21 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-24	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	24 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-25	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	25 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-28	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	28 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-30	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-32	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	32 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-40	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	40 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-42	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	42 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-50	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	50 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0327-60	J7506			05/10/2004	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	60 EA	BO	PO	EA		5 MG		2	05/10/2004	12/31/2015						
63874-0370-01	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-08	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	8 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-10	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-12	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-15	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-20	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-24	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	24 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-30	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-40	Q0170			05/07/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	40 EA	BO	PO	EA		25 MG		1	05/07/2004	12/31/2013						
63874-0370-60	Q0170			03/02/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60 EA	NA	PO	EA		25 MG		1	03/02/2006	12/31/2013						
63874-0373-01	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	100 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-02	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	1000 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-10	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	10 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-15	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	15 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-20	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	20 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-21	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	21 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-30	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	30 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-33	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	33 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-36	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	36 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-40	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	40 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-50	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	50 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0373-60	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	60 EA	BO	PO	EA		5 MG		1	01/15/2006	12/31/2015						
63874-0392-01	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	100 EA	BO	PO	EA		5 MG		4	01/15/2006	12/31/2015						
63874-0392-02	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	1000 EA	BO	PO	EA		5 MG		4	01/15/2006	12/31/2015						
63874-0392-06	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	60 EA	BO	PO	EA		5 MG		4	01/15/2006	12/31/2015						
63874-0392-10	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	10 EA	BO	PO	EA		5 MG		4	01/15/2006	12/31/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63874-0392-14	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	14 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0392-15	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	15 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0392-20	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	20 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0392-21	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	21 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0392-24	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	24 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0392-28	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	28 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0392-30	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	30 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0392-40	J7506			01/15/2006	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	40 EA	BO PO EA	5 MG			4			01/15/2006	12/31/2015						
63874-0404-01	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	100 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-10	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	10 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-14	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	14 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-15	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	15 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-20	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	20 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-24	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	24 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-25	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-30	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-35	J8499			01/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	35 EA	BO PO EA	1 EA			1			01/15/2006	02/03/2016						
63874-0404-40	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-50	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	50 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0404-60	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	60 EA	BO PO EA	1 EA			1			01/23/2002	02/03/2016						
63874-0405-01	J8499			01/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100 EA	BO PO EA	1 EA			1			01/15/2006	02/03/2016						
63874-0405-10	J8499			01/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	10 EA	BO PO EA	1 EA			1			01/15/2006	02/03/2016						
63874-0405-20	J8499			01/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	20 EA	BO PO EA	1 EA			1			01/15/2006	02/03/2016						
63874-0405-25	J8499			01/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	25 EA	BO PO EA	1 EA			1			01/15/2006	02/03/2016						
63874-0405-30	J8499			01/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30 EA	BO PO EA	1 EA			1			01/15/2006	02/03/2016						
63874-0405-35	J8499			01/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	35 EA	BO PO EA	1 EA			1			01/15/2006	02/03/2016						
63874-0413-21	J7509			01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21 EA	DP PO EA	4 MG			1			01/01/2002	99/99/9999						
63874-0442-02	Q0177			05/11/2004	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	1000 EA	NA PO EA	25 MG			1			05/11/2004	99/99/9999						
63874-0442-03	Q0177			05/11/2004	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	500 EA	NA PO EA	25 MG			1			05/11/2004	99/99/9999						
63874-0442-04	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	120 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-05	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	5 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-09	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	9 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-10	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	10 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-14	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	14 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-15	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	15 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-20	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-25	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	25 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						
63874-0442-28	Q0177			05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	28 EA	BO PO EA	25 MG			1			05/11/2004	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63874-0442-30		Q0177		05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	BO	PO	EA	25	MG	1	05/11/2004	02/03/2016						
63874-0442-40		Q0177		05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	40	EA	BO	PO	EA	25	MG	1	05/11/2004	02/03/2016						
63874-0442-45		Q0177		05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	45	EA	BO	PO	EA	25	MG	1	05/11/2004	02/03/2016						
63874-0442-60		Q0177		05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	60	EA	BO	PO	EA	25	MG	1	05/11/2004	02/03/2016						
63874-0442-90		Q0177		05/11/2004	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	90	EA	BO	PO	EA	25	MG	1	05/11/2004	02/03/2016						
63874-0444-01	J8540			01/01/2006	02/03/2016	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (DOSE PAK) 0.75 MG	100	EA	BO	PO	EA	0.25	MG	3	01/01/2006	02/03/2016						
63874-0444-12	J8540			01/01/2006	02/03/2016	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (DOSE PAK) 0.75 MG	12	EA	BO	PO	EA	0.25	MG	3	01/01/2006	02/03/2016						
63874-0444-15	J8540			01/01/2006	02/03/2016	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (DOSE PAK) 0.75 MG	15	EA	BO	PO	EA	0.25	MG	3	01/01/2006	02/03/2016						
63874-0444-20	J8540			01/01/2006	02/03/2016	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (DOSE PAK) 0.75 MG	20	EA	BO	PO	EA	0.25	MG	3	01/01/2006	02/03/2016						
63874-0444-21	J8540			01/01/2006	02/03/2016	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (DOSE PAK) 0.75 MG	12	EA	BO	PO	EA	0.25	MG	3	01/01/2006	02/03/2016						
63874-0444-30	J8540			01/01/2006	02/03/2016	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (DOSE PAK) 0.75 MG	30	EA	BO	PO	EA	0.25	MG	3	01/01/2006	02/03/2016						
63874-0490-01		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-06		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	6	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-08		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	8	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-10		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-12		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	12	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-15		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-20		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-28		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	28	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-30		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0490-60		Q0165		05/10/2004	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	10	MG	1	05/10/2004	12/31/2013						
63874-0500-01	J8499			03/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100	EA	BO	PO	EA	1	EA	1	03/15/2006	02/03/2016						
63874-0500-15	J8499			01/23/2002	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15	EA	BO	PO	EA	1	EA	1	01/23/2002	02/03/2016						
63874-0500-20	J8499			03/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	20	EA	BO	PO	EA	1	EA	1	03/15/2006	02/03/2016						
63874-0500-21	J8499			03/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	21	EA	BO	PO	EA	1	EA	1	03/15/2006	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63874-0500-25	J8499			03/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	BO PO EA				1 EA		1	03/15/2006	02/03/2016						
63874-0500-30	J8499			03/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30 EA	BO PO EA				1 EA		1	03/15/2006	02/03/2016						
63874-0500-40	J8499			03/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	40 EA	BO PO EA				1 EA		1	03/15/2006	02/03/2016						
63874-0500-60	J8499			03/15/2006	02/03/2016	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	60 EA	NA PO EA				1 EA		1	03/15/2006	02/03/2016						
63874-0708-20	J7611			04/01/2008	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 1 MG PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ALBUTEROL SULFATE 0.5%	20 ML	NA IH ML				1 MG		5	04/01/2008	99/99/9999						
63874-0712-12	Q0170			01/01/2002	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120 ML	NA PO ML				25 MG		0.05	01/01/2002	12/31/2013						
63874-0757-01	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-04	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	120 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-10	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	10 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-15	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	15 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-20	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-21	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	21 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-24	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	24 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-28	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	28 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-30	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-60	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	60 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0757-90	Q0178			01/15/2006	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	90 EA	BO PO EA				50 MG		1	01/15/2006	12/31/2013						
63874-0806-12	J8498			01/15/2006	99/99/9999	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	12 EA	NA RC EA				1 EA		1	01/15/2006	99/99/9999						
64019-0750-85	J1230			01/01/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL	1 EA	BO NA GM				10 MG		100	01/01/2002	99/99/9999						
64019-0750-88	J1230			01/01/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL	1 EA	BO NA GM				10 MG		100	01/01/2002	99/99/9999						
64116-0011-12	J9216			01/01/2002	11/12/2013	INJECTION, INTERFERON, GAMMA 1-B, 3 MILLION UNITS	ACTIMMUNE (VIAL) 2 Million IU/0.5 ML	0.5 ML	VL SC ML				3 MU		1.33333	01/01/2002	11/12/2013						
64253-0111-21	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,6 ML W/LUER LOCK,PF) 0.9%	1 ML	SR IV ML				10 ML		0.1	01/01/2007	02/03/2016						
64253-0111-22	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,6 ML W/LUER LOCK,PF) 0.9%	2 ML	SR IV ML				10 ML		0.1	01/01/2007	02/03/2016						
64253-0111-23	A4216			01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,6 ML W/LUER LOCK,PF) 0.9%	3 ML	SR IV ML				10 ML		0.1	01/01/2007	99/99/9999						
64253-0111-25	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,6 ML W/LUER LOCK,PF) 0.9%	5 ML	SR IV ML				10 ML		0.1	01/01/2007	02/03/2016						
64253-0111-30	A4216			01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN W/LUER LOCK,PF) 0.9%	10 ML	SR IV ML				10 ML		0.1	01/01/2007	99/99/9999						
64253-0111-33	A4216			01/01/2007	02/03/2016	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,12 ML W/LUER LOK,PF) 0.9%	3 ML	SR IV ML				10 ML		0.1	01/01/2007	02/03/2016						
64253-0111-35	A4216			01/01/2007	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	NORMAL SALINE FLUSH (SRN,12 ML W/LUER LOK,PF) 0.9%	5 ML	SR IV ML				10 ML		0.1	01/01/2007	99/99/9999						
64253-0222-21	J1642			01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 10 U/ML-0.9%	1 ML	SR IV ML				10 U		1	01/01/2002	02/03/2016						
64253-0222-22	J1642			01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 10 U/ML-0.9%	2 ML	SR IV ML				10 U		1	01/01/2002	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
64253-0222-23		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 10 U/ML-0.9%	3 ML	SR	IV	ML		10 U		1	01/01/2002	02/03/2016							
64253-0222-25		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 10 U/ML-0.9%	5 ML	SR	IV	ML		10 U		1	01/01/2002	02/03/2016							
64253-0222-30		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,12 ML W/LUER LOCK) 10 U/ML-0.9%	10 ML	SR	IV	ML		10 U		1	01/01/2002	02/03/2016							
64253-0222-33		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,12 ML W/LUER LOCK) 10 U/ML-0.9%	3 ML	SR	IV	ML		10 U		1	01/01/2002	02/03/2016							
64253-0222-35		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 10 U/ML-0.9%	5 ML	SR	IV	ML		10 U		1	01/01/2002	99/99/9999							
64253-0333-21		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 100 U/ML-0.9%	1 ML	SR	IV	ML		10 U		10	01/01/2002	02/03/2016							
64253-0333-22		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 100 U/ML-0.9%	2 ML	SR	IV	ML		10 U		10	01/01/2002	02/03/2016							
64253-0333-23		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 100 U/ML-0.9%	3 ML	SR	IV	ML		10 U		10	01/01/2002	02/03/2016							
64253-0333-25		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,6 ML W/LUER LOCK) 100 U/ML-0.9%	5 ML	SR	IV	ML		10 U		10	01/01/2002	02/03/2016							
64253-0333-30		J1642		01/01/2002	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,12 ML W/LUER LOCK) 100 U/ML-0.9%	10 ML	SR	IV	ML		10 U		10	01/01/2002	02/03/2016							
64253-0333-33		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,12 ML W/LUER LOCK) 100 U/ML-0.9%	3 ML	SR	IV	ML		10 U		10	01/01/2002	99/99/9999							
64253-0333-35		J1642		01/01/2002	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (SRN,12 ML W/LUER LOCK) 100 U/ML-0.9%	5 ML	SR	IV	ML		10 U		10	01/01/2002	99/99/9999							
64253-0444-25		J1642		10/10/2003	12/08/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (6ML PRE-FILLED SYRINGE) 1 U/ML	5 ML	SR	IV	ML		10 U		0.1	10/10/2003	12/08/2016							
64281-0100-06		J7674		01/01/2005	99/99/9999	METHACHOLINE CHLORIDE ADMINISTERED AS INHALATION SOLUTION THROUGH A NEBULIZER, PER 1 MG	PROVOCHOLINE 100 MG NAVELBINE (1X1ML,SINGLE USE,PF) 10 MG/ML	1 EA	VL	IH	EA		1 MG		100	01/01/2005	99/99/9999							
64370-0532-01		J9390		06/23/2008	99/99/9999	INJECTION, VINORELBINE TARTRATE, 10 MG	NAVELBINE (1X5ML,SINGLE USE,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	06/23/2008	99/99/9999							
64370-0532-02		J9390		06/23/2008	99/99/9999	INJECTION, VINORELBINE TARTRATE, 10 MG	NAVELBINE (1X5ML,SINGLE USE,PF) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	06/23/2008	99/99/9999							
64679-0662-01		J1626		04/25/2008	05/31/2014	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (5X1ML,PF) 0.1 MG/ML	1 ML	VL	IV	ML		100 MCG		1	04/25/2008	05/31/2014							
64679-0701-02		J0696		05/18/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 250 MG	1 EA	VL	IJ	EA		250 MG		1	05/18/2007	99/99/9999							
64679-0701-03		J0696		05/18/2007	05/31/2014	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 250 MG	1 EA	VL	IJ	EA		250 MG		1	05/18/2007	05/31/2014							
64679-0702-02		J0696		05/18/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 500 MG	1 EA	VL	IJ	EA		250 MG		2	05/18/2007	99/99/9999							
64679-0703-01		J0696		05/18/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 2 GM	1 EA	VL	IJ	EA		250 MG		8	05/18/2007	99/99/9999							
64679-0726-01		J2405		12/26/2006	08/19/2013	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (5X2ML,SDV,USP) 2 MG/ML	2 ML	VL	IJ	ML		1 MG		2	12/26/2006	08/19/2013							
64679-0727-01		J2405		12/26/2006	08/19/2013	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV,USP) 2 MG/ML	20 ML	VL	IJ	ML		1 MG		2	12/26/2006	08/19/2013							
64679-0757-01		J1885		04/12/2007	08/19/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP,SDV) 15 MG/ML	1 ML	VL	IJ	ML		15 MG		1	04/12/2007	08/19/2013							
64679-0757-02		J1885		04/12/2007	08/19/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP,SDV) 15 MG/ML	1 ML	VL	IJ	ML		15 MG		1	04/12/2007	08/19/2013							
64679-0758-01		J1885		04/12/2007	08/19/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP,SDV) 30 MG/ML	1 ML	VL	IJ	ML		15 MG		2	04/12/2007	08/19/2013							
64679-0758-02		J1885		04/12/2007	08/19/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP,SDV,25X2ML) 30 MG/ML	2 ML	VL	IJ	ML		15 MG		2	04/12/2007	08/19/2013							
64679-0758-04		J1885		04/12/2007	08/19/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP,SDV) 30 MG/ML	1 ML	VL	IJ	ML		15 MG		2	04/12/2007	08/19/2013							
64679-0758-06		J1885		04/12/2007	08/19/2013	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (ASP,SDV,2X10ML) 30 MG/ML	2 ML	VL	IJ	ML		15 MG		2	04/12/2007	08/19/2013							
64679-0964-03		Q0144		02/14/2008	05/31/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM COATED) 500 MG	3 EA	BX	PO	EA		1 GM		0.5	02/14/2008	05/31/2014							
64679-0983-02		J0698		05/26/2006	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 1 GM	1 EA	VL	IJ	EA		250 MG		4	05/26/2006	99/99/9999							
64679-0986-01		J0698		09/20/2006	05/31/2014	INJECTION, CEFOTAXIME SODIUM, PER GM	CEFOTAXIME (USP) 1 GM	1 EA	VL	IJ	EA		1 GM		1	09/20/2006	05/31/2014							
64679-0986-02		J0698		09/20/2006	05/31/2014	INJECTION, CEFOTAXIME SODIUM, PER GM	CEFOTAXIME (USP) 1 GM	1 EA	VL	IJ	EA		1 GM		1	09/20/2006	05/31/2014							
64679-0986-03		J0698		09/20/2006	05/31/2014	INJECTION, CEFOTAXIME SODIUM, PER GM	CEFOTAXIME (USP) 1 GM	1 EA	VL	IJ	EA		1 GM		1	09/20/2006	05/31/2014							
64679-0986-04		J0698		09/20/2006	05/31/2014	INJECTION, CEFOTAXIME SODIUM, PER GM	CEFOTAXIME (USP) 1 GM	1 EA	VL	IJ	EA		1 GM		1	09/20/2006	05/31/2014							
64720-0198-02		Q0166		12/29/2007	08/20/2014	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (FILM-COATED) 1 MG	20 EA	BO	PO	EA		1 MG		1	12/29/2007	08/20/2014							
64720-0198-98		Q0166		12/29/2007	08/20/2014	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (FILM-COATED) 1 MG	2 EA	DP	PO	EA		1 MG		1	12/29/2007	08/20/2014							
65293-0001-01		J0583		01/01/2004	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	ANGIOMAX (VIAL, GLASS) 250 MG	1 EA	VL	IV	EA		1 MG		250	01/01/2004	99/99/9999							
65483-0590-10		J7500		01/01/2002	12/31/2017	AZATHIOPRINE, ORAL, 50 MG	IMURAN 50 MG	100 EA	BO	PO	EA		50 MG		1	01/01/2002	12/31/2017							
65580-0251-01		J7510		05/09/2002	09/28/2012	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (DYE-FREE,RASPBERRY) 6.7 MG/5 ML	120 ML	BO	PO	ML		5 MG		0.268	05/09/2002	09/28/2012							
65649-0231-41		J7500		10/31/2003	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZASAN 75 MG	100 EA	BO	PO	EA		50 MG		1.5	10/31/2003	99/99/9999							
65649-0241-41		J7500		10/31/2003	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZASAN 100 MG	100 EA	BO	PO	EA		50 MG		2	10/31/2003	99/99/9999							
65847-0205-25		J2325		01/01/2006	99/99/9999	INJECTION, NESIRITIDE, 0.1 MG	NATRECOR (S.D.V.) 1.5 MG	1 EA	VL	IV	EA		0.1 MG		15	01/01/2006	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
66105-0507-01		Q0144		08/22/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	10 EA	BO PO EA				1 GM		0.25	08/22/2006	99/99/9999						
66105-0507-03		Q0144		01/01/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	30 EA	BO PO EA				1 GM		0.25	01/01/2006	99/99/9999						
66105-0507-06		Q0144		08/22/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	60 EA	BO PO EA				1 GM		0.25	08/22/2006	99/99/9999						
66105-0507-09		Q0144		08/22/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	90 EA	BO PO EA				1 GM		0.25	08/22/2006	99/99/9999						
66105-0507-10		Q0144		08/22/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	100 EA	BO PO EA				1 GM		0.25	08/22/2006	99/99/9999						
66105-0549-10		J7507		01/01/2006	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	PROGRAF 1 MG	100 EA	NA PO EA				1 MG		1	01/01/2006	99/99/9999						
66105-0653-01		Q0144		09/13/2006	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	10 EA	BO PO EA				1 GM		0.5	09/13/2006	02/03/2016						
66105-0653-03		Q0144		09/13/2006	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	30 EA	BO PO EA				1 GM		0.5	09/13/2006	02/03/2016						
66105-0653-05		Q0144		09/13/2006	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	50 EA	BO PO EA				1 GM		0.5	09/13/2006	02/03/2016						
66105-0653-06		Q0144		09/13/2006	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	60 EA	BO PO EA				1 GM		0.5	09/13/2006	02/03/2016						
66105-0653-19		Q0144		09/13/2006	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	9 EA	BO PO EA				1 GM		0.5	09/13/2006	02/03/2016						
66105-0670-01		Q0144		09/13/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	10 EA	BO PO EA				1 GM		0.25	09/13/2006	99/99/9999						
66105-0670-03		Q0144		09/13/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	30 EA	BO PO EA				1 GM		0.25	09/13/2006	99/99/9999						
66105-0670-05		Q0144		09/13/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	50 EA	BO PO EA				1 GM		0.25	09/13/2006	99/99/9999						
66105-0670-06		Q0144		09/13/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	60 EA	BO PO EA				1 GM		0.25	09/13/2006	99/99/9999						
66105-0670-18		Q0144		09/13/2006	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	18 EA	BO PO EA				1 GM		0.25	09/13/2006	99/99/9999						
66105-0832-01		J8999		09/13/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	NOLVADEX 10 MG	10 EA	BO PO EA				1 EA		1	09/13/2006	99/99/9999						
66105-0832-03		J8999		09/13/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	NOLVADEX 10 MG	30 EA	BO PO EA				1 EA		1	09/13/2006	99/99/9999						
66105-0832-06		J8999		09/13/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	NOLVADEX 10 MG	60 EA	BO PO EA				1 EA		1	09/13/2006	99/99/9999						
66105-0832-09		J8999		09/13/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	NOLVADEX 10 MG	90 EA	BO PO EA				1 EA		1	09/13/2006	99/99/9999						
66105-0832-10		J8999		09/13/2006	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	NOLVADEX 10 MG	100 EA	BO PO EA				1 EA		1	09/13/2006	99/99/9999						
66267-0006-25		J8499		04/08/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	25 EA	BO PO EA				1 EA		1	04/08/2002	99/99/9999						
66267-0006-40		J8499		08/01/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40 EA	BO PO EA				1 EA		1	08/01/2002	99/99/9999						
66267-0006-50		J8499		04/08/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	50 EA	BO PO EA				1 EA		1	04/08/2002	99/99/9999						
66267-0007-15		J8499		04/08/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15 EA	BO PO EA				1 EA		1	04/08/2002	99/99/9999						
66267-0007-21		J8499		04/08/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	BO PO EA				1 EA		1	04/08/2002	99/99/9999						
66267-0007-25		J8499		04/08/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25 EA	BO PO EA				1 EA		1	04/08/2002	99/99/9999						
66267-0007-30		J8499		04/08/2002	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	30 EA	BO PO EA				1 EA		1	04/08/2002	99/99/9999						
66267-0066-12		J8540		01/01/2006	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	12 EA	BO PO EA				0.25 MG		3	01/01/2006	99/99/9999						
66267-0080-15		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15 EA	BO PO EA				50 MG		0.5	01/01/2002	99/99/9999						
66267-0080-20		Q0163		04/05/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20 EA	BO PO EA				50 MG		0.5	04/05/2002	99/99/9999						
66267-0080-30		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30 EA	BO PO EA				50 MG		0.5	01/01/2002	99/99/9999						
66267-0080-60		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60 EA	BO PO EA				50 MG		0.5	01/01/2002	99/99/9999						
66267-0081-15		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15 EA	BO PO EA				50 MG		1	01/01/2002	99/99/9999						
66267-0081-20		Q0163		04/05/2002	10/17/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	20 EA	BO PO EA				50 MG		1	04/05/2002	10/17/2016						
66267-0081-30		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30 EA	BO PO EA				50 MG		1	01/01/2002	99/99/9999						
66267-0081-60		Q0163		09/04/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	60 EA	BO PO EA				50 MG		1	09/04/2002	99/99/9999						
66267-0171-15		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	15 EA	BO PO EA				5 MG		2	01/01/2002	12/31/2015						
66267-0171-20		J7506		04/04/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	20 EA	BO PO EA				5 MG		2	04/04/2002	12/31/2015						
66267-0171-21		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	21 EA	BO PO EA				5 MG		2	01/01/2002	12/31/2015						
66267-0171-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30 EA	BO PO EA				5 MG		2	01/01/2002	12/31/2015						
66267-0171-40		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	40 EA	BO PO EA				5 MG		2	01/01/2002	12/31/2015						
66267-0171-42		J7506		04/04/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	42 EA	BO PO EA				5 MG		2	04/04/2002	12/31/2015						
66267-0172-10		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	10 EA	BO PO EA				5 MG		4	01/01/2002	12/31/2015						
66267-0172-15		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	15 EA	BO PO EA				5 MG		4	01/01/2002	12/31/2015						
66267-0172-20		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	20 EA	BO PO EA				5 MG		4	01/01/2002	12/31/2015						
66267-0172-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	30 EA	BO PO EA				5 MG		4	01/01/2002	12/31/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
66267-0173-20		J7506		04/04/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	20	EA	BO	PO	EA	5	MG	1	04/04/2002	12/31/2015						
66267-0173-30		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	30	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
66267-0173-40		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	40	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
66267-0173-42		J7506		03/24/2003	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	42	EA	BO	PO	EA	5	MG	1	03/24/2003	12/31/2015						
66267-0173-60		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	60	EA	BO	PO	EA	5	MG	1	01/01/2002	12/31/2015						
66267-0208-10		Q0173		01/01/2002	10/17/2016	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	10	EA	BO	PO	EA	250	MG	1	01/01/2002	10/17/2016						
66267-0208-20		Q0173		01/01/2002	10/17/2016	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	20	EA	BO	PO	EA	250	MG	1	01/01/2002	10/17/2016						
66267-0399-30		J8499		03/15/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	30	EA	BO	PO	EA	1	EA	1	03/15/2005	99/99/9999						
66267-0928-06		Q0144		01/01/2002	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	ZITHROMAX 250 MG	6	EA	BO	PO	EA	1	GM	0.25	01/01/2002	99/99/9999						
66267-0948-21		J7506		01/01/2002	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE (DOSEPACK) 5 MG	21	EA	DP	PO	EA	5	MG	1	01/01/2002	12/31/2015						
66267-0961-21		J7509		01/01/2002	99/99/9999	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21	EA	BO	PO	EA	4	MG	1	01/01/2002	99/99/9999						
66267-0977-04		Q0163		01/01/2002	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	120	ML	BO	PO	ML	50	MG	0.05	01/01/2002	99/99/9999						
66288-1100-01		J0690		10/01/2002	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM 100 GM	1	EA	FC	IJ	GM	500	MG	2	10/01/2002	99/99/9999						
66288-1300-01		J0690		10/01/2002	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN SODIUM 300 GM	1	EA	FC	IJ	GM	500	MG	2	10/01/2002	99/99/9999						
66302-0101-01		J3285		01/01/2006	99/99/9999	INJECTION, TREPROSTINIL, 1 MG	REMODULIN (M.D.V.) 1 MG/ML	20	ML	VL	IJ	ML	1	MG	1	01/01/2006	99/99/9999						
66302-0102-01		J3285		01/01/2006	99/99/9999	INJECTION, TREPROSTINIL, 1 MG	REMODULIN (M.D.V.) 2.5 MG/ML	20	ML	VL	IJ	ML	1	MG	2.5	01/01/2006	99/99/9999						
66302-0105-01		J3285		01/01/2006	99/99/9999	INJECTION, TREPROSTINIL, 1 MG	REMODULIN (M.D.V.) 5 MG/ML	20	ML	VL	IJ	ML	1	MG	5	01/01/2006	99/99/9999						
66302-0110-01		J3285		01/01/2006	99/99/9999	INJECTION, TREPROSTINIL, 1 MG	REMODULIN (M.D.V.) 10 MG/ML	20	ML	VL	IJ	ML	1	MG	10	01/01/2006	99/99/9999						
66336-0045-06		Q0163		10/22/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	6	EA	BO	PO	EA	50	MG	1	10/22/2004	06/01/2014						
66336-0045-15		Q0163		10/22/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15	EA	BO	PO	EA	50	MG	1	10/22/2004	06/01/2014						
66336-0045-30		Q0163		11/23/2003	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	30	EA	BO	PO	EA	50	MG	1	11/23/2003	06/01/2014						
61755-0008-01		J9999		09/28/2018	99/99/9999	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	LIBTAYO 50 MG/1 ML	7	ML	VL	IV	ML	1	MG	1	09/28/2018	99/99/9999						
66336-0058-10		J7506		10/22/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	10	EA	BO	PO	EA	5	MG	2	10/22/2004	06/01/2014						
66336-0058-12		J7506		11/04/2005	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	12	EA	BO	PO	EA	5	MG	2	11/04/2005	06/01/2014						
66336-0058-20		J7506		10/22/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	20	EA	BO	PO	EA	5	MG	2	10/22/2004	06/01/2014						
66336-0058-21		J7506		10/22/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	21	EA	BO	PO	EA	5	MG	2	10/22/2004	06/01/2014						
66336-0058-30		J7506		04/16/2002	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	30	EA	BO	PO	EA	5	MG	2	04/16/2002	06/01/2014						
66336-0058-60		J7506		10/22/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	60	EA	BO	PO	EA	5	MG	2	10/22/2004	06/01/2014						
66336-0085-10		Q0170		10/22/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10	EA	BO	PO	EA	25	MG	1	10/22/2004	12/31/2013						
66336-0085-12		Q0170		10/22/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	12	EA	BO	PO	EA	25	MG	1	10/22/2004	12/31/2013						
66336-0085-20		Q0170		05/29/2008	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	20	EA	BO	PO	EA	25	MG	1	05/29/2008	12/31/2013						
66336-0085-25		Q0170		05/29/2008	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	25	EA	BO	PO	EA	25	MG	1	05/29/2008	12/31/2013						
66336-0085-30		Q0170		10/22/2004	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	30	EA	BO	PO	EA	25	MG	1	10/22/2004	12/31/2013						
66336-0085-60		Q0170		05/29/2008	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60	EA	BO	PO	EA	25	MG	1	05/29/2008	12/31/2013						
66336-0094-10		J7506		10/22/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	10	EA	BO	PO	EA	5	MG	4	10/22/2004	06/01/2014						
66336-0094-18		J7506		10/22/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	18	EA	BO	PO	EA	5	MG	4	10/22/2004	06/01/2014						
66336-0094-20		J7506		10/22/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	20	EA	BO	PO	EA	5	MG	4	10/22/2004	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
66336-0094-30	J7506			10/22/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	30	EA	BO	PO	EA	5	MG	4	10/22/2004	06/01/2014						
66336-0150-03	J8498			01/01/2006	06/01/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	3	EA	BO	RC	EA	1	EA	1	01/01/2006	06/01/2014						
66336-0150-06	J8498			04/20/2007	06/01/2014	ANTIEMETIC DRUG, RECTAL/SUPPOSITORY, NOT OTHERWISE SPECIFIED	PROCHLORPERAZINE 25 MG	6	EA	BX	RC	EA	1	EA	1	04/20/2007	06/01/2014						
66336-0208-20	Q0177			10/22/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20	EA	BO	PO	EA	25	MG	1	10/22/2004	06/01/2014						
66336-0208-30	Q0177			10/22/2004	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30	EA	BO	PO	EA	25	MG	1	10/22/2004	06/01/2014						
66336-0400-05	Q0144			12/03/2007	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 500 MG	5	EA	BO	PO	EA	1	GM	0.5	12/03/2007	06/01/2014						
66336-0434-06	Q0164			10/22/2004	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 5 MG	6	EA	BO	PO	EA	5	MG	1	10/22/2004	06/01/2014						
66336-0434-10	Q0164			08/18/2005	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 5 MG	10	EA	BO	PO	EA	5	MG	1	08/18/2005	06/01/2014						
66336-0479-06	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 4 MG	6	EA	BO	PO	EA	0.25	MG	16	01/01/2006	06/01/2014						
66336-0515-21	J7506			10/22/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	21	EA	BO	PO	EA	5	MG	1	10/22/2004	06/01/2014						
66336-0515-30	J7506			10/22/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	30	EA	BO	PO	EA	5	MG	1	10/22/2004	06/01/2014						
66336-0515-40	J7506			10/22/2004	06/01/2014	PREDNISONE, ORAL, PER 5MG	PREDNISONE 5 MG	40	EA	BO	PO	EA	5	MG	1	10/22/2004	06/01/2014						
66336-0550-12	J8540			01/01/2006	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 0.75 MG	12	EA	BO	PO	EA	0.25	MG	3	01/01/2006	06/01/2014						
66336-0589-15	Q0163			01/01/2002	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	15	EA	BO	PO	EA	50	MG	0.5	01/01/2002	06/01/2014						
66336-0589-20	Q0163			10/22/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	20	EA	BO	PO	EA	50	MG	0.5	10/22/2004	06/01/2014						
66336-0589-30	Q0163			10/22/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	30	EA	BO	PO	EA	50	MG	0.5	10/22/2004	06/01/2014						
66336-0589-60	Q0163			10/22/2004	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 25 MG	60	EA	BO	PO	EA	50	MG	0.5	10/22/2004	06/01/2014						
66336-0642-30	J8499			06/22/2005	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	30	EA	BO	PO	EA	1	EA	1	06/22/2005	06/01/2014						
66336-0642-40	J8499			10/22/2004	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	40	EA	BO	PO	EA	1	EA	1	10/22/2004	06/01/2014						
66336-0642-50	J8499			01/07/2008	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	50	EA	BO	PO	EA	1	EA	1	01/07/2008	06/01/2014						
66336-0735-15	J8499			10/22/2004	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	15	EA	BO	PO	EA	1	EA	1	10/22/2004	06/01/2014						
66336-0735-25	J8499			10/22/2004	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	25	EA	BO	PO	EA	1	EA	1	10/22/2004	06/01/2014						
66336-0735-40	J8499			10/22/2004	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	40	EA	BO	PO	EA	1	EA	1	10/22/2004	06/01/2014						
66336-0862-50	J8499			05/01/2006	06/01/2014	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	DISPENSEQUICK ACYCLOVIR 800 MG	50	EA	BO	PO	EA	1	EA	1	05/01/2006	06/01/2014						
66336-0921-15	Q0165			12/03/2007	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	10	MG	1	12/03/2007	12/31/2013						
66336-0921-60	Q0165			05/29/2008	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	10	MG	1	05/29/2008	12/31/2013						
66479-0520-01	J0735			06/28/2006	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	DURACLON (SDV,PF) 0.1 MG/ML	10	ML	VL	EP	ML	1	MG	0.1	06/28/2006	99/99/9999						
66479-0521-01	J0735			06/14/2006	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	DURACLON (SDV,PF) 0.5 MG/ML	10	ML	VL	EP	ML	1	MG	0.5	06/14/2006	99/99/9999						
66490-0041-01	J1110			12/31/2002	99/99/9999	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG	D.H.E. 45 (AMP) 1 MG/ML	1	ML	AM	IJ	ML	1	MG	1	12/31/2002	99/99/9999						
66657-0301-05	J1457			01/01/2005	09/05/2013	INJECTION, GALLIUM NITRATE, 1 MG	GANITE (PF) 25 MG/ML	20	ML	VL	IV	ML	1	MG	25	01/01/2005	09/05/2013						
66689-0681-55	J1230			02/01/2002	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL	1	EA	BO	NA	GM	10	MG	100	02/01/2002	99/99/9999						
50090-0294-09	None			06/08/2018	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	36	EA	BO	PO	EA	2.5	MG	1	06/08/2018	99/99/9999						
66733-0948-23	J9055			01/01/2005	99/99/9999	INJECTION, CETUXIMAB, 10 MG	ERBITUX (PF) 2 MG/ML	50	ML	VL	IV	ML	10	MG	0.2	01/01/2005	99/99/9999						
66733-0958-23	J9055			05/03/2007	99/99/9999	INJECTION, CETUXIMAB, 10 MG	ERBITUX (PF) 2 MG/ML	100	ML	VL	IV	ML	10	MG	0.2	05/03/2007	99/99/9999						
66758-0016-04	J2370			06/08/2005	03/31/2016	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	PHENYLEPHRINE HCL (USP,25X5ML,PF) 10 MG/ML	5	ML	VL	IJ	ML	1	ML	1	06/08/2005	03/31/2016						
66758-0017-01	J2370			01/08/2004	03/31/2016	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	PHENYLEPHRINE HCL (USP, BULK PACKAGE,PF) 10 MG/ML	10	ML	VL	IJ	ML	1	ML	1	01/08/2004	03/31/2016						
66758-0043-01	J9265			01/11/2008	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (USP,1X5ML,MULTI-DOSE) 6 MG/ML	5	ML	VL	IV	ML	30	MG	0.2	01/11/2008	12/31/2014						
66758-0043-02	J9265			01/11/2008	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (USP,1X16.7ML,MULTI-DOSE) 6 MG/ML	16.7	ML	VL	IV	ML	30	MG	0.2	01/11/2008	12/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
66758-0043-03	J9265			01/11/2008	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (USP,1X50ML,MULTI-DOSE) 6 MG/ML	50 ML	VL	IV	ML		30 MG		0.2	01/11/2008	12/31/2014						
66758-0045-01	J9390			03/05/2008	10/06/2014	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE (1X1ML,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	03/05/2008	10/06/2014						
66758-0045-02	J9390			03/05/2008	10/06/2014	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE (1X5ML,PF) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	03/05/2008	10/06/2014						
66758-0046-01	J9185			10/12/2007	99/99/9999	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE (SDV,PF) 25 MG/ML	2 ML	VL	IV	ML		50 MG		0.5	10/12/2007	99/99/9999						
47781-0606-94	J9045			04/02/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	60 ML	VL	IV	ML		50 MG		0.2	04/02/2018	99/99/9999						
50242-0082-03	J2778			04/23/2018	99/99/9999	INJECTION, RANIBIZUMAB, 0.1 MG	LUCENTIS (INTRAVITREAL,PF) 0.3 MG/0.05 ML	0.05 ML	VL	IO	ML		0.1 MG		60	04/23/2018	99/99/9999						
67253-0101-10	J8499			10/01/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	100 EA	BO	PO	EA		1 EA		1	10/01/2003	99/99/9999						
67253-0101-11	J8499			07/15/2003	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 400 MG	1000 EA	BO	PO	EA		1 EA		1	07/15/2003	99/99/9999						
50742-0428-02	J9171			04/13/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (1X2ML,SINGLE-USE) 10 MG/1 ML	2 ML	VL	IV	ML		1 MG		10	04/13/2018	99/99/9999						
50742-0431-08	J9171			04/13/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (1X8ML,SINGLE-USE) 10 MG/1 ML	8 ML	VL	IV	ML		1 MG		10	04/13/2018	99/99/9999						
50742-0463-16	J9171			04/13/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (1X16ML,SINGLE-USE) 10 MG/1 ML	16 ML	VL	IV	ML		1 MG		10	04/13/2018	99/99/9999						
51754-1000-04	J3475			04/24/2018	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (SDV,PF) 500 MG/1 ML	10 ML	VL	IJ	ML		500 MG		1	04/24/2018	99/99/9999						
67253-0320-10	None			12/30/2005	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE (USP) 2.5 MG	100 EA	BO	PO	EA		2.5 MG		1	10/29/2007	99/99/9999	12/30/2005	01/01/2007	1			
67253-0580-42	None			07/01/2003	09/23/2016	METHOTREXATE, 2.5 MG, ORAL	RHEUMATREX DOSE PACK (4X2) 2.5 MG	8 EA	DP	PO	EA		2.5 MG		1	07/01/2003	09/23/2016						
67253-0580-43	None			07/01/2003	09/23/2016	METHOTREXATE, 2.5 MG, ORAL	RHEUMATREX DOSE PACK (4X3) 2.5 MG	12 EA	DP	PO	EA		2.5 MG		1	07/01/2003	09/23/2016						
67253-0580-44	None			07/01/2003	09/23/2016	METHOTREXATE, 2.5 MG, ORAL	RHEUMATREX DOSE PACK (4X4) 2.5 MG	16 EA	DP	PO	EA		2.5 MG		1	07/01/2003	09/23/2016						
67253-0580-45	None			07/01/2003	09/23/2016	METHOTREXATE, 2.5 MG, ORAL	RHEUMATREX DOSE PACK (4X5) 2.5 MG	20 EA	DP	PO	EA		2.5 MG		1	07/01/2003	09/23/2016						
67253-0580-46	None			07/01/2003	09/23/2016	METHOTREXATE, 2.5 MG, ORAL	RHEUMATREX DOSE PACK (4X6) 2.5 MG	24 EA	DP	PO	EA		2.5 MG		1	07/01/2003	09/23/2016						
67386-0501-52	J2515			06/11/2003	05/31/2012	INJECTION, PENTOBARBITAL SODIUM, PER 50 MG	NEMBUTAL SODIUM 50 MG/ML	20 ML	VL	IJ	ML		50 MG		1	06/11/2003	05/31/2012						
67386-0501-55	J2515			06/11/2003	05/31/2012	INJECTION, PENTOBARBITAL SODIUM, PER 50 MG	NEMBUTAL SODIUM (VIAL) 50 MG/ML	50 ML	VL	IJ	ML		50 MG		1	06/11/2003	05/31/2012						
67386-0611-52	J0515			01/21/2006	06/19/2012	INJECTION, BENZTROPINE MESYLATE, PER 1 MG	COGENTIN (5X2ML) 1 MG/ML	2 ML	AM	IJ	ML		1 MG		1	01/21/2006	06/19/2012						
67386-0701-54	J1640			01/01/2006	10/31/2013	INJECTION, HEMIN, 1 MG	PANHEMATIN 313 MG	1 EA	VL	IV	EA		1 MG		313	01/01/2006	10/31/2013						
67386-0811-55	J9120			01/21/2006	10/31/2013	INJECTION, DACTINOMYCIN, 0.5 MG	COSMEGEN 0.5 MG	1 EA	VL	IV	EA		0.5 MG		1	01/21/2006	10/31/2013						
67386-0911-51	J9230			01/21/2006	10/31/2013	INJECTION, MECHLORETHAMINE HYDROCHLORIDE, (NITROGEN MUSTARD), 10 MG	MUSTARGEN 10 MG	1 EA	VL	IV	EA		10 MG		1	01/21/2006	10/31/2013						
67425-0002-10	J3470			01/28/2005	04/21/2013	INJECTION, HYALURONIDASE, UP TO 150 UNITS	VITRASE (LYOPHILIZED,OVINE,SDV) 200 U/ML	1.2 ML	VL	SC	ML		150 U		1.33333	01/28/2005	04/21/2013						
67457-0124-10	J1200			05/01/2007	99/99/9999	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HYDROCHLORIDE (MDV,USP) 50 MG/ML	10 ML	VL	IJ	ML		50 MG		1	05/01/2007	99/99/9999						
67457-0153-03	J0282			07/01/2005	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL 50 MG/ML	3 ML	VL	IV	ML		30 MG		1.66666	07/01/2005	99/99/9999						
67457-0153-09	J0282			11/29/2005	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HYDROCHLORIDE (9X10ML) 50 MG/ML	9 ML	VL	IV	ML		30 MG		1.66666	11/29/2005	99/99/9999						
67457-0153-18	J0282			11/29/2005	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HYDROCHLORIDE 50 MG/ML	18 ML	VL	IV	ML		30 MG		1.66666	11/29/2005	99/99/9999						
67457-0177-50	J1212			06/22/2007	99/99/9999	INJECTION, DMSO, DIMETHYL SULFOXIDE, 50%, 50 ML	RIMSO-50 (ODORLESS) 50%	50 ML	VL	IL	ML		50 %		0.02	06/22/2007	99/99/9999						
67871-0007-10	J9175			01/01/2006	06/04/2013	INJECTION, ELLIOTT'S B SOLUTION, 1 ML	ELLIOTT'S B (FOR INTRATHECAL USE,PF)	10 ML	AM	IN	ML		1 ML		1	01/01/2006	06/04/2013						
67871-4790-06	J1430			01/01/2006	99/99/9999	INJECTION, ETHANOLAMINE OLEATE, 100 MG	ETHAMOLIN (10X2ML AMP) 50 MG/ML	2 ML	AM	IV	ML		100 MG		0.5	01/01/2006	99/99/9999						
67919-0011-01	J0878			01/01/2005	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	CUBICIN (PF) 500 MG	1 EA	VL	IV	EA		1 MG		500	01/01/2005	99/99/9999						
67979-0001-01	J9357			10/31/2007	99/99/9999	INJECTION, VALZUROBIN, INTRAVESICAL, 200 MG	VALSTAR (4X5ML,PF) 40 MG/ML	5 ML	VL	IL	ML		200 MG		0.2	06/03/2009	99/99/9999	10/31/2007	03/03/2009	0.2			
67979-0002-01	J9226			01/01/2008	99/99/9999	HISTRELIN IMPLANT (SUPPRELIN LA), 50 MG	SUPPRELIN LA 50 MG	1 EA	BX	SC	EA		50 MG		1	01/01/2008	99/99/9999						
67979-0500-01	J9226			01/01/2008	99/99/9999	HISTRELIN IMPLANT (SUPPRELIN LA), 50 MG	VANTAS 50 MG	1 EA	BX	SC	EA		50 MG		1	01/01/2008	99/99/9999						
68094-0518-59	J8999			07/01/2007	04/30/2015	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (1X20ML,LEMON-LIME) 40 MG/ML	20 ML	CP	PO	ML		1 EA		1	07/01/2007	04/30/2015						
68094-0518-62	J8999			11/28/2006	04/30/2015	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (30X20ML,LEMON-LIME) 40 MG/ML	20 ML	CP	PO	ML		1 EA		1	11/28/2006	04/30/2015						
68094-0528-59	J8999			07/01/2007	12/31/2014	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (1X10ML,LEMON-LIME) 40 MG/ML	10 ML	CP	PO	ML		1 EA		1	07/01/2007	12/31/2014						
68094-0528-61	J8999			02/26/2004	12/31/2014	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE (10X10) 40 MG/ML	10 ML	CP	PO	ML		1 EA		1	02/26/2004	12/31/2014						
68094-0528-62	J8999			02/26/2004	12/31/2014	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	MEGESTROL ACETATE 40 MG/ML	10 ML	CP	PO	ML		1 EA		1	02/26/2004	12/31/2014						
68115-0770-02	J3030			01/20/2004	02/03/2016	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	IMITREX (SRN,PREFILLED,UNIT/USE) 6 MG/0.5 ML	0.5 ML	BX	SC	ML		6 MG		2	01/20/2004	02/03/2016						
68135-0020-01	J1458			01/01/2007	99/99/9999	INJECTION, GALSULFASE, 1 MG	NAGLAZYME (PF) 1 MG/ML	5 ML	VL	IV	ML		1 MG		1	01/01/2007	99/99/9999						
68180-0611-10	J0696			07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 250 MG	1 EA	VL	IJ	EA		250 MG		1	07/20/2005	99/99/9999						
68180-0611-10	J0696			07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 250 MG	1 EA	VL	IJ	EA		250 MG		1	07/20/2005	99/99/9999						
68180-0622-01	J0696			07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 500 MG	1 EA	NA	IJ	EA		250 MG		2	07/20/2005	99/99/9999						
68180-0622-10	J0696			07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 500 MG	1 EA	NA	IJ	EA		250 MG		2	07/20/2005	99/99/9999						
68180-0633-01	J0696			07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 1 GM	1 EA	VL	IJ	EA		250 MG		4	07/20/2005	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
68180-0633-10		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 1 GM	10	EA	VL	IJ	EA	250	MG	4	07/20/2005	99/99/9999							
68180-0644-01		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 2 GM	1	EA	NA	IJ	EA	250	MG	8	07/20/2005	99/99/9999							
68180-0644-10		J0696		07/20/2005	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE 2 GM	1	EA	NA	IJ	EA	250	MG	8	07/20/2005	99/99/9999							
68330-0001-01		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 250 MG	1	EA	VL	IJ	EA	250	MG	1	09/15/2007	99/99/9999							
68330-0001-10		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 250 MG	1	EA	VL	IJ	EA	250	MG	1	09/15/2007	99/99/9999							
68330-0002-01		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 500 MG	1	EA	VL	IJ	EA	250	MG	2	09/15/2007	99/99/9999							
68330-0002-10		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 500 MG	1	EA	VL	IJ	EA	250	MG	2	09/15/2007	99/99/9999							
68330-0003-01		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 1 GM	1	EA	VL	IJ	EA	250	MG	4	09/15/2007	99/99/9999							
68330-0003-10		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 1 GM	1	EA	VL	IJ	EA	250	MG	4	09/15/2007	99/99/9999							
68330-0004-01		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 2 GM	1	EA	VL	IJ	EA	250	MG	8	09/15/2007	99/99/9999							
68330-0004-10		J0696		09/15/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP) 2 GM	1	EA	VL	IJ	EA	250	MG	8	09/15/2007	99/99/9999							
68330-0005-01		J0696		11/05/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP,PIGGYBACK) 1 GM	1	EA	GC	IJ	EA	250	MG	4	11/05/2007	99/99/9999							
68330-0006-01		J0696		11/05/2007	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (USP,PIGGYBACK) 2 GM	1	EA	GC	IJ	EA	250	MG	8	11/05/2007	99/99/9999							
68382-0003-01		J7500		05/01/2007	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE (USP) 50 MG	100	EA	BO	PO	EA	50	MG	1	05/01/2007	99/99/9999							
68382-0003-05		J7500		05/01/2007	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE (USP) 50 MG	500	EA	BO	PO	EA	50	MG	1	05/01/2007	99/99/9999							
68382-0040-01		Q0169		12/01/2005	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 12.5 MG	100	EA	BO	PO	EA	12.5	MG	1	12/01/2005	99/99/9999							
68382-0041-01		Q0170		12/01/2005	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	100	EA	BO	PO	EA	25	MG	1	12/01/2005	12/31/2013							
68382-0041-10		Q0170		02/27/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	1000	EA	BO	PO	EA	25	MG	1	02/27/2007	12/31/2013							
68387-0170-01		J7509		03/26/2004	06/01/2014	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE 4 MG	21	EA	DP	PO	EA	4	MG	1	03/26/2004	06/01/2014							
68387-0240-25		J7506		03/26/2004	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	25	EA	BO	PO	EA	5	MG	4	03/26/2004	06/01/2014							
68387-0468-30		Q0178		03/01/2007	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30	EA	BO	PO	EA	50	MG	1	03/01/2007	12/31/2013							
68387-0469-30		Q0178		03/01/2007	12/31/2013	HYDROXYZINE PAMOATE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	30	EA	BO	PO	EA	50	MG	2	03/01/2007	12/31/2013							
68387-0536-12		Q0170		03/08/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	12	EA	BO	PO	EA	25	MG	1	03/08/2006	12/31/2013							
68387-0536-30		Q0170		05/01/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	30	EA	BO	PO	EA	25	MG	1	05/01/2006	12/31/2013							
68387-0536-60		Q0170		05/04/2007	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60	EA	BO	PO	EA	25	MG	1	05/04/2007	12/31/2013							
68387-0536-90		Q0170		05/01/2006	12/31/2013	PROMETHAZINE HYDROCHLORIDE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	90	EA	BO	PO	EA	25	MG	1	05/01/2006	12/31/2013							
68387-0541-30		Q0163		05/01/2006	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HYDROCHLORIDE 25 MG	30	EA	BO	PO	EA	50	MG	0.5	05/01/2006	06/01/2014							
68387-0565-06		Q0144		05/01/2006	06/01/2014	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6	EA	BX	PO	EA	1	GM	0.25	05/01/2006	06/01/2014							
68546-0317-30		J1595		04/28/2008	99/99/9999	INJECTION, GLATIRAMER ACETATE, 20 MG	COPAXONE 20 MG/ML	1	ML	DP	MR	EA	20	MG	30	04/28/2008	99/99/9999							
68817-0134-50		J9264		01/01/2006	99/99/9999	INJECTION, PACLITAXEL PROTEIN-BOUND PARTICLES, 1 MG	ABRAXANE 100 MG	1	EA	VL	IV	EA	1	MG	100	01/01/2006	99/99/9999							
68883-0010-03		J1642		01/05/2006	08/17/2012	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (IN 6ML SD SYRINGE,PF) 10 U/ML	3	ML	SR	IV	ML	10	U	1	01/05/2006	08/17/2012							
68883-0010-05		J1642		01/05/2006	08/17/2012	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (IN 12ML SD SYRINGE,PF) 10 U/ML	5	ML	SR	IV	ML	10	U	1	01/05/2006	08/17/2012							
68883-0010-06		J1642		01/05/2006	08/17/2012	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (IN 3ML SD SYRINGE,PF) 10 U/ML	2.5	ML	SR	IV	ML	10	U	1	01/05/2006	08/17/2012							
68883-0100-03		J1642		01/05/2006	08/17/2012	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (IN 6ML SD SYRINGE,PF) 100 U/ML	3	ML	SR	IV	ML	10	U	10	01/05/2006	08/17/2012							
68883-0100-04		J1642		01/05/2006	08/17/2012	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (IN 12ML SD SYRINGE,PF) 100 U/ML	3	ML	SR	IV	ML	10	U	10	01/05/2006	08/17/2012							
68883-0100-05		J1642		01/05/2006	08/17/2012	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (IN 12ML SD SYRINGE,PF) 100 U/ML	5	ML	SR	IV	ML	10	U	10	01/05/2006	08/17/2012							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
68883-0100-06	J1642			01/05/2006	08/17/2012	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (IN 6ML SD SYRINGE,PF) 100 U/ML	5 ML	SR	IV	ML		10 U		10	01/05/2006	08/17/2012						
68883-0900-01	A4216			01/01/2007	08/17/2012	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE FLUSH (IN 3ML SD SYRINGE,PF) 0.9%	2.5 ML	SR	IV	ML		10 ML		0.1	01/01/2007	08/17/2012						
68883-0900-03	A4216			01/01/2007	08/17/2012	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE FLUSH (IN 6ML SD SYRINGE,PF) 0.9%	3 ML	SR	IV	ML		10 ML		0.1	01/01/2007	08/17/2012						
68883-0900-04	A4216			01/01/2007	08/17/2012	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE FLUSH (IN 6ML SD SYRINGE,PF) 0.9%	5 ML	SR	IV	ML		10 ML		0.1	01/01/2007	08/17/2012						
75137-0212-15	Q0163			01/01/2002	02/16/2016	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPOZ NIGHTTIME SLEEP AID (GELCAPLET) 50 MG	16 EA	BO	PO	EA		50 MG		1	01/01/2002	02/16/2016						
68387-0240-10	J7506			05/29/2008	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	10 EA	DP	PO	EA		4 MG		4	05/29/2008	06/01/2014						
60505-6145-04	J0692			03/15/2018	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME NOVAPLUS (USP) 2 GM	10 EA	VL	IJ	EA		500 MG		4	03/15/2018	99/99/9999						
52959-0127-18	J7506			06/18/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	18 EA	BO	PO	EA		5 MG		4	06/18/2008	12/31/2015						
21695-0765-48	J7506			06/09/2008	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	48 EA	NA	PO	EA		5 MG		2	06/09/2008	06/01/2014						
63323-0318-01	J1626			06/25/2008	99/99/9999	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X1ML,SDV,PF) 1 MG/ML	1 ML	VL	IV	ML		100 MCG		10	06/25/2008	99/99/9999						
00703-7973-01	J1626			06/30/2008	04/30/2012	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X4ML) 1 MG/ML	4 ML	VL	IV	ML		100 MCG		10	06/30/2008	04/30/2012						
63323-0319-04	J1626			06/25/2008	99/99/9999	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X4ML,MDV) 1 MG/ML	4 ML	VL	IV	ML		100 MCG		10	06/25/2008	99/99/9999						
66758-0035-01	J1626			06/30/2008	99/99/9999	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X1ML,SINGLE-USE) 1 MG/ML	1 ML	VL	IV	ML		100 MCG		10	06/30/2008	99/99/9999						
66758-0036-01	J1626			06/30/2008	99/99/9999	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X4ML,MULTI-USE) 1 MG/ML	4 ML	VL	IV	ML		100 MCG		10	06/30/2008	99/99/9999						
50090-2345-09	None			06/08/2018	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	36 EA	BO	PO	EA		2.5 MG		1	06/08/2018	99/99/9999						
51407-0121-01	None			06/07/2018	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	100 EA	BO	PO	EA		2.5 MG		1	06/07/2018	99/99/9999						
68382-0751-67	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 5 MG	14 EA	BO	PO	EA		5 MG		1	06/01/2018	99/99/9999						
68382-0751-96	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 5 MG	5 EA	BO	PO	EA		5 MG		1	06/01/2018	99/99/9999						
42023-0118-01	J3250			08/01/2008	99/99/9999	INJECTION, TRIMETHOENZAMIDE HCL, UP TO 200 MG	TRIMETHOENZAMIDE HCL (MDV,1X20ML) 100 MG/ML	20 ML	VL	IM	ML		200 MG		0.5	08/01/2008	99/99/9999						
00054-3177-57	J8540			07/31/2008	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (1X240ML)	240 ML	BO	PO	ML		0.25 MG		2	07/31/2008	99/99/9999						
21695-0573-30	Q0177			08/14/2008	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	30 EA	BO	PO	EA		25 MG		1	08/14/2008	06/01/2014						
21695-0573-20	Q0177			08/14/2008	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 25 MG	20 EA	BO	PO	EA		25 MG		1	08/14/2008	06/01/2014						
21695-0307-21	J7506			08/14/2008	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	21 EA	BO	PO	EA		5 MG		4	08/14/2008	06/01/2014						
21695-0307-15	J7506			09/03/2008	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 20 MG	15 EA	BO	PO	EA		5 MG		4	09/03/2008	06/01/2014						
68382-0752-67	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 20 MG	14 EA	BO	PO	EA		20 MG		1	06/01/2018	99/99/9999						
68382-0752-96	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 20 MG	5 EA	BO	PO	EA		20 MG		1	06/01/2018	99/99/9999						
68382-0753-67	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14 EA	BO	PO	EA		100 MG		1	06/01/2018	99/99/9999						
68387-0241-15	J7506			07/23/2008	06/01/2014	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	15 EA	BO	PO	EA		5 MG		2	07/23/2008	06/01/2014						
55289-0330-07	J7506			09/16/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 50 MG	7 EA	BO	PO	EA		5 MG		10	09/16/2008	12/31/2015						
21695-0571-30	Q0164			08/22/2008	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 5 MG	30 EA	BO	PO	EA		5 MG		1	08/22/2008	06/01/2014						
35356-0359-30	J8540			08/08/2008	01/01/2015	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 1 MG	1 EA	BO	PO	EA		0.3 MG		4	08/08/2008	01/01/2015						
49999-0028-21	J7506			08/08/2008	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 10 MG	21 EA	BO	PO	EA		5 MG		2	08/08/2008	12/31/2015						
35356-0325-00	Q0165			08/01/2008	12/31/2013	PROCHLORPERAZINE MALEATE, 10 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	1 EA	BO	PO	EA		10 MG		1	08/01/2008	12/31/2013						
42023-0119-25	J3250			07/22/2008	99/99/9999	INJECTION, TRIMETHOENZAMIDE HCL, UP TO 200 MG	TIGAN (SDV, 25X2ML) 100 MG/ML	2 ML	VL	IM	ML		200 MG		1	07/22/2008	99/99/9999						
54868-5511-00	J3535			10/21/2008	99/99/9999	DRUG ADMINISTERED THROUGH A METERED DOSE INHALER	IPRATROPIUM BROMIDE (0.017 MG/ACTUATION)	12.9 GM	PC	IH	GM		1 MG		0.017	10/21/2008	99/99/9999						
64679-0661-02	J1626			07/01/2008	04/30/2014	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X4ML) 1 MG/ML MG/ML	4 MG	VL	IV	ML		100 MCG		10	07/01/2008	04/30/2014						
64679-0661-03	J1626			07/01/2008	04/30/2014	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X1ML) 1 MG/ML	1 MG	VL	IV	ML		100 MCG		10	07/01/2008	04/30/2014						
49502-0605-61	KO	J7606	KO	01/01/2009	99/99/9999	MICROGRAMS	PERFORMIST 20 MCG/2 ML	2 ML	PC	IH	ML		20 MCG		0.5	01/01/2009	99/99/9999						
54868-4319-00	J1750			01/01/2009	99/99/9999	INJECTION, IRON DEXTRAN, 50 MG	INFED (2MLX10) 50 MG/ML	2 ML	VL	IJ	ML		50 MG		1	01/01/2009	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
44206-0436-05	J1459			01/01/2009	99/99/9999	INJECTION, IMMUNE GLOBULIN (PRIVIGEN), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	PRIVIGEN (PF,LATEX-FREE) 10%	1	ML	VL	IV	ML	500	MG	0.2	01/01/2009	99/99/9999						
44206-0437-10	J1459			01/01/2009	99/99/9999	INJECTION, IMMUNE GLOBULIN (PRIVIGEN), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	PRIVIGEN (PF,LATEX-FREE) 10%	1	ML	VL	IV	ML	500	MG	0.2	01/01/2009	99/99/9999						
44206-0438-20	J1459			01/01/2009	99/99/9999	INJECTION, IMMUNE GLOBULIN (PRIVIGEN), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	PRIVIGEN (PF,LATEX-FREE) 10% TKASE (VIAL W/DILUENT,SRN,PADS) 50 MG	1	ML	VL	IV	ML	500	MG	0.2	01/01/2009	99/99/9999						
50242-0038-61	J3101			01/01/2009	06/30/2012	INJECTION, TENECTEPLASE, 1 MG	ETOPOSIDE (U.S.P.) 1 GM	1	EA	BX	IV	EA	1	MG	50	01/01/2009	06/30/2012						
51927-2772-00	J9181			01/01/2009	99/99/9999	INJECTION, ETOPOSID, 10 MG	ETOPOSIDE (U.S.P.) 1 GM	1	EA	BO	NA	GM	10	MG	100	01/01/2009	99/99/9999						
08881-5801-21	J1642			03/14/2002	05/01/2017	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	MONOJECT PREFILL HEPARIN LOCK FLUSH (SRN, 12 ML,LATEX-FREE) 10 U/ML (10 ML 180S)	10	ML	SR	IV	U	10	U		1	03/14/2002	05/01/2017					
08881-5801-23	J1642			03/14/2002	05/01/2017	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	MONOJECT PREFILL HEPARIN LOCK FLUSH (SRN,12 ML, PF, LATEX-FREE) 10 U/ML (2.5 ML 180S)	2.5	ML	SR	IV	U	10	U		1	03/14/2002	05/01/2017					
08881-5801-25	J1642			08/23/2006	05/01/2017	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	MONOJECT PREFILL HEPARIN LOCK FLUSH (SRN,12 ML,PF,LATEX-FREE) 10 U/ML (5 ML 180S)	10	ML	SR	IV	U	10	U		1	08/23/2006	05/01/2017					
08881-5901-21	J1642			03/14/2002	05/01/2017	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	MONOJECT PREFILL HEPARIN LOCK FLUSH (SRN,12 ML,PF,LATEX-FREE) 100 U/ML (10ML 180S)	10	ML	SR	IV	U	10	U		10	03/14/2002	05/01/2017					
08881-5901-23	J1642			03/14/2002	01/01/2017	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	MONOJECT PREFILL ADVANCED HEPARIN LOCK FLUSH (SRN,12 ML,PF,LATEX-FREE) 100 U/ML (2.5 ML 180S)	2.5	ML	SR	IV	U	10	U		10	03/14/2002	01/01/2017					
08881-5901-25	J1642			08/23/2006	05/01/2017	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	MONOJECT PREFILL HEPARIN LOCK FLUSH (SRN,12 ML,PF,LATEX-FREE) 100 U/ML (5 ML 180S)	5	ML	SR	IV	U	10	U		10	08/23/2006	05/01/2017					
10019-0176-39	J2270			08/21/1998	10/31/2013	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (1X1ML,SDV,USP) 5MG/ML	1	ML	VL	IJ	ML	10	MG	0.5	08/21/1998	10/31/2013						
10019-0177-39	J2270			09/13/2001	10/17/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (1X1ML,USP) 8MG/ML	1	ML	VL	IJ	ML	10	MG	0.8	09/13/2001	10/17/2016						
10019-0178-37	J2270			08/21/1998	02/03/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (1X1ML,USP) 10MG/ML	1	ML	VL	IJ	ML	10	MG	1	08/21/1998	02/03/2016						
10019-0179-39	J2270			05/05/1999	02/03/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (1X1ML,SDV, USP) 15MG/ML	1	ML	VL	IJ	ML	10	MG	1.5	05/05/1999	02/03/2016						
00088-2502-05	J1817			03/04/2009	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	APIDRA SOLOSTAR (5X3ML) 100U/ML	3	ML	EA	IJ	ML	50	U	2	03/04/2009	99/99/9999						
68382-0753-96	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5	EA	BO	PO	EA	100	MG	1	06/01/2018	99/99/9999						
61553-0651-76	J2271			03/03/2005	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	MORPHINE SULFATE (5X55ML) 50 MG/ML	55	ML	EA	IJ	ML	100	MG	0.5	03/03/2005	12/31/2014						
00007-4205-11	None			07/01/2009	07/30/2017	TOPOTECAN, ORAL, 0.25 MG	HYCAMTIN 0.25 MG	10	EA	BO	PO	EA	0.25	MG	1	07/01/2009	07/30/2017						
00007-4207-11	None			07/01/2009	03/20/2017	TOPOTECAN, ORAL, 0.25 MG	HYCAMTIN 1 MG	10	EA	BO	PO	EA	0.25	MG	4	07/01/2009	03/20/2017						
51991-0940-17	J3370			07/06/2017	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (USP,PF,LATEX-FREE) 500 MG	10	EA	VL	IV	EA	500	MG	1	07/06/2017	99/99/9999						
51991-0941-17	J3370			07/06/2017	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (USP,PF,LATEX-FREE) 1 GM	10	EA	VL	IV	EA	500	MG	2	07/06/2017	99/99/9999						
68382-0754-67	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 140 MG MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	14	EA	BO	PO	EA	20	MG	7	06/01/2018	99/99/9999						
00093-7477-05	J7517			05/05/2009	06/04/2018	MYCOPHENOLATE MOFETIL , ORAL, 250 MG	MORPHINE SULFATE (5X50ML,LATEX-FREE) 50 MG/ML	500	EA	BO	PO	EA	250	MG	2	05/05/2009	06/04/2018						
61553-0649-75	J2271			03/03/2005	12/31/2014	INJECTION, MORPHINE SULFATE, 100MG	TACROLIMUS (HARD GELATIN) 0.5 MG	50	ML	EA	IJ	ML	100	MG	0.5	03/03/2005	12/31/2014						
00781-2102-01	J7507			08/10/2009	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	BO	PO	EA	1	MG	0.5	08/10/2009	99/99/9999						
00781-2103-01	J7507			08/10/2009	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	BO	PO	EA	1	MG	1	08/10/2009	99/99/9999						
00781-2104-01	J7507			08/10/2009	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	100	EA	BO	PO	EA	1	MG	5	08/10/2009	99/99/9999						
68382-0754-96	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 140 MG MYCOPHENOLATE MOFETIL (FILM-COATED), 500MG	5	EA	BO	PO	EA	20	MG	7	06/01/2018	99/99/9999						
68382-0131-01	J7517			05/04/2009	08/31/2013	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED), 500MG	100	EA	BO	PO	EA	250	MG	2	05/04/2009	08/31/2013						
68382-0131-05	J7517			05/04/2009	09/30/2013	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED), 500MG	500	EA	BO	PO	EA	250	MG	2	05/04/2009	09/30/2013						
24385-0431-26	Q0163			08/03/2009	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	NIGHTTIME SLEEP AID (CAPLET) 25 MG HEPARIN LOCK FLUSH (FTV,25X10ML) 10 U/ML	24	EA	NA	PO	EA	50	MG	0.5	08/03/2009	99/99/9999						
00409-1151-70	J1642			10/01/2009	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (VIAL,FLUPTOP,LIFESHIELD) 100 U/ML	10	ML	VL	IV	ML	10	U	1	10/01/2009	02/03/2016						
00409-1152-12	J1642			10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (VIAL,FLUPTOP,LIFESHIELD) 100 U/ML	10	ML	VL	IV	ML	10	U	10	10/01/2009	99/99/9999						
00409-1152-78	J1642			10/01/2009	02/03/2016	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (VIAL,FLUPTOP,LATEX-FREE) 100 U/ML	30	ML	VL	IV	ML	10	U	10	10/01/2009	02/03/2016						
00409-1280-31	J1642			10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,LATEX-FREE) 10 U/ML	1	ML	SR	IV	ML	10	U	1	10/01/2009	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-1280-32		J1642		10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,LATEX-FREE) 10 U/ML	2	ML	SR	IV	ML	10	U	1	10/01/2009	99/99/9999						
00409-1280-33		J1642		10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,LATEX-FREE) 10 U/ML	3	ML	CR	IV	ML	10	U	1	10/01/2009	99/99/9999						
00409-1280-35		J1642		03/03/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,LATEX-FREE) 10 U/ML	5	ML	CR	IV	ML	10	U	1	03/03/2009	99/99/9999						
00409-1281-31		J1642		10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,50X1ML) 100 U/ML	1	ML	CR	IV	ML	10	U	10	10/01/2009	99/99/9999						
00409-1281-32		J1642		10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,CARPUJECT) 100 U/ML	2	ML	CR	IV	ML	10	U	10	10/01/2009	99/99/9999						
00409-1281-33		J1642		10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,25X3ML) 100 U/ML	3	ML	CR	IV	ML	10	U	10	10/01/2009	99/99/9999						
00409-1281-35		J1642		10/01/2009	99/99/9999	INJECTION, HEPARIN SODIUM, (HEPARIN LOCK FLUSH), PER 10 UNITS	HEPARIN LOCK FLUSH (LUER LOCK,CARPUJECT) 100 U/ML	5	ML	CR	IV	ML	10	U	10	10/01/2009	99/99/9999						
00378-1005-01		J7500		12/22/2009	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE, 50 MG	100	EA	BO	PO	EA	50	MG	1	12/22/2009	99/99/9999						
50580-0843-10		Q0163		02/02/2009	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SIMPLY SLEEP (CAPLET) 25 MG	100	EA	BO	PO	EA	50	MG	0.5	02/02/2009	99/99/9999						
50580-0843-24		Q0163		02/02/2009	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	SIMPLY SLEEP (CAPLET) 25 MG	24	EA	BO	PO	EA	50	MG	0.5	02/02/2009	99/99/9999						
00378-6989-62		J7644		10/07/2009	04/02/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	25	EA	SOL	IH	ML	1	MG	0.2	10/07/2009	04/02/2013						
00378-6989-62	KO	J7644	KO	10/07/2009	04/02/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	25	EA	SOL	IH	ML	1	MG	0.2	10/07/2009	04/02/2013						
00378-6989-66		J7644		10/07/2009	03/03/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML,PF) 0.02%	60	EA	SOL	IH	ML	1	MG	0.2	10/07/2009	03/03/2013						
00378-6989-66	KO	J7644	KO	10/07/2009	03/03/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML,PF) 0.02%	60	EA	SOL	IH	ML	1	MG	0.2	10/07/2009	03/03/2013						
00378-6989-93		J7644		10/07/2009	10/07/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	30	EA	SOL	IH	ML	1	MG	0.2	10/07/2009	10/07/2013						
00378-6989-93	KO	J7644	KO	10/07/2009	10/07/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	30	EA	SOL	IH	ML	1	MG	0.2	10/07/2009	10/07/2013						
00378-6990-52		J7613		10/07/2009	12/12/2012	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (5X5) 0.083%	25	EA	SOL	IH	ML	1	MG	0.83333	10/07/2009	12/12/2012						
00378-6990-52	KO	J7613	KO	10/07/2009	12/12/2012	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (5X5) 0.083%	25	EA	SOL	IH	ML	1	MG	0.83333	10/07/2009	12/12/2012						
00378-6990-58		J7613		10/07/2009	01/21/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (6X5) 0.083%	30	EA	SOL	IH	ML	1	MG	0.83333	10/07/2009	01/21/2013						
00378-6990-58	KO	J7613	KO	10/07/2009	01/21/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (6X5) 0.083%	30	EA	SOL	IH	ML	1	MG	0.83333	10/07/2009	01/21/2013						
00378-6990-91		J7613		10/07/2009	04/10/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (12X5) 0.083%	60	EA	SOL	IH	ML	1	MG	0.83333	10/07/2009	04/10/2013						
00378-6990-91	KO	J7613	KO	10/07/2009	04/10/2013	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (12X5) 0.083%	60	EA	SOL	IH	ML	1	MG	0.83333	10/07/2009	04/10/2013						
00378-6993-93		J7612		08/28/2009	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (USP,PF) 1.25 MG/0.5 ML	30	EA	SOL	IH	ML	0.5	MG	5	08/28/2009	99/99/9999						
00378-6993-93	KO	J7612	KO	08/28/2009	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (USP,PF) 1.25 MG/0.5 ML	30	EA	SOL	IH	ML	0.5	MG	5	08/28/2009	99/99/9999						
66336-0045-60		Q0163		04/01/2010	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	60	EA	BO	PO	EA	50	MG	1	04/01/2010	06/01/2014						
66215-0401-01		J1325		08/27/2007	99/99/9999	INJECTION, EPOPROSTENOL, 0.5 MG	EPOPROSTENOL (SINGLE DOSE,LYOPHILIZED) 1.5 MG	1	EA	EA	IV	EA	0.5	MG	3	08/27/2007	99/99/9999						
66336-0045-20		Q0163		04/01/2010	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	15	EA	BO	PO	EA	50	MG	1	04/01/2010	06/01/2014						
66336-0045-90		Q0163		04/01/2010	06/01/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 50 MG	90	EA	BO	PO	EA	50	MG	1	04/01/2010	06/01/2014						
66336-0515-10		J7506		04/01/2010	06/01/2014	PREDNISON, ORAL, PER SMG	PREDNISON 5 MG	10	EA	TAB	PO	EA	5	MG	1	04/01/2010	06/01/2014						
66336-0629-10		Q0173		04/01/2010	06/01/2014	TRIMETHOENZAMIDE HYDROCHLORIDE, 250 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	TRIMETHOENZAMIDE HCL 250 MG	10	EA	NA	PO	EA	250	MG	1	04/01/2010	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
63628-1472-02		None		02/01/2009	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	12	EA	BO	PO	EA	2.5	MG	1	02/01/2009	99/99/9999							
00008-1040-05		J7520		04/09/2010	99/99/9999	SIROLIMUS, ORAL, 1 MG	RAPAMUNE 0.5 MG	100	EA	EA	PO	EA	1	MG	0.5	04/09/2010	99/99/9999							
00378-6991-52		J7613		11/02/2009	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 0.63 MG/3 ML	3	ML	EA	IH	ML	1	MG	0.21	11/02/2009	99/99/9999							
00378-6991-52	KO	J7613	KO	11/02/2009	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 0.63 MG/3 ML	3	ML	EA	IH	ML	1	MG	0.21	11/02/2009	99/99/9999							
00378-6992-52		J7613		11/02/2009	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 1.25 MG/3 ML	3	ML	EA	IH	ML	1	MG	0.4166	11/02/2009	99/99/9999							
00378-6992-52	KO	J7613	KO	11/02/2009	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML,PF) 1.25 MG/3 ML	3	ML	EA	IH	ML	1	MG	0.4166	11/02/2009	99/99/9999							
66336-0479-15		J8540		04/01/2010	06/01/2014	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE, 4 MG	15	EA	TAB	PO	EA	0.25	MG	16	04/01/2010	06/01/2014							
55111-0525-01		J7507		05/14/2010	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 0.5 MG	100	EA	CAP	PO	EA	1	MG	0.5	05/14/2010	99/99/9999							
55111-0526-01		J7507		05/14/2010	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	CAP	PO	EA	1	MG	1	05/14/2010	99/99/9999							
55111-0527-01		J7507		05/14/2010	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	100	EA	CAP	PO	EA	1	MG	5	05/14/2010	99/99/9999							
68382-0130-05		J7517		05/04/2009	08/31/2013	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	500	EA	CAP	PO	EA	250	MG	1	05/04/2009	08/31/2013							
00378-6989-64		J7644		10/07/2009	02/18/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	30	EA	PC	IH	ML	1	MG	0.2	10/07/2009	02/18/2013							
00378-6989-64	KO	J7644	KO	10/07/2009	02/18/2013	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	30	EA	PC	IH	ML	1	MG	0.2	10/07/2009	02/18/2013							
00008-1040-10		J7520		04/09/2010	99/99/9999	SIROLIMUS, ORAL, 1 MG	RAPAMUNE 0.5 MG	100	EA	BX	PO	EA	1	MG	0.5	04/09/2010	99/99/9999							
00173-0517-00		J1325		07/27/2010	99/99/9999	INJECTION, EPOPROSTENOL, 0.5 MG	FLOLAN 0.5 MG	1	EA	VL	IV	EA	0.5	MG	1	07/27/2010	99/99/9999							
00173-0519-00		J1325		07/27/2010	99/99/9999	INJECTION, EPOPROSTENOL, 0.5 MG	FLOLAN 1.5 MG	1	EA	VL	IV	EA	0.5	MG	3	07/27/2010	99/99/9999							
54569-4026-04		J7506		08/24/2010	12/31/2015	PREDNISONE, ORAL, PER SMG	PREDNISONE 5 MG	40	EA	TAB	PO	EA	5	MG	1	08/24/2010	12/31/2015							
00378-2045-01		J7507		09/23/2010	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 0.5 MG	100	EA	EA	PO	EA	1	MG	0.5	09/23/2010	99/99/9999							
00378-2046-01		J7507		09/23/2010	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	EA	PO	EA	1	MG	1	09/23/2010	99/99/9999							
00378-2047-01		J7507		09/23/2010	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	100	EA	EA	PO	EA	1	MG	5	09/23/2010	99/99/9999							
44206-0451-01		J1559		01/01/2011	99/99/9999	INJECTION, IMMUNE GLOBULIN (HIZENTRA), 100 MG	HIZENTRA (SINGLE-USE VIAL,PF) 20%	5	ML	VL	SC	ML	100	MG	2	01/01/2011	99/99/9999							
44206-0452-02		J1559		01/01/2011	99/99/9999	INJECTION, IMMUNE GLOBULIN (HIZENTRA), 100 MG	HIZENTRA (SINGLE-USE VIAL,PF) 20%	10	ML	VL	SC	ML	100	MG	2	01/01/2011	99/99/9999							
44206-0454-04		J1559		01/01/2011	99/99/9999	INJECTION, IMMUNE GLOBULIN (HIZENTRA), 100 MG	HIZENTRA (SINGLE-USE VIAL,PF) 20%	20	ML	VL	SC	ML	100	MG	2	01/01/2011	99/99/9999							
00591-3797-83		J7613		11/04/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (2.5 MG/3ML) 0.083% (25X3ML)	25	EA	SOL	IH	ML	1	MG	0.83	11/04/2010	99/99/9999							
00591-3797-60		J7613		11/04/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (2.5 MG/3ML) 0.083% (60X3ML)	60	EA	SOL	IH	ML	1	MG	0.83	11/04/2010	99/99/9999							
00591-3797-83	KO	J7613	KO	11/04/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (2.5 MG/3ML) 0.083% (25X3ML)	25	EA	SOL	IH	ML	1	MG	0.83	11/04/2010	99/99/9999							
00591-3797-60	KO	J7613	KO	11/04/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (2.5 MG/3ML) 0.083% (60X3ML)	60	EA	SOL	IH	ML	1	MG	0.83	11/04/2010	99/99/9999							
00093-6815-73		J7626		12/15/2009	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.25 MG/2 ML	30	EA	PC	IH	ML	0.25	MG	0.5	12/15/2009	99/99/9999							
00093-6815-73	KO	J7626	KO	12/15/2009	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.25 MG/2 ML	30	EA	PC	IH	ML	0.25	MG	0.5	12/15/2009	99/99/9999							
00093-6816-73		J7626		12/15/2009	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.5 MG/2 ML	30	EA	PC	IH	ML	0.5	MG	0.5	12/15/2009	99/99/9999							
00093-6816-73	KO	J7626	KO	12/15/2009	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.5 MG/2 ML	30	EA	PC	IH	ML	0.5	MG	0.5	12/15/2009	99/99/9999							
00093-7477-01		J7517		05/05/2009	06/04/2018	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	100	EA	BO	PO	EA	250	MG	2	05/05/2009	06/04/2018							
00487-0301-01		J7613		07/19/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML,LDPE VIAL,PF) 0.63 MG/3 ML	30	EA	PC	IH	ML	1	MG	0.21	07/19/2010	99/99/9999							
00487-0301-01	KO	J7613	KO	07/19/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML,LDPE VIAL,PF) 0.63 MG/3 ML	30	EA	PC	IH	ML	1	MG	0.21	07/19/2010	99/99/9999							
00409-1283-10		J1170		05/15/2009	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (USP,ISECURE SINGLE-DOSE) 1 MG/ML	10	EA	SR	IJ	ML	4	MG	0.25	05/15/2009	99/99/9999							
00409-1312-10		J1170		10/01/2010	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (USP,ISECURE SINGLE-DOSE) 2 MG/ML	10	EA	SR	IJ	ML	4	MG	0.5	10/01/2010	99/99/9999							
00093-7334-05		J7517		05/06/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	500	EA	BO	PO	EA	250	MG	1	05/06/2009	99/99/9999							
12496-0757-01		J0592		01/01/2003	01/18/2015	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENEX (AMP) 0.3 MG/ML	1	ML	AM	IJ	ML	0.1	MG	3.24	01/01/2003	01/18/2015							
18860-0720-10		J2278		01/31/2011	99/99/9999	INJECTION, ZICONOTIDE, 1 MICROGRAM	PRIALT (1X1ML,SINGLE-USE VIAL) 100 MCG/ML	1	ML	VL	IN	ML	1	MCG	100	01/31/2011	99/99/9999							
18860-0722-10		J2278		01/31/2011	99/99/9999	INJECTION, ZICONOTIDE, 1 MICROGRAM	PRIALT (1X5ML,SINGLE-USE VIAL) 100 MCG/ML	1	ML	VL	IN	ML	1	MCG	100	01/31/2011	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
18860-0723-10	J2278			01/31/2011	99/99/9999	INJECTION, ZICONOTIDE, 1 MICROGRAM	PRIALT (1X20ML,SINGLE-USE VIAL) 25 MCG/ML	1 ML	VL	IN	ML		1 MCG		25	01/31/2011	99/99/9999						
54868-3826-08	None			06/29/2010	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	40 EA	BO	PO	EA		2.5 MG		1	06/29/2010	99/99/9999						
54868-3826-09	None			09/13/2010	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	2 EA	BO	PO	EA		2.5 MG		1	09/13/2010	99/99/9999						
13533-0800-12	J1561			12/07/2010	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX-C (1X100ML,SINGLE-USE) 100 MG/1 ML	10 ML	VL	IJ	ML		500 MG		0.2	12/07/2010	99/99/9999						
13533-0800-15	J1561			12/07/2010	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX-C (1X25ML,SINGLE-USE) 100 MG/1 ML	25 ML	VL	IJ	ML		500 MG		0.2	12/07/2010	99/99/9999						
13533-0800-20	J1561			12/07/2010	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX-C (1X50ML,SINGLE-USE) 100 MG/1 ML	50 ML	VL	IJ	ML		500 MG		0.2	12/07/2010	99/99/9999						
13533-0800-24	J1561			12/07/2010	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX-C (1X200ML,SINGLE-USE) 100 MG/1 ML	200 ML	VL	IJ	ML		500 MG		0.2	12/07/2010	99/99/9999						
13533-0800-71	J1561			12/07/2010	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMUNEX-C (1X100ML,SINGLE-USE) 100 MG/1 ML	100 ML	VL	IJ	ML		500 MG		0.2	12/07/2010	99/99/9999						
00944-2700-07	J1569			03/18/2011	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAGARD LIQUID), NON-LYOPHILIZED,(E.G. LIQUID), 500 MG	GAMMAGARD LIQUID (1X300ML, PF, LATEX-FREE) 100 MG/ML	1 ML	VL	IV	ML		500 MG		0.2	03/18/2011	99/99/9999						
68382-0755-67	None			06/01/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 180 MG	14 EA	BO	PO	EA		20 MG		9	06/01/2018	99/99/9999						
54868-3826-00	None			02/07/2011	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	16 EA	DP	PO	EA		2.5 MG		1	02/07/2011	99/99/9999						
00004-1101-75	None			03/29/2011	12/31/2013	CAPECITABINE, 500 MG, ORAL	XELODA (10 X 12.FILM COATED) 500MG	120 EA	BP	PO	EA		500 MG		1	03/29/2011	12/31/2013						
00591-3797-30	J7613			11/04/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	11/04/2010	99/99/9999						
00591-3797-30	KO	J7613	KO	11/04/2010	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	11/04/2010	99/99/9999						
52609-0001-05	None			05/20/2011	99/99/9999	MELPHALAN, ORAL, 2 MG	ALKERAN (FILM-COATED) 2 MG	50 EA	BO	PO	EA		2 MG		1	05/20/2011	99/99/9999						
00143-9708-01	J2260			03/29/2011	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE, 1 MG/ML	1 ML	VL	IV	ML		5 MG		0.2	03/29/2011	99/99/9999						
00143-9709-10	J2260			03/29/2011	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE, 1 MG/ML	10 ML	VL	IV	ML		5 MG		0.2	03/29/2011	99/99/9999						
00069-3030-20	J9000			05/19/2011	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (PF) 2 MG/ML	1 ML	VL	IV	ML		10 MG		0.2	05/19/2011	99/99/9999						
00069-3031-20	J9000			05/19/2011	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (PF) 2 MG/ML	1 ML	VL	IV	ML		10 MG		0.2	05/19/2011	99/99/9999						
00069-3032-20	J9000			05/19/2011	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (PF) 2 MG/ML	1 ML	VL	IV	ML		10 MG		0.2	05/19/2011	99/99/9999						
00069-3033-20	J9000			05/19/2011	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (PF) 2 MG/ML	1 ML	VL	IV	ML		10 MG		0.2	05/19/2011	99/99/9999						
00069-3034-20	J9000			05/19/2011	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (PF) 2 MG/ML	1 ML	VL	IV	ML		10 MG		0.2	05/19/2011	99/99/9999						
33261-0759-20	None			06/01/2010	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE, 2.5 MG	20 EA	BO	PO	EA		2.5 MG		1	06/01/2010	99/99/9999						
33261-0759-30	None			06/01/2010	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE, 2.5 MG	30 EA	BO	PO	EA		2.5 MG		1	06/01/2010	99/99/9999						
33261-0759-40	None			06/01/2010	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE, 2.5 MG	40 EA	BO	PO	EA		2.5 MG		1	06/01/2010	99/99/9999						
33261-0759-60	None			06/01/2010	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE, 2.5 MG	60 EA	BO	PO	EA		2.5 MG		1	06/01/2010	99/99/9999						
00591-2223-15	J7502			12/23/2008	08/02/2016	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (USP, MODIFIED) 100 MG	30 EA	BX	PO	EA		100 MG		1	12/23/2008	08/02/2016						
00904-6012-60	None			10/12/2009	12/04/2012	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE (USP) 2.5 MG	100 EA	BO	PO	EA		2.5 MG		1	10/12/2009	12/04/2012						
67253-0320-36	None			06/25/2009	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	36 EA	BO	PO	EA		2.5 MG		1	06/25/2009	99/99/9999						
54868-5980-00	None			01/26/2009	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR 180 MG	14 EA	BO	PO	EA		20 MG		9	01/26/2009	99/99/9999						
49999-0380-36	None			12/23/2009	01/01/2015	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	36 EA	BO	PO	EA		2.5 MG		1	12/23/2009	01/01/2015						
21695-0111-00	None			02/02/2009	06/01/2014	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	100 EA	BO	PO	EA		2.5 MG		1	02/02/2009	06/01/2014						
00378-2250-01	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250MG	100 EA	BO	PO	EA		250 MG		1	05/04/2009	99/99/9999						
00378-4472-01	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	100 EA	BO	PO	EA		250 MG		2	05/04/2009	99/99/9999						
00378-4472-05	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	500 EA	BO	PO	EA		250 MG		2	05/04/2009	99/99/9999						
53270-0051-01	J1573			08/01/2010	12/31/2016	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAVENOUS, 0.5 ML	HEPAGAM B (1X5ML->312IU/ML,SDV)	1 ML	VL	IJ	ML		0.5 ML		2	08/01/2010	12/31/2016						
53270-0052-01	J1573			08/01/2010	12/31/2016	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAVENOUS, 0.5 ML	HEPAGAM B (1X1ML->312IU/ML,SDV)	1 ML	VL	IJ	ML		0.5 ML		2	08/01/2010	12/31/2016						
53270-0053-01	J1573			08/01/2010	12/31/2016	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAVENOUS, 0.5 ML	NOVAPLUS HEPAGAM B (1X1ML->312IU/ML,SDV)	1 ML	VL	IJ	ML		0.5 ML		2	08/01/2010	12/31/2016						
53270-0054-01	J1573			08/01/2010	12/31/2016	INJECTION, HEPATITIS B IMMUNE GLOBULIN (HEPAGAM B), INTRAVENOUS, 0.5 ML	NOVAPLUS HEPAGAM B (1X1ML->312IU/ML,SDV)	1 ML	VL	IJ	ML		0.5 ML		2	08/01/2010	12/31/2016						
53270-3000-01	J2792			06/01/2010	12/31/2016	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT 6 DETERGENT, 100 IU	WINRHO SDF (SDV) 15000 IU	1 ML	VL	IV	ML		100 IU		150	06/01/2010	12/31/2016						
53270-3100-01	J2792			06/01/2010	12/31/2016	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT 6 DETERGENT, 100 IU	WINRHO SDF (1X4.AML,SDV) 5000 IU	1 ML	VL	IV	ML		100 IU		50	06/01/2010	12/31/2016						
53270-3300-01	J2792			06/01/2010	12/31/2016	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT 6 DETERGENT, 100 IU	WINRHO SDF (1X1.3ML,SDV) 1500 IU	1 ML	VL	IV	ML		100 IU		15	06/01/2010	12/31/2016						
53270-3500-01	J2792			06/01/2010	12/31/2016	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT 6 DETERGENT, 100 IU	WINRHO SDF (1X2.2ML,SDV) 2500 IU	1 ML	VL	IV	ML		100 IU		25	06/01/2010	12/31/2016						
00591-3798-30	J7644			06/24/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	30 ML	PC	IH	ML		1 MG		0.2	06/24/2011	99/99/9999						
00591-3798-30	KO	J7644	KO	06/24/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	30 ML	PC	IH	ML		1 MG		0.2	06/24/2011	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00591-3798-60		J7644		05/23/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML.LDPE.PF) 0.02%	60	ML	PC	IH	ML	1	MG	0.2	05/23/2011	99/99/9999						
00591-3798-60	KO	J7644	KO	05/23/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML.LDPE.PF) 0.02%	60	ML	PC	IH	ML	1	MG	0.2	05/23/2011	99/99/9999						
00591-2222-15		J7515		12/23/2008	07/17/2016	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE (USP,MODIFIED) 25 MG	30	EA	BX	PO	EA	25	MG	1	12/23/2008	07/17/2016						
21695-0111-30		None		10/04/2011	06/01/2014	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	30	EA	BO	PO	EA	2.5	MG	1	10/04/2011	06/01/2014						
00093-7236-56		Q0162		01/01/2012	10/05/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	10/05/2016						
00143-2423-30		Q0162		01/01/2012	04/10/2012	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	04/10/2012						
00173-0446-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
00173-0447-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
00173-0447-02		Q0162		01/01/2012	08/21/2013	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	100	EA	BX	PO	EA	1	MG	8	01/01/2012	08/21/2013						
00173-0447-04		Q0162		01/01/2012	04/01/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN (1X3 DAILY PACK) 8 MG	3	EA	BX	PO	EA	1	MG	8	01/01/2012	04/01/2014						
00173-0569-00		Q0162		01/01/2012	08/29/2017	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 4 MG	30	EA	BX	PO	EA	1	MG	4	01/01/2012	08/29/2017						
00173-0570-00		Q0162		01/01/2012	09/18/2017	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	30	EA	BX	PO	EA	1	MG	8	01/01/2012	09/18/2017						
00378-0315-93		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
00378-0344-93		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
00378-7732-93		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
00378-7734-93		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
00378-7734-97		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP) 8 MG	10	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
00490-0075-00		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON ODT 4 MG	100	EA	BX	PO	EA	1	MG	4	01/01/2012	01/31/2014						
00490-0075-30		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON ODT 4 MG	30	EA	BX	PO	EA	1	MG	4	01/01/2012	01/31/2014						
00490-0075-60		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON ODT 4 MG	60	EA	BX	PO	EA	1	MG	4	01/01/2012	01/31/2014						
00490-0075-90		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON ODT 4 MG	90	EA	BX	PO	EA	1	MG	4	01/01/2012	01/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00781-1681-31		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
33358-0369-02		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	2	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
33358-0370-02		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	2	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
45802-0127-14		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	3	EA	BX	PO	EA	1	MG	4	01/01/2012	99/99/9999						
45802-0127-65		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
45802-0205-14		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	3	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
45802-0205-65		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
49999-0783-30		Q0162		01/01/2012	01/01/2015	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN (CAPLET) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	01/01/2015						
51079-0524-20		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (USP,10X10,FILM-COATED) 4 MG	100	EA	BX	PO	EA	1	MG	4	01/01/2012	99/99/9999						
51079-0525-20		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (USP,10X10,FILM-COATED) 8 MG	100	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
54569-5873-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	4	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
54868-3508-00		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN (1X3 DAILY PACK) 4 MG	3	EA	BX	PO	EA	1	MG	4	01/01/2012	02/03/2016						
54868-3508-01		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	02/03/2016						
54868-3508-02		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	10	EA	BO	PO	EA	1	MG	4	01/01/2012	02/03/2016						
54868-3509-00		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN (1X3 DAILY PACK) 8 MG	3	EA	BX	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-3509-01		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	15	EA	BO	PO	EA	1	MG	8	01/01/2012	02/03/2016						
00173-0489-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN (BERRY) 4 MG/5 ML	1	ML	BO	PO	ML	1	MG	0.8	01/01/2012	99/99/9999						
51079-0524-01		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	1	EA	BP	PO	EA	1	MG	4	01/01/2012	99/99/9999						
51079-0525-01		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	1	EA	BP	PO	EA	1	MG	8	01/01/2012	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51672-4091-03		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (1x50ML) 4MG/5ML	1	ML	BO	PO	ML	1	MG	0.8	01/01/2012	99/99/9999						
54868-3509-02		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	10	EA	BO	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-3509-03		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	20	EA	BO	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-5089-00		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	2	EA	BO	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-5089-01		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	15	EA	BO	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-5089-02		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	10	EA	BX	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-5089-03		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	3	EA	BO	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-5089-04		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	20	EA	BO	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-5089-05		Q0162		01/01/2012	02/03/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	30	EA	BX	PO	EA	1	MG	8	01/01/2012	02/03/2016						
54868-5738-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE 8 MG	10	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
54868-5749-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON ODT 8 MG	10	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
54868-5749-01		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON ODT 8 MG	15	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
54868-5801-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
54868-5801-01		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON 4 MG	15	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
54868-5887-00		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (STRAWBERRY) 4 MG	10	EA	BX	PO	EA	1	MG	4	01/01/2012	99/99/9999						
55045-3729-03		Q0162		01/01/2012	06/01/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	06/01/2014						
55045-3815-01		Q0162		01/01/2012	06/01/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON 8 MG	10	EA	BX	PO	EA	1	MG	8	01/01/2012	06/01/2014						
55111-0153-13		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (1X3.FILM-COATED) 4 MG	3	EA	BX	PO	EA	1	MG	4	01/01/2012	99/99/9999						
55111-0153-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55111-0154-13		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (1X3,FILM-COATED) 8 MG	3	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
55111-0154-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
55111-0156-11		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (1X1,FILM-COATED) 24 MG	1	EA	BP	PO	EA	1	MG	24	01/01/2012	99/99/9999						
55289-0559-03		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,STRAWBERRY) 4 MG	3	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
55289-0559-05		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (STRAWBERRY) 4 MG	5	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
55289-0559-06		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,STRAWBERRY) 4 MG	6	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
58016-0084-00		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	100	EA	BO	PO	EA	1	MG	8	01/01/2012	01/31/2014						
58016-0084-10		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	10	EA	BO	PO	EA	1	MG	8	01/01/2012	01/31/2014						
58016-0084-30		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	01/31/2014						
58016-0084-60		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	60	EA	BO	PO	EA	1	MG	8	01/01/2012	01/31/2014						
58016-0084-90		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 8 MG	90	EA	BO	PO	EA	1	MG	8	01/01/2012	01/31/2014						
58016-0826-00		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	100	EA	BO	PO	EA	1	MG	4	01/01/2012	01/31/2014						
58016-0826-30		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	01/31/2014						
58016-0826-60		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	60	EA	BO	PO	EA	1	MG	4	01/01/2012	01/31/2014						
58016-0826-90		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN 4 MG	90	EA	BO	PO	EA	1	MG	4	01/01/2012	01/31/2014						
60505-0381-05		Q0162		01/01/2012	01/31/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP, 1X50ML) 4 MG/5 ML	1	EA	BO	PO	ML	1	MG	0.8	01/01/2012	01/31/2014						
62756-0130-01		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
62756-0131-01		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
62756-0240-64		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON 4 MG	30	EA	BX	PO	EA	1	MG	4	01/01/2012	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
62756-0356-64		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON 8 MG	30	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
62756-0356-66		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON 8 MG	10	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
63304-0458-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
63304-0459-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
65862-0187-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
65862-0188-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
66336-0268-03		Q0162		01/01/2012	06/01/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	3	EA	BO	PO	EA	1	MG	8	01/01/2012	06/01/2014						
66336-0793-03		Q0162		01/01/2012	06/01/2014	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	3	EA	BO	PO	EA	1	MG	4	01/01/2012	06/01/2014						
68462-0105-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	01/01/2012	99/99/9999						
68462-0106-30		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/01/2012	99/99/9999						
68462-0157-13		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (STRAWBERRY) 4 MG	30	EA	BX	PO	EA	1	MG	4	01/01/2012	99/99/9999						
68462-0158-11		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (STRAWBERRY) 8 MG	30	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
68462-0158-13		Q0162		01/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (STRAWBERRY) 8 MG	10	EA	BX	PO	EA	1	MG	8	01/01/2012	99/99/9999						
00078-0414-20	J8561			01/01/2012	12/31/2012	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (6X10) 0.5 MG	60	EA	EA	PO	EA	0.25	MG	2	01/01/2012	12/31/2012						
00078-0414-61	J8561			01/01/2012	12/31/2012	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (1X1) 0.5 MG	1	EA	EA	PO	EA	0.25	MG	2	01/01/2012	12/31/2012						
00078-0415-20	J8561			01/01/2012	12/31/2012	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (6X10) 0.75 MG	60	EA	EA	PO	EA	0.25	MG	3	01/01/2012	12/31/2012						
00078-0415-61	J8561			01/01/2012	12/31/2012	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (1X1) 0.75 MG	1	EA	EA	PO	EA	0.25	MG	3	01/01/2012	12/31/2012						
00078-0417-20	J8561			01/01/2012	12/31/2012	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (6X10) 0.25 MG	60	EA	EA	PO	EA	0.25	MG	1	01/01/2012	12/31/2012						
00078-0417-61	J8561			01/01/2012	12/31/2012	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (1X1) 0.25 MG	1	EA	EA	PO	EA	0.25	MG	1	01/01/2012	12/31/2012						
64208-8234-01	J1557			01/01/2012	01/31/2015	INJECTION, IMMUNE GLOBULIN (GAMMAPLEX), INTRAVENOUS, NONLYOPHILIZED (E.G., LIQUID) 500 MG	GAMMAPLEX (1X50ML,SINGLE USE) 2.5 GM/50 ML	1	ML	VL	IV	ML	1	EA	0.1	01/01/2012	01/31/2015						
64208-8234-02	J1557			01/01/2012	99/99/9999	INJECTION, IMMUNE GLOBULIN (GAMMAPLEX), INTRAVENOUS, NONLYOPHILIZED (E.G., LIQUID) 500 MG	GAMMAPLEX (1X100ML,SINGLE USE) 5 GM/ 100 ML	1	ML	VL	IV	ML	1	EA	0.1	01/01/2012	99/99/9999						
64208-8234-03	J1557			01/01/2012	99/99/9999	INJECTION, IMMUNE GLOBULIN (GAMMAPLEX), INTRAVENOUS, NONLYOPHILIZED (E.G., LIQUID) 500 MG	GAMMAPLEX (1X200ML,SINGLE USE) 10 GM/ 200 ML	1	ML	VL	IV	ML	1	EA	0.1	01/01/2012	99/99/9999						
00378-2046-05	J7507			07/13/2011	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	500	EA	BO	PO	EA	1	MG	1	07/13/2011	99/99/9999						
16729-0041-01	J7507			09/30/2011	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 0.5 MG	100	EA	BO	PO	EA	1	MG	0.5	09/30/2011	99/99/9999						
67467-0843-01	J1568			11/04/2011	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (1GM/VIAL,S/D TREATED) 50 MG/ML	1	ML	VL	IV	ML	500	MG	0.1	11/04/2011	09/14/2015						
67467-0843-02	J1568			11/04/2011	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (2.5GM/VIAL,S/D TREATED) 50 MG/ML	1	ML	VL	IV	ML	500	MG	0.1	11/04/2011	09/14/2015						
67467-0843-03	J1568			11/04/2011	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (5GM/VIAL,S/D TREATED) 50 MG/ML	1	ML	VL	IV	ML	500	MG	0.1	11/04/2011	09/14/2015						
67467-0843-04	J1568			11/04/2011	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (10GM/VIAL,S/D TREATED) 50 MG/ML	1	ML	VL	IV	ML	500	MG	0.1	11/04/2011	09/14/2015						
67467-0843-05	J1568			11/04/2011	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (LATEX-FREE) 50 MG/ML	1	ML	VL	IV	ML	500	MG	0.1	11/04/2011	09/14/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00078-0240-61		J7515		01/05/2012	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	SANDIMMUNE (INNER PACK, SOFTGEL) 25 MG	1 EA	BP	PO	EA		25 MG		1	01/05/2012	99/99/9999						
00078-0241-61		J7502		01/05/2012	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	SANDIMMUNE (INNER PACK, SOFTGEL) 100 MG	1 EA	BP	PO	EA		100 MG		1	01/05/2012	99/99/9999						
00078-0467-61		J0895		01/05/2012	99/99/9999	INJECTION, DEFEROXAMINE MESYLATE, 500 MG	DEFERAL (INNER PACK) 500 MG	1 EA	VL	IJ	EA		500 MG		1	01/05/2012	99/99/9999						
00641-6068-01		J2270		02/08/2012	09/16/2015	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (M.D.V.) 10MG/ML	1 ML	VL	IJ	ML		10 MG		1	02/08/2012	09/16/2015						
00641-6070-25		J2270		02/08/2012	09/16/2015	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (S.D.V., 25X1ML) 10MG/ML	25 ML	VL	IJ	ML		10 MG		1	02/08/2012	09/16/2015						
00641-6071-25		J2271		02/08/2012	12/31/2014	INJECTION, MORPHINE SULFATE, 100 MG	MORPHINE SULFATE, (S.D.V., 1MLx25) 15MG/ML	25 ML	VL	IJ	ML		100 MG		0.15	02/08/2012	12/31/2014						
00641-6072-01		J2271		02/08/2012	12/31/2014	INJECTION, MORPHINE SULFATE, 100 MG	MORPHINE SULFATE (M.D.V.) 15MG/ML	1 ML	VL	IJ	ML		100 MG		0.15	02/08/2012	12/31/2014						
00641-6073-25		J2270		02/08/2012	09/16/2015	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (S.D.V.) 5 MG/ ML MORPHINE SULFATE (VIAL, DOSETTE) 8MG/ML	25 ML	VL	IJ	ML		10 MG		0.5	02/08/2012	09/16/2015						
00641-6075-25		J2270		02/08/2012	06/30/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (S.D.V.) 5 MG/ ML MORPHINE SULFATE (VIAL, DOSETTE) 8MG/ML	25 ML	VL	IJ	ML		10 MG		0.8	02/08/2012	06/30/2016						
00054-0163-25		J7517		05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG,	MYCOPHENOLATE MOFETIL, 250 MG	100 EA	BO	PO	EA		250 MG		1	05/04/2009	99/99/9999						
00054-0166-25		J7517		05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG,	MYCOPHENOLATE MOFETIL, 500 MG	100 EA	BO	PO	EA		250 MG		2	05/04/2009	99/99/9999						
00078-0616-05		J7507		02/07/2012	02/11/2015	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	HECORIA (HARD GELATIN) 0.5 MG	100 EA	BO	PO	EA		1 MG		0.5	02/07/2012	02/11/2015						
00078-0618-05		J7507		02/07/2012	02/11/2015	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	HECORIA 5 MG	100 EA	BO	PO	EA		1 MG		5	02/07/2012	02/11/2015						
42254-0110-30		None		01/10/2012	06/01/2014	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE, 2.5 MG	30 EA	BO	PO	EA		2.5 MG		1	01/10/2012	06/01/2014						
62991-1003-01		J7608		10/31/2011	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1 GM	BO	NA	GM		1 GM		1	10/31/2011	99/99/9999						
62991-1003-01	KO	J7608	KO	10/31/2011	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (U.S.P.)	1 GM	BO	NA	GM		1 GM		1	10/31/2011	99/99/9999						
62991-1041-01		J7638		10/31/2011	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 GM	BO	NA	GM		1 MG		1000	10/31/2011	99/99/9999						
62991-1041-01	KO	J7638	KO	10/31/2011	99/99/9999	DEXAMETHASONE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	DEXAMETHASONE SODIUM PHOSPHATE (U.S.P.)	1 GM	BO	NA	GM		1 MG		1000	10/31/2011	99/99/9999						
62991-1486-01		J9190		08/17/2011	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (U.S.P.)	1 GM	BO	NA	GM		500 MG		2	08/17/2011	99/99/9999						
00078-0617-05		J7507		02/07/2012	02/11/2015	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	HECORIA 1 MG	100 EA	BO	PO	EA		1 MG		1	02/07/2012	02/11/2015						
76204-0100-25	KO	J7644	KO	02/01/2012	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	25 ML	SOL	IH	ML		1 MG		0.2	02/01/2012	99/99/9999						
76204-0100-25		J7644		02/01/2012	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	25 ML	SOL	IH	ML		1 MG		0.2	02/01/2012	99/99/9999						
76204-0100-30	KO	J7644	KO	02/01/2012	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	25 ML	SOL	IH	ML		1 MG		0.2	02/01/2012	99/99/9999						
76204-0100-30		J7644		02/01/2012	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	25 ML	SOL	IH	ML		1 MG		0.2	02/01/2012	99/99/9999						
76204-0100-60	KO	J7644	KO	02/01/2012	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML,PF) 0.02%	25 ML	SOL	IH	ML		1 MG		0.2	02/01/2012	99/99/9999						
76204-0100-60		J7644		02/01/2012	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML,PF) 0.02%	25 ML	SOL	IH	ML		1 MG		0.2	02/01/2012	99/99/9999						
76204-0200-25	KO	J7613	KO	02/01/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	02/01/2012	99/99/9999						
76204-0200-25		J7613		02/01/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	02/01/2012	99/99/9999						
76204-0200-30	KO	J7613	KO	02/01/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	02/01/2012	99/99/9999						
76204-0200-30		J7613		02/01/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (30X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	02/01/2012	99/99/9999						
76204-0200-60	KO	J7613	KO	02/01/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (60X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	02/01/2012	99/99/9999						
76204-0200-60		J7613		02/01/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (60X3ML) 0.083%	30 ML	PC	IH	ML		1 MG		0.83	02/01/2012	99/99/9999						
66336-0338-21		None		03/01/2012	06/01/2014	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE, 2.5 MG	21 EA	BO	PO	EA		2.5 MG		1	03/01/2012	06/01/2014						
66336-0338-30		None		04/01/2012	06/01/2014	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM, 2.5 MG	30 EA	BO	PO	EA		2.5 MG		1	04/01/2012	06/01/2014						
00703-5747-11		J9060		06/19/2000	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (M.D.V.) 1 MG/ML	1 ML	VL	IV	ML		10 MG		0.1	06/19/2000	99/99/9999						
49502-0605-30		J7606		07/02/2012	99/99/9999	FORMOTEROL FUMARATE, INHALATION SOLUTION, FDA APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 20 MICROGRAMS	PERFORMIST, 20 MCG/2 ML	30 ML	PC	IH	ML		20 MCG		0.5	07/02/2012	99/99/9999						
49502-0605-30	KO	J7606	KO	07/02/2012	99/99/9999	FORMOTEROL FUMARATE, INHALATION SOLUTION, FDA APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 20 MICROGRAMS	PERFORMIST, 20 MCG/2 ML	30 ML	PC	IH	ML		20 MCG		0.5	07/02/2012	99/99/9999						
76388-0713-25		None		06/22/2012	99/99/9999	BUSULFAN; ORAL, 2 MG	MYLERAN, (FILM-COATED), 2 MG	25 EA	BO	PO	EA		2 MG		1	06/22/2012	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
52152-0538-30		Q0162		07/10/2012	07/11/2012	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE, (FILM-COATED), 4 MG	30	EA	BO	PO	EA	1	MG	4	07/10/2012	07/11/2012						
52152-0539-30		Q0162		07/10/2012	07/11/2012	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE, (FILM-COATED), 8 MG	30	EA	BO	PO	EA	1	MG	8	07/10/2012	07/11/2012						
00781-2067-01	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	100	EA	BO	PO	EA	250	MG	1	05/04/2009	99/99/9999						
00781-2067-05	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	500	EA	BO	PO	EA	250	MG	1	05/04/2009	99/99/9999						
00781-2067-89	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (12X120, HARD GELATIN) 250 MG	1440	EA	BO	PO	EA	250	MG	1	05/04/2009	99/99/9999						
00781-5175-01	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	100	EA	BO	PO	EA	250	MG	2	05/04/2009	99/99/9999						
00781-5175-05	J7517			05/04/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	500	EA	BO	PO	EA	250	MG	2	05/04/2009	99/99/9999						
63323-0690-30	J7608			09/19/2012	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (PDF) 20%	3	ML	SOL	IH	ML	1	GM	0.2	09/19/2012	99/99/9999						
63323-0690-30	KO	J7608	KO	09/19/2012	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (PDF) 20%	3	ML	SOL	IH	ML	1	GM	0.2	09/19/2012	99/99/9999						
66215-0402-01	J1325			10/01/2012	99/99/9999	INJECTION, EPOPROSTENOL, 0.5 MG	VELETRI (SINGLE DOSE, LYOPHILIZED) 1.5 MG	1	EA	VL	IV	EA	0.5	MG	3	10/01/2012	99/99/9999						
66215-0403-01	J1325			10/01/2012	99/99/9999	INJECTION, EPOPROSTENOL, 0.5 MG	VELETRI (SINGLE DOSE, LYOPHILIZED) 0.5 MG	1	EA	VL	IV	EA	0.5	MG	1	10/01/2012	99/99/9999						
00409-1890-01	J2275			08/23/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 2 MG/ML	10	ML	SR	IV	ML	10	MG	0.2	08/23/2012	12/31/2014						
00409-1891-01	J2275			08/06/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 4 MG/ML	10	ML	SR	IV	ML	10	MG	0.4	08/06/2012	12/31/2014						
00409-1894-01	J2275			08/10/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 15 MG/ML	10	ML	SR	IV	ML	10	MG	1.5	08/10/2012	12/31/2014						
50742-0208-01	J7507			10/01/2012	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	EA	PO	EA	1	MG	1	10/01/2012	99/99/9999						
00591-2918-23	KO	J7614	KO	08/20/2012	06/09/2014	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.31 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.20666	08/20/2012	06/09/2014						
00591-2918-23	J7614			08/20/2012	06/09/2014	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.31 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.20666	08/20/2012	06/09/2014						
00591-2919-23	KO	J7614	KO	08/20/2012	08/06/2014	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.63 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.42	08/20/2012	08/06/2014						
00591-2919-23	J7614			08/20/2012	08/06/2014	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.63 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.42	08/20/2012	08/06/2014						
00591-2920-23	KO	J7614	KO	08/20/2012	06/30/2014	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 1.25 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.83333	08/20/2012	06/30/2014						
00591-2920-23	J7614			08/20/2012	06/30/2014	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 1.25 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.83333	08/20/2012	06/30/2014						
00409-1893-01	J2275			08/15/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 10 MG/ML	10	ML	SR	IV	ML	10	MG	1	08/15/2012	12/31/2014						
66993-0021-27	J7614			08/23/2012	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.31 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.20667	08/23/2012	99/99/9999						
66993-0021-27	KO	J7614	KO	08/23/2012	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.31 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.20667	08/23/2012	99/99/9999						
66993-0022-27	J7614			08/23/2012	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.63 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.42	08/23/2012	99/99/9999						
66993-0022-27	KO	J7614	KO	08/23/2012	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 0.63 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.42	08/23/2012	99/99/9999						
66993-0023-27	J7614			08/23/2012	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 1.25 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.83333	08/23/2012	99/99/9999						
66993-0023-27	KO	J7614	KO	08/23/2012	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 1.25 MG/3 ML	24	ML	PC	IH	ML	0.5	MG	0.83333	08/23/2012	99/99/9999						
76125-0900-50	J1561			02/24/2012	99/99/9999	LYOPHILIZED (E.G. LIQUID), 500 MG	GAMMAKED (1X50ML, SINGLE-USE) 10%	1	ML	VL	IJ	ML	500	MG	0.002	02/24/2012	99/99/9999						
00078-0414-20	J7527			01/01/2013	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (6X10) 0.5 MG	60	EA	EA	PO	EA	0.25	MG	2	01/01/2013	99/99/9999						
00078-0414-61	J7527			01/01/2013	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (1X1) 0.5 MG	1	EA	EA	PO	EA	0.25	MG	2	01/01/2013	99/99/9999						
00078-0415-20	J7527			01/01/2013	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (6X10) 0.75 MG	60	EA	EA	PO	EA	0.25	MG	3	01/01/2013	99/99/9999						
00078-0415-61	J7527			01/01/2013	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (1X1) 0.75 MG	1	EA	EA	PO	EA	0.25	MG	3	01/01/2013	99/99/9999						
00078-0417-20	J7527			01/01/2013	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (6X10) 0.25 MG	60	EA	EA	PO	EA	0.25	MG	1	01/01/2013	99/99/9999						
00078-0417-61	J7527			01/01/2013	99/99/9999	EVEROLIMUS, ORAL, 0.25 MG	ZORTRESS (1X1) 0.25 MG	1	EA	EA	PO	EA	0.25	MG	1	01/01/2013	99/99/9999						
38779-0312-03	J7501			10/01/2012	99/99/9999	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (U.S.P.)	5	GM	BO	NA	GM	100	MG	10	10/01/2012	99/99/9999						
38779-0312-04	J7501			10/01/2012	99/99/9999	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (U.S.P.)	25	GM	BO	NA	GM	100	MG	10	10/01/2012	99/99/9999						
38779-0312-06	J7501			10/01/2012	99/99/9999	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (U.S.P.)	1	GM	BO	NA	GM	100	MG	10	10/01/2012	99/99/9999						
00085-1366-03	None			12/05/2012	99/99/9999	TEMODAR, 100 MG, ORAL	TEMODAR, 100 MG	5	EA	BX	PO	EA	100	MG	1	12/05/2012	99/99/9999						
00085-1366-04	None			12/05/2012	99/99/9999	TEMODAR, 100 MG, ORAL	TEMODAR, 100 MG	14	EA	BX	PO	EA	100	MG	1	12/05/2012	99/99/9999						
00085-1417-02	None			12/05/2012	99/99/9999	TEMODAR, 250 MG, ORAL	TEMODAR, 250 MG	5	EA	BX	PO	EA	250	MG	1	12/05/2012	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00085-1425-03		None		12/05/2012	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR, 140 MG	5	EA	BX	PO	EA	20	MG	7	12/05/2012	99/99/9999							
00085-1425-04		None		12/05/2012	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR, 140 MG	14	EA	BX	PO	EA	20	MG	7	12/05/2012	99/99/9999							
00085-1430-03		None		12/05/2012	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR, 180 MG	5	EA	BX	PO	EA	20	MG	9	12/05/2012	99/99/9999							
00085-1430-04		None		12/05/2012	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR, 180 MG	14	EA	BX	PO	EA	20	MG	9	12/05/2012	99/99/9999							
00085-1519-03		None		12/05/2012	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR, 20 MG	5	EA	BX	PO	EA	20	MG	1	12/05/2012	99/99/9999							
00085-1519-04		None		12/05/2012	99/99/9999	TEMODAR, 20 MG, ORAL	TEMODAR, 20 MG	14	EA	BX	PO	EA	20	MG	1	12/05/2012	99/99/9999							
00085-3004-03		None		12/05/2012	99/99/9999	TEMODAR, 5 MG, ORAL	TEMODAR, 5 MG	5	EA	BX	PO	EA	5	MG	1	12/05/2012	99/99/9999							
00085-3004-04		None		12/05/2012	99/99/9999	TEMODAR, 5 MG, ORAL	TEMODAR, 5 MG	14	EA	BX	PO	EA	5	MG	1	12/05/2012	99/99/9999							
62175-0381-37		J7507		09/28/2012	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	BO	PO	EA	1	MG	1	09/28/2012	99/99/9999							
							NOVAPLUS CLADRIBINE (1X10ML,SDV,PF) 1 MG/ML	10	ML	VL	IV	ML	1	MG	1	01/14/2013	10/13/2014							
00069-0201-01		J9065		01/14/2013	10/13/2014	INJECTION, CLADRIBINE, PER 1 MG	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	3	ML	PC	IH	ML	1	MG	0.83333	12/13/2012	99/99/9999							
00378-8270-52		J7613		12/13/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML) 0.083%	3	ML	PC	IH	ML	1	MG	0.83333	12/13/2012	99/99/9999							
00378-8270-52	KO	J7613	KO	12/13/2012	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (25X3ML) 0.083% MYCOPHENOLATE MOFETIL (FILM COATED) 500 MG	3	ML	PC	IH	ML	1	MG	0.83333	12/13/2012	99/99/9999							
16729-0019-01		J7517		05/05/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	100	EA	BO	PO	EA	250	MG	2	05/05/2009	99/99/9999							
16729-0094-01		J7517		05/05/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE, (30 x 3 ML) 3 MG/3 ML-0.5 MG/3 ML	100	EA	BO	PO	EA	250	MG	1	05/05/2009	99/99/9999							
76204-0600-05		J7620		01/01/2013	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE, (60 x 3 ML) 3 MG/3 ML-0.5 MG/3 ML	3	ML	PC	IH	ML	3	MG	0.33333	01/01/2013	99/99/9999							
76204-0600-12		J7620		01/01/2013	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	3	ML	PC	IH	ML	3	MG	0.33333	01/01/2013	99/99/9999							
76204-0002-24		J7614		02/01/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HYDROCHLORIDE, 0.63 MG/3ML,(24X3ML, PF)	3	ML	BO	IH	ML	0.5	MG	0.42	02/01/2013	99/99/9999							
76204-0002-24	KO	J7614	KO	02/01/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HYDROCHLORIDE, 0.63 MG/3ML,(24X3ML, PF)	3	ML	BO	IH	ML	0.5	MG	0.42	02/01/2013	99/99/9999							
76204-0003-24		J7614		02/18/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HYDROCHLORIDE, 1.25 MG/3ML,(24X3ML, PF)	3	ML	BO	IH	ML	0.5	MG	0.83333	02/01/2013	99/99/9999							
76204-0003-24	KO	J7614	KO	02/18/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HYDROCHLORIDE, 1.25 MG/3ML,(24X3ML, PF)	3	ML	BO	IH	ML	0.5	MG	0.83333	02/01/2013	99/99/9999							
		J2275		07/03/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	DURAMORPH (10X10ML,PF) 0.5 MG/ML	10	ML	AM	IJ	ML	10	MG	0.05	07/03/2012	12/31/2014							
00641-6020-10		J2275		07/03/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	DURAMORPH (10X10ML,PF) 1 MG/ML	10	ML	AM	IJ	ML	10	MG	0.1	07/03/2012	12/31/2014							
00641-6019-10		J2275		07/03/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	FENTANYL CITRATE (SINGLE DOSE, 10X2ML) 0.05 MG/ML	10	ML	AM	IJ	ML	0.1	MG	0.5	10/10/2012	99/99/9999							
00641-6024-10		J3010		10/10/2012	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (SINGLE DOSE, 20ML X5) 0.05 MG/ML	5	ML	AM	IJ	ML	0.1	MG	0.5	10/10/2012	99/99/9999							
00641-6026-05		J3010		10/10/2012	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE 0.05 MG/ML	10	ML	AM	IJ	ML	0.1	MG	0.5	11/13/2012	99/99/9999							
00641-6025-10		J3010		11/13/2012	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (25X2ML,USP,SDV,PF) 0.05 MG/ML	25	ML	VL	IJ	ML	0.1	MG	0.5	07/25/2012	99/99/9999							
00641-6027-25		J3010		07/25/2012	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (25X5ML,USP,SDV,PF) 0.05 MG/ML	25	ML	VL	IJ	ML	0.1	MG	0.5	07/25/2012	99/99/9999							
00641-6028-25		J3010		07/25/2012	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (25X20ML,SDV,PF) 0.05 MG/ML	25	ML	VL	IJ	ML	0.1	MG	0.5	10/10/2012	99/99/9999							
00641-6029-25		J3010		10/10/2012	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (S.D.V) 0.05 MG/ML	1	ML	VL	IJ	ML	0.1	MG	0.5	07/25/2012	99/99/9999							
00641-6030-01		J3010		07/25/2012	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	INFUMORPH 200 (1X20ML,PF) 10 MG/ML	1	ML	AM	IJ	ML	10	MG	1	07/25/2012	12/31/2014							
00641-6039-01		J2275		07/25/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	INFUMORPH 500 (1X20ML,PF) 25 MG/ML	1	ML	AM	IJ	ML	10	MG	2.5	07/25/2012	12/31/2014							
00641-6040-01		J2275		07/25/2012	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MILRINONE LACTATE IN DEXTROSE (10X100ML, SINGLE DOSE) 5%-20 MG/100 ML	10	ML	FC	IV	ML	5	MG	0.04	02/23/2011	99/99/9999							
00143-9719-10		J2260		02/23/2011	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE IN DEXTROSE (10X200ML, SINGLE DOSE) 5%-20 MG/100 ML	10	ML	FC	IV	ML	5	MG	0.04	02/23/2011	99/99/9999							
00143-9718-10		J2260		02/23/2011	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	3	ML	PC	IH	ML	0.5	MG	0.20666	03/15/2013	99/99/9999							
00378-9680-44		J7614		03/15/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12,PF) 0.31 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.20666	03/15/2013	99/99/9999							
00378-9680-44	KO	J7614	KO	03/15/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12,PF) 0.31 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.20666	03/15/2013	99/99/9999							
00378-9681-44		J7614		03/15/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	03/15/2013	99/99/9999							
00378-9681-44	KO	J7614	KO	03/15/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	03/15/2013	99/99/9999							
00378-9682-44		J7614		03/15/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	03/15/2013	99/99/9999							
00378-9682-44	KO	J7614	KO	03/15/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	03/15/2013	99/99/9999							
00409-1283-05		J1170		10/22/2012	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HYDROCHLORIDE (USP,ISECURE SINGLE-DOSE) 1 MG/ML	0.5	ML	SR	IJ	ML	4	MG	0.25	10/22/2012	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00944-2656-03	J1566			01/24/2013	99/99/9999	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	GAMMAGARD S/D (IGA<1UG/ML) (SINGLE DOSE) 5 GM	1	EA	VL	IV	EA	500	MG	10	01/24/2013	99/99/9999							
00944-2658-04	J1566			01/24/2013	99/99/9999	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT OTHERWISE SPECIFIED, 500 MG	GAMMAGARD S/D (IGA<1UG/ML) 10 GM	1	EA	VL	IV	EA	500	MG	20	01/24/2013	99/99/9999							
16729-0043-01	J7507			09/30/2011	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	100	EA	BO	PO	EA	1	MG	5	09/30/2011	99/99/9999							
00378-7970-52	J7644			04/03/2013	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	04/03/2013	99/99/9999							
00378-7970-52	KO J7644	KO		04/03/2013	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	04/03/2013	99/99/9999							
43063-0439-30	None			03/14/2013	99/99/9999	METHOTREXATE SODIUM, 2.5 MG, ORAL	METHOTREXATE SODIUM, 2.5 MG	30	EA	BO	PO	EA	2.5	MG	1	03/14/2013	99/99/9999							
44206-0439-40	J1459			06/01/2013	99/99/9999	INJECTION, IMMUNE GLOBULIN (PRIVIGEN), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	PRIVIGEN, (PF,LATEX-FREE), 10%	400	ML	VL	IV	ML	500	MG	0.2	06/01/2013	99/99/9999							
00591-3817-30	J7620			05/13/2013	02/24/2016	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	3	ML	PC	IH	ML	3	MG	0.33333	05/13/2013	02/24/2016							
00591-3817-60	J7620			05/13/2013	02/24/2016	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	3	ML	PC	IH	ML	3	MG	0.33333	05/13/2013	02/24/2016							
67877-0225-01	J7517			03/20/2012	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	100	EA	BO	PO	EA	250	MG	2	03/20/2012	99/99/9999							
45963-0539-30	Q0162			08/29/2011	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	08/29/2011	99/99/9999							
00093-7600-41	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 20 MG	14	EA	BO	PO	EA	20	MG	1	08/12/2013	99/99/9999							
00469-0647-73	J7599			08/06/2013	12/31/2013	IMMUNOSUPPRESSIVE DRUG, NOT OTHERWISE CLASSIFIED	ASTAGRAF XL 0.5 MG	30	EA	BO	PO	EA	1	MG	1	08/06/2013	12/31/2013							
00469-0677-73	J7599			08/06/2013	12/31/2013	IMMUNOSUPPRESSIVE DRUG, NOT OTHERWISE CLASSIFIED	ASTAGRAF XL 1 MG	30	EA	BO	PO	EA	1	MG	1	08/06/2013	12/31/2013							
00469-0687-73	J7599			08/06/2013	12/31/2013	IMMUNOSUPPRESSIVE DRUG, NOT OTHERWISE CLASSIFIED	ASTAGRAF XL 5 MG	30	EA	BO	PO	EA	1	MG	1	08/06/2013	12/31/2013							
00781-2691-44	None			08/12/2013	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	14	EA	BO	PO	EA	5	MG	1	08/12/2013	99/99/9999							
00781-2691-75	None			08/12/2013	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	5	EA	BO	PO	EA	5	MG	1	08/12/2013	99/99/9999							
00781-2692-44	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14	EA	BO	PO	EA	20	MG	1	08/12/2013	99/99/9999							
00781-2692-75	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5	EA	BO	PO	EA	20	MG	1	08/12/2013	99/99/9999							
00781-2693-44	None			08/12/2013	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14	EA	BO	PO	EA	100	MG	1	08/12/2013	99/99/9999							
00781-2693-75	None			08/12/2013	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5	EA	BO	PO	EA	100	MG	1	08/12/2013	99/99/9999							
00781-2694-44	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	14	EA	BO	PO	EA	20	MG	7	08/12/2013	99/99/9999							
00781-2694-75	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	5	EA	BO	PO	EA	20	MG	7	08/12/2013	99/99/9999							
00781-2695-44	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	14	EA	BO	PO	EA	20	MG	9	08/12/2013	99/99/9999							
00781-2695-75	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	5	EA	BO	PO	EA	20	MG	9	08/12/2013	99/99/9999							
00603-1567-56	J7510			07/01/2013	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (CHERRY) 15 MG/5 ML	240	ML	BO	PO	ML	5	MG	0.6	07/01/2013	99/99/9999							
00603-1567-58	J7510			07/01/2013	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (CHERRY) 15 MG/5 ML	480	ML	BO	PO	ML	5	MG	0.6	07/01/2013	99/99/9999							
00904-5789-61	J8499			09/13/2013	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (10X10,USP,HARD GELATIN) 200 MG	100	EA	BX	PO	EA	1	MG	1	09/13/2013	99/99/9999							
00904-5790-61	J8499			09/13/2013	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (10X10,USP) 400 MG	100	EA	BX	PO	EA	1	MG	1	09/13/2013	99/99/9999							
00781-7146-64	J7620			07/30/2013	03/14/2017	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	30	ML	VL	IH	ML	3	MG	0.33333	07/30/2013	03/14/2017							
00093-7602-57	None			08/12/2013	99/99/9999	TEMODAR, 250 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 250 MG	5	EA	BO	PO	EA	250	MG	1	08/12/2013	99/99/9999							
00093-7601-41	None			08/12/2013	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 100 MG	14	EA	BO	PO	EA	100	MG	1	08/12/2013	99/99/9999							
00093-7601-57	None			08/12/2013	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 100 MG	5	EA	BO	PO	EA	100	MG	1	08/12/2013	99/99/9999							
00093-7638-41	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 140 MG	14	EA	BO	PO	EA	20	MG	7	08/12/2013	99/99/9999							
00093-7638-57	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 140 MG	5	EA	BO	PO	EA	20	MG	7	08/12/2013	99/99/9999							
00093-7639-57	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 180 MG	5	EA	BO	PO	EA	20	MG	9	08/12/2013	99/99/9999							
00093-7639-41	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 180 MG	14	EA	BO	PO	EA	20	MG	9	08/12/2013	99/99/9999							
00093-7600-57	None			08/12/2013	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 20 MG	5	EA	BO	PO	EA	20	MG	1	08/12/2013	99/99/9999							
00093-7599-41	None			08/12/2013	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 5 MG	14	EA	BO	PO	EA	5	MG	1	08/12/2013	99/99/9999							
00093-7599-57	None			08/12/2013	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE (UNIT-OF-USE) 5MG	5	EA	BO	PO	EA	5	MG	1	08/12/2013	99/99/9999							
25021-0207-05	J9000			11/01/2013	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HYDROCHLORIDE (USP,STERILE,SDV) 2 MG/ML	5	ML	VL	IV	ML	10	MG	0.2	11/01/2013	99/99/9999							
25021-0207-25	J9000			11/01/2013	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HYDROCHLORIDE (USP,STERILE,SDV) 2 MG/ML	25	ML	VL	IV	ML	10	MG	0.2	11/01/2013	99/99/9999							
25021-0207-51	J9000			11/01/2013	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HYDROCHLORIDE (USP,STERILE,SDV) 2 MG/ML	100	ML	VL	IV	ML	10	MG	0.2	11/01/2013	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00378-9671-58		J7620		09/26/2013	01/27/2016	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	IPRATROPIUM BROMIDE AND ALBUTEROL SULFATE (30X3ML,5 VIALS/POUCH) 3 MG/3 ML-0.5 MG/3 ML	3	ML	PC	IH	ML	3	MG	0.33333	09/26/2013	01/27/2016							
10122-0820-56		J7682		09/20/2013	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	BETHKIS 300 MG/4 ML	56	EA	PC	IH	ML	300	MG	0.25	09/20/2013	99/99/9999							
10122-0820-56	KO	J7682	KO	09/20/2013	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	BETHKIS 300 MG/4 ML	56	EA	PC	IH	ML	300	MG	0.25	09/20/2013	99/99/9999							
00781-5238-64		Q0162		12/18/2008	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,3X10,STRAWBERRY) 4 MG	30	EA	BX	PO	EA	1	MG	4	12/18/2008	99/99/9999							
00781-2696-75		None		09/30/2013	99/99/9999	TEMODAR, 250 MG, ORAL	TEMOZOLOMIDE 250 MG	5	EA	BO	PO	EA	250	MG	1	09/30/2013	99/99/9999							
00469-0647-73		J7508		01/01/2014	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ASTAGRAF XL), ORAL, 0.1 MG	ASTAGRAF XL 0.5 MG	30	EA	BO	PO	EA	0.1	MG	5	01/01/2014	99/99/9999							
00469-0677-73		J7508		01/01/2014	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ASTAGRAF XL), ORAL, 0.1 MG	ASTAGRAF XL 1 MG	30	EA	BO	PO	EA	0.1	MG	10	01/01/2014	99/99/9999							
00469-0687-73		J7508		01/01/2014	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ASTAGRAF XL), ORAL, 0.1 MG	ASTAGRAF XL 5 MG	30	EA	BO	PO	EA	0.1	MG	50	01/01/2014	99/99/9999							
00093-9652-01		Q0164		01/01/2014	04/18/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (USP,FILM-COATED) 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2014	04/16/2018							
00378-5110-01		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999							
00781-5021-01		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999							
16590-0327-10		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (FILM-COATED) 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2014	06/01/2014							
21695-0572-30		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (FILM-COATED) 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	06/01/2014							
33358-0300-10		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999							
33358-0300-20		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	20	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999							
33358-0300-30		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999							
33358-0300-60		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	60	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999							
35356-0325-00		Q0164		01/01/2014	01/01/2015	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (FILM-COATED) 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2014	01/01/2015							
38779-0180-04		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	25	GM	BO	NA	GM	5	MG	200	01/01/2014	99/99/9999							
38779-0180-05		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	100	GM	BO	NA	GM	5	MG	200	01/01/2014	99/99/9999							
38779-0180-08		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	500	GM	BO	NA	GM	5	MG	200	01/01/2014	99/99/9999							
51079-0542-01		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (USP) 10 MG	1	EA	BP	PO	WA	5	MG	2	01/01/2014	99/99/9999							
51079-0542-20		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (10X10) 10 MG	100	EA	BX	PO	EA	5	MG	2	01/01/2014	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51552-0074-05		Q0164		01/01/2014	01/01/2015	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	100	GM	BO	NA	GM	5	MG	200	01/01/2014	01/01/2015						
51552-0074-09		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	25	GM	BO	NA	GM	5	MG	200	01/01/2014	99/99/9999						
51927-2134-00		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	1	GM	BO	NA	GM	5	MG	200	01/01/2014	99/99/9999						
52959-0391-15		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	COMPAZINE 10 MG	15	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
52959-0476-02		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	120	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
52959-0476-10		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
52959-0476-15		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
52959-0476-20		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (FILM-COATED) 10 MG	20	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
52959-0476-24		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	24	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
52959-0476-30		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
52959-0476-60		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54569-0355-00		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE (FILM-COATED) 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54569-0355-02		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE (FILM-COATED) 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54868-1082-00		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54868-1082-01		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54868-1082-02		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54868-1082-03		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54868-1082-04		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
54868-1082-05		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Scouts of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-1082-06		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	90 EA	BO	PO	EA		5 MG		2	01/01/2014	99/99/9999						
54888-1082-03		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100 EA	NA	PO	EA		5 MG		2	01/01/2014	99/99/9999						
55045-1126-02		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10 EA	BO	PO	EA		5 MG		2	01/01/2014	06/01/2014						
55045-1126-03		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	5 EA	BO	PO	EA		5 MG		2	01/01/2014	06/01/2014						
55045-1126-04		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE 10 MG	12 EA	BO	PO	EA		5 MG		2	01/01/2014	06/01/2014						
55045-1126-06		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60 EA	BO	PO	EA		5 MG		2	01/01/2014	06/01/2014						
55045-1126-07		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20 EA	BO	PO	EA		5 MG		2	01/01/2014	06/01/2014						
55045-1126-08		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30 EA	BO	PO	EA		5 MG		2	01/01/2014	06/01/2014						
55289-0224-04		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	4 EA	BO	PO	EA		5 MG		2	01/01/2014	99/99/9999						
55289-0224-06		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10 EA	BO	PO	EA		5 MG		2	01/01/2014	99/99/9999						
55289-0224-12		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	12 EA	BO	PO	EA		5 MG		2	01/01/2014	99/99/9999						
58016-0706-00		Q0164		01/01/2014	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100 EA	BO	PO	EA		5 MG		2	01/01/2014	01/31/2014						
58016-0706-02		Q0164		01/01/2014	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	120 EA	BO	PO	EA		5 MG		2	01/01/2014	01/31/2014						
58016-0706-03		Q0164		01/01/2014	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	150 EA	BO	PO	EA		5 MG		2	01/01/2014	01/31/2014						
58016-0706-08		Q0164		01/01/2014	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	8 EA	NA	PO	EA		5 MG		2	01/01/2014	01/31/2014						
58016-0706-30		Q0164		01/01/2014	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30 EA	BO	PO	EA		5 MG		2	01/01/2014	01/31/2014						
58016-0706-60		Q0164		01/01/2014	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	6 EA	BO	PO	EA		5 MG		2	01/01/2014	01/31/2014						
58016-0706-90		Q0164		01/01/2014	01/31/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	90 EA	BO	PO	EA		5 MG		2	01/01/2014	01/31/2014						
59746-0115-06		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100 EA	BO	PO	EA		5 MG		2	01/01/2014	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
62991-1122-02		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	100	GM	BO	NA	GM	5	MG	200	01/01/2014	99/99/9999						
63629-1335-03		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
63874-0490-01		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	100	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63874-0490-06		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	6	EA	NP	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63874-0490-08		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	8	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63874-0490-10		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
63874-0490-15		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63874-0490-20		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	20	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63874-0490-28		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	28	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63874-0490-30		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63874-0490-60		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
66336-0921-15		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	5	MG	2	01/01/2014	06/01/2014						
66336-0921-60		Q0164		01/01/2014	06/01/2014	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	60	EA	BO	PO	EA	5	MG	2	01/01/2014	06/01/2014						
54868-3189-00		Q0167		01/01/2014	02/03/2016	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	25	EA	BO	PO	EA	2.5	MG	2	01/01/2014	02/03/2016						
54868-3189-01		Q0167		01/01/2014	02/03/2016	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL 5 MG	100	EA	BO	PO	EA	2.5	MG	2	01/01/2014	02/03/2016						
54868-3189-02		Q0167		01/01/2014	02/03/2016	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL 5 MG	60	EA	BO	PO	EA	2.5	MG	2	01/01/2014	02/03/2016						
58016-0951-30		Q0167		01/01/2014	01/31/2014	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	30	EA	BO	PO	EA	2.5	MG	2	01/01/2014	01/31/2014						
58016-0951-60		Q0167		01/01/2014	01/31/2014	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	60	EA	BO	PO	EA	2.5	MG	2	01/01/2014	01/31/2014						
58016-0951-90		Q0167		01/01/2014	01/31/2014	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	90	EA	BO	PO	EA	2.5	MG	2	01/01/2014	01/31/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00069-5420-66		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	VISTARIL 50 MG	100	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
00185-0615-01		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
00185-0615-05		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
00463-6156-10		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMACOT 25 MG	1000	EA	NA	PO	WA	12.5	MG	2	01/01/2014	02/03/2016						
00555-0302-02		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
00555-0302-04		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
00555-0324-02		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	100	EA	BO	PO	EA	25	MG	4	01/01/2014	99/99/9999						
00591-0801-05		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
00591-5307-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
00591-5307-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	1000	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
00591-5319-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	100	EA	BO	PO	EA	12.5	MG	4	01/01/2014	99/99/9999						
00603-1584-54		Q0169		01/01/2014	06/11/2018	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE PLAIN (USP) 6.25 MG/5 ML	118	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	06/11/2018						
00603-1584-58		Q0169		01/01/2014	06/11/2018	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE PLAIN (USP) 6.25 MG/5 ML	473	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	06/11/2018						
00603-5438-21		Q0169		01/01/2014	01/09/2017	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	100	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/09/2017						
00603-5438-32		Q0169		01/01/2014	01/09/2017	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	1000	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/09/2017						
00603-5439-21		Q0169		01/01/2014	01/09/2017	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 50 MG	100	EA	BO	PO	EA	12.5	MG	4	01/01/2014	01/09/2017						
00781-1048-01		Q0175		01/01/2014	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	100	EA	BO	PO	EA	4	MG	2	01/01/2014	99/99/9999						
00781-1048-13		Q0175		01/01/2014	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	100	EA	BX	PO	EA	4	MG	2	01/01/2014	99/99/9999						
00781-1049-01		Q0175		01/01/2014	99/99/9999	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 16 MG	100	EA	BO	PO	EA	4	MG	4	01/01/2014	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00781-1830-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PI	EA	12.5	MG	2	01/01/2014	99/99/9999						
00781-1830-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	1000	EA	BO	PI	EA	12.5	MG	2	01/01/2014	99/99/9999						
00781-1832-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	100	EA	BO	PO	EA	12.5	MG	4	01/01/2014	99/99/9999						
00904-5840-61		Q0169		01/01/2014	08/14/2015	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	100	EA	BX	PO	EA	12.5	MG	2	01/01/2014	08/14/2015						
10702-0003-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	100	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
10702-0003-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	1000	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
10702-0004-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 50 MG	100	EA	BO	PO	EA	12.5	MG	4	01/01/2014	99/99/9999						
16590-0191-10		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
16590-0191-15		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	15	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
16590-0191-20		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
16590-0191-30		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	30	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
16590-0191-60		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
16590-0191-90		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	90	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
21695-0415-60		Q0175		01/01/2014	06/01/2014	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE (FILM-COATED) 8 MG	60	EA	BO	PO	EA	4	MG	2	01/01/2014	06/01/2014						
21695-0453-10		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
21695-0453-15		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	15	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
21695-0453-20		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
21695-0453-25		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	25	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
21695-0703-04		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (1X120ML TROPICAL FRUIT) 6.25 MG/5 ML	120	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
33358-0302-08		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	8 EA	BO	PO	EA	12.5 MG			2	01/01/2014	99/99/9999						
33358-0302-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10 EA	BO	PO	EA	12.5 MG			2	01/01/2014	99/99/9999						
33358-0302-30		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	30 EA	BO	PO	EA	12.5 MG			2	01/01/2014	99/99/9999						
33358-0302-60		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	60 EA	BO	PO	EA	12.5 MG			2	01/01/2014	99/99/9999						
35356-0096-60		Q0169		01/01/2014	01/01/2015	PERPHENAZINE, 4MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	60 EA	BO	PO	EA	4 MG			2	01/01/2014	01/01/2015						
35356-0098-90		Q0169		01/01/2014	01/01/2015	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE 100 MG	90 EA	BO	PO	EA	12.5 MG			8	01/01/2014	01/01/2015						
49999-0036-12		Q0169		01/01/2014	01/01/2015	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	12 EA	BO	PO	EA	12.5 MG			8	01/01/2014	01/01/2015						
49999-0036-60		Q0169		01/01/2014	01/01/2015	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	60 EA	BO	PO	EA	12.5 MG			8	01/01/2014	01/01/2015						
49999-0090-05		Q0169		01/01/2014	12/31/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	5 EA	BO	PO	EA	12.5 MG			2	01/01/2014	12/31/2016						
49999-0090-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10 EA	BO	PO	EA	12.5 MG			2	01/01/2014	99/99/9999						
49999-0090-12		Q0169		01/01/2014	12/31/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12 EA	BO	PO	EA	12.5 MG			2	01/01/2014	12/31/2016						
49999-0090-15		Q0169		01/01/2014	12/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15 EA	BO	PO	EA	12.5 MG			2	01/01/2014	12/31/2014						
49999-0090-20		Q0169		01/01/2014	06/01/2017	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20 EA	BO	PO	EA	12.5 MG			2	01/01/2014	06/01/2017						
49999-0090-30		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30 EA	BO	PO	EA	12.5 MG			2	01/01/2014	99/99/9999						
49999-0090-60		Q0169		01/01/2014	12/31/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60 EA	BO	PO	EA	12.5 MG			2	01/01/2014	12/31/2016						
49999-0262-04		Q0169		01/01/2014	01/01/2015	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120 ML	BO	PO	ML	12.5 MG			0.1	01/01/2014	01/01/2015						
50383-0801-16		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (CHERRY) 6.25 MG/5 ML	473 ML	BO	PO	ML	12.5 MG			0.1	01/01/2014	99/99/9999						
51079-0078-01		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (USP) 50 MG	1 EA	NA	PO	EA	25 MG			2	01/01/2014	99/99/9999						
51079-0078-20		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (10X10) 50 MG	100 EA	BX	PO	EA	25 MG			2	01/01/2014	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
51079-0895-01		Q0169		01/01/2014	09/02/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	1	EA	BX	PO	EA	12.5	MG	2	01/01/2014	09/02/2016						
51079-0895-20		Q0169		01/01/2014	09/02/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (10X10) 25 MG	100	EA	BX	PO	EA	12.5	MG	2	01/01/2014	09/02/2016						
51552-0979-04		Q0177		01/01/2014	01/01/2015	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (U.S.P.)	25	GM	BO	NA	GM	25	MG	40	01/01/2014	01/01/2015						
51927-2316-00		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (U.S.P.)	1	GM	JR	NA	GM	25	MG	40	01/01/2014	99/99/9999						
52959-0804-04		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	120	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	99/99/9999						
52959-0804-08		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	240	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	99/99/9999						
52959-0833-06		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	6	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
52959-0833-20		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
54569-1046-00		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE PLAIN 6.25 MG/5 ML	120	ML	BO	PO	ML	12.5	ML	0.1	01/01/2014	99/99/9999						
54569-1754-00		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	12	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54569-1754-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54569-1754-05		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54569-1754-06		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54569-1754-09		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	30	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54569-2571-01		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
54569-4168-00		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	5	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1323-00		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1323-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1323-02		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54868-1323-04		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1323-05		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1323-06		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1323-07		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1323-08		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	50	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
54868-1854-04		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	25	MG	2	01/01/2014	02/03/2016						
54868-1867-00		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	99/99/9999						
54868-2302-00		Q0161		01/01/2014	02/03/2016	CHLORPROMAZINE HYDROCHLORIDE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 50 MG	10	EA	BO	PO	EA	5	MG	10	01/01/2014	02/03/2016						
54868-2302-02		Q0161		01/01/2014	02/03/2016	CHLORPROMAZINE HYDROCHLORIDE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 50 MG	100	EA	BO	PO	EA	5	MG	10	01/01/2014	02/03/2016						
54868-2347-00		Q0161		01/01/2014	02/03/2016	CHLORPROMAZINE HYDROCHLORIDE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 100 MG	100	EA	BO	PO	EA	5	MG	20	01/01/2014	02/03/2016						
54868-2464-00		Q0161		01/01/2014	99/99/9999	CHLORPROMAZINE HYDROCHLORIDE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 25 MG	30	EA	BO	PO	EA	5	MG	5	01/01/2014	99/99/9999						
54868-2464-02		Q0161		01/01/2014	99/99/9999	CHLORPROMAZINE HYDROCHLORIDE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE HCL 25 MG	60	EA	NA	PO	EA	5	MG	5	01/01/2014	99/99/9999						
54868-2684-01		Q0161		01/01/2014	02/03/2016	CHLORPROMAZINE HYDROCHLORIDE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CHLORPROMAZINE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
54868-2687-01		Q0175		01/01/2014	02/03/2016	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	100	EA	BO	PO	EA	4	MG	2	01/01/2014	02/03/2016						
54868-2687-02		Q0175		01/01/2014	02/03/2016	PERPHENAZINE, 4 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PERPHENAZINE 8 MG	60	EA	BO	PO	EA	4	MG	2	01/01/2014	02/03/2016						
54868-2844-00		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	60	EA	BO	PO	EA	12.5	MG	4	01/01/2014	99/99/9999						
54868-2844-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	30	EA	BO	PO	EA	12.5	MG	4	01/01/2014	99/99/9999						
54868-4109-00		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	100	EA	BO	PO	EA	12.5	MG	8	01/01/2014	99/99/9999						
55045-1596-00		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55045-1596-01		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	120	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1596-02		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1596-03		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1596-04		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	NA	pO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1596-05		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1596-06		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1596-08		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1596-09		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	90	EA	BO	PO	EA	12.5	MG	2	01/01/2014	06/01/2014						
55045-1643-09		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (CHERRY) 6.25 MG/5 ML	118	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	06/01/2014						
55045-1661-00		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	NA	PO	EA	25	MG	2	01/01/2014	06/01/2014						
55045-1661-01		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	120	EA	NA	PO	EA	25	MG	2	01/01/2014	06/01/2014						
55045-1661-02		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20	EA	NA	PO	EA	25	MG	2	01/01/2014	06/01/2014						
55045-1661-03		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	40	EA	NA	PO	EA	25	MG	2	01/01/2014	06/01/2014						
55045-1661-06		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	60	EA	NA	PO	EA	25	MG	2	01/01/2014	06/01/2014						
55045-1661-08		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30	EA	BO	PO	EA	25	MG	2	01/01/2014	06/01/2014						
55045-1661-09		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	90	EA	NA	PO	EA	25	MG	2	01/01/2014	06/01/2014						
55289-0354-10		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	10	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
55289-0464-15		Q0169		01/01/2014	04/12/2018	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	12.5	MG	2	01/01/2014	04/12/2018						
55289-0464-79		Q0169		01/01/2014	04/12/2018	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	1	EA	BO	PO	EA	12.5	MG	2	01/01/2014	04/12/2018						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
55289-0531-04		Q0169		01/01/2014	07/12/2017	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 50 MG	4	EA	BO	PO	EA	12.5	MG	4	01/01/2014	07/12/2017							
58016-0424-00		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-02		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	120	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-03		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	150	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-10		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-12		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-15		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-20		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-30		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-40		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	40	EA	NA	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-48		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	48	EA	NA	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-50		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	50	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-60		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	60	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-73		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	300	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-89		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	200	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0424-90		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	90	EA	BO	PO	EA	12.5	MG	2	01/01/2014	01/31/2014							
58016-0464-10		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	10	EA	BO	PO	EA	12.5	MG	4	01/01/2014	01/31/2014							
58016-0464-15		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	15	EA	BO	PO	EA	12.5	MG	4	01/01/2014	01/31/2014							
58016-0464-20		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20	EA	BO	PO	EA	12.5	MG	4	01/01/2014	01/31/2014							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
58016-0464-30		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30	EA	BO	PO	EA	12.5	MG	4	01/01/2014	01/31/2014						
58016-4008-01		Q0169		01/01/2014	01/31/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120	ML	NA	PO	ML	12.5	MG	0.1	01/01/2014	01/31/2014						
58864-0761-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (REDI-SCRIPT) 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
58864-0761-30		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
58864-0761-42		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	42	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
60432-0608-04		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (TROPICAL FRUIT) 6.25 MG/5 ML	118	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	99/99/9999						
60432-0608-16		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (TROPICAL FRUIT) 6.25 MG/5 ML	473	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	99/99/9999						
60760-0830-20		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
63629-1742-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	15	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
63629-1742-02		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	30	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
63629-1742-03		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
63629-1742-04		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	20	EA	BO	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
63629-1870-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	120	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	99/99/9999						
63629-1870-02		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 6.25 MG/5 ML	240	ML	BO	PO	ML	12.5	MG	0.1	01/01/2014	99/99/9999						
63739-0213-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE (USP) 25 MG	100	EA	BX	PO	EA	12.5	MG	2	01/01/2014	99/99/9999						
63874-0370-01		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	100	EA	BO	PO	EA	12.5	MG	2	01/01/2014	02/03/2016						
63874-0370-08		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	8	EA	BO	PO	EA	12.5	MG	2	01/01/2014	02/03/2016						
63874-0370-10		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	10	EA	BO	PO	EA	12.5	MG	2	01/01/2014	02/03/2016						
63874-0370-12		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	12	EA	BO	PO	EA	12.5	MG	2	01/01/2014	02/03/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63874-0370-15		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	15 EA	BO	PO	EA	12.5 MG			2	01/01/2014	02/03/2016						
63874-0370-20		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20 EA	BO	PO	EA	12.5 MG			2	01/01/2014	02/03/2016						
63874-0370-24		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	24 EA	BO	PO	EA	12.5 MG			2	01/01/2014	02/03/2016						
63874-0370-30		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	30 EA	BO	PO	EA	12.5 MG			2	01/01/2014	02/03/2016						
63874-0370-40		Q0169		01/01/2014	02/03/2016	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	40 EA	BO	PO	EA	12.5 MG			2	01/01/2014	02/03/2016						
63874-0370-60		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60 EA	BO	PO	EA	12.5 MG			2	01/01/2014	99/99/9999						
63874-0712-12		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 6.25 MG/5 ML	120 ML	NA	PO	ML	12.5 MG	0.1		01/01/2014	99/99/9999							
63874-0757-01		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-04		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	120 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-10		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	10 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-15		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	15 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-20		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	20 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-21		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	21 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-24		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	24 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-28		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	28 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-30		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-60		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	60 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
63874-0757-90		Q0177		01/01/2014	02/03/2016	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	90 EA	BO	PO	EA	25 MG			2	01/01/2014	02/03/2016						
66336-0085-10		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	10 EA	BO	PO	EA	12.5 MG			2	01/01/2014	06/01/2014						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
66336-0085-12		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	12 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
66336-0085-20		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	20 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
66336-0085-25		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	25 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
66336-0085-30		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE 25 MG	30 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
66336-0085-60		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
68382-0041-01		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	100 EA	BO	PO	EA	12.5 MG		2	01/01/2014	99/99/9999							
68382-0041-10		Q0169		01/01/2014	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	1000 EA	BO	PO	EA	12.5 MG		2	01/01/2014	99/99/9999							
68387-0468-30		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	30 EA	BO	PO	EA	25 MG		2	01/01/2014	06/01/2014							
68387-0469-30		Q0177		01/01/2014	06/01/2014	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 100 MG	30 EA	BO	PO	EA	25 MG		4	01/01/2014	06/01/2014							
68387-0536-12		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	12 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
68387-0536-30		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	30 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
68387-0536-60		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	60 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
68387-0536-90		Q0169		01/01/2014	06/01/2014	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HYDROCHLORIDE 25 MG	90 EA	BO	PO	EA	12.5 MG		2	01/01/2014	06/01/2014							
00378-8270-93		J7613		01/22/2013	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (3MLX30) 0.083%	3 ML	PC	IH	ML	1 MG	0.83		01/22/2013	99/99/9999							
00378-8270-93	KO	J7613	KO	01/22/2013	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (3MLX30) 0.083%	3 ML	PC	IH	ML	1 MG	0.83		01/22/2013	99/99/9999							
00378-8270-91		J7613		04/11/2013	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (60X3ML) 0.083%	3 ML	PC	IH	ML	1 MG	0.83		04/11/2013	99/99/9999							
00378-8270-91	KO	J7613	KO	04/11/2013	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (60X3ML) 0.083%	3 ML	PC	IH	ML	1 MG	0.83		04/11/2013	99/99/9999							
00378-6990-93		J7613		10/07/2009	03/06/2014	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (1X30) 0.083%	3 ML	PC	IH	ML	1 MG	0.83		10/07/2009	03/06/2014							
00378-6990-93	KO	J7613	KO	10/07/2009	03/06/2014	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE (1X30) 0.083%	3 ML	PC	IH	ML	1 MG	0.83		10/07/2009	03/06/2014							
00378-9671-93		J7620		06/13/2013	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	ALBUTEROL SULFATE (30X3ML, 1 VIAL/POUCH) 3 MG/3 ML-0.5 MG/3 ML	3 ML	PC	IH	ML	3 MG	0.33333		06/13/2013	99/99/9999							
00378-7970-93		J7644		02/19/2013	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5 ML	PC	IH	ML	1 MG	0.2		02/19/2013	99/99/9999							
00378-7970-93	KO	J7644	KO	02/19/2013	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5 ML	PC	IH	ML	1 MG	0.2		02/19/2013	99/99/9999							
00378-2047-05		J7507		07/13/2011	10/13/2015	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	500 EA	BO	PO	EA	1 MG		5	07/13/2011	10/13/2015							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63874-0490-12		Q0164		01/01/2014	02/03/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	12	EA	BO	PO	EA	5	MG	2	01/01/2014	02/03/2016						
63629-1335-02		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	30	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
63629-1335-01		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	10	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
58864-0644-42		Q0164		01/01/2014	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (REDI-SCRIPT) 10 MG	42	EA	BO	PO	EA	5	MG	2	01/01/2014	99/99/9999						
58016-0951-00		Q0167		01/01/2014	01/31/2014	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 5 MG	100	EA	BO	PO	EA	2.5	MG	2	01/01/2014	01/31/2014						
00591-0801-01		Q0177		01/01/2014	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	100	EA	BO	PO	EA	25	MG	2	01/01/2014	99/99/9999						
59730-6502-01		J1556		12/19/2012	99/99/9999	INJECTION, IMMUNE GLOBULIN (BIVIGAM), 500 MG	BIVIGAM (LATEX-FREE) 100 MG/ML	50	ML	VL	IV	ML	500	MG	0.2	12/19/2012	99/99/9999						
59730-6503-01		J1556		12/19/2012	99/99/9999	INJECTION, IMMUNE GLOBULIN (BIVIGAM), 500 MG	BIVIGAM (LATEX-FREE) 100 MG/ML	100	ML	VL	IV	ML	500	MG	0.2	12/19/2012	99/99/9999						
51079-0028-20		J7507		08/06/2013	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (10X10;HARD GELATIN) 5 MG	100	EA	BX	PO	EA	1	MG	5	08/06/2013	99/99/9999						
51079-0817-20		J7507		08/06/2013	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (10X10;HARD GELATIN) 0.5 MG	100	EA	BX	PO	EA	1	MG	0.5	08/06/2013	99/99/9999						
00093-4085-63		J7682		11/19/2013	99/99/9999	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5	ML	PC	IH	ML	300	ML	0.2	11/19/2013	99/99/9999						
00093-4085-63	KO	J7682	KO	11/19/2013	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5	ML	PC	IH	ML	300	ML	0.2	11/19/2013	99/99/9999						
00143-9738-05		J7506		07/03/2013	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE 20 MG	500	EA	BO	PO	EA	5	MG	4	07/03/2013	12/31/2015						
00378-4201-78		J7518		01/08/2014	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (FILM-COATED) 180 MG	120	EA	BO	PO	EA	180	MG	1	01/08/2014	99/99/9999						
00378-4202-78		J7518		01/08/2014	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (FILM-COATED) 360 MG	120	EA	BO	PO	EA	180	MG	2	01/08/2014	99/99/9999						
68209-0843-01		J1568		03/21/2012	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGRAM (1GM/1VIAL,SD TREATED) 50MG/ML	20	ML	VL	IV	ML	500	MG	0.1	03/21/2012	09/14/2015						
68209-0843-02		J1568		03/21/2012	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGRAM (PF.SUCROSE-FREE) 50MG/ML	50	ML	VL	IV	ML	500	MG	0.1	03/21/2012	09/14/2015						
68209-0843-03		J1568		03/21/2012	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGRAM (PF.SUCROSE-FREE) 50MG/ML	100	ML	VL	IV	ML	500	MG	0.1	03/21/2012	09/14/2015						
68209-0843-04		J1568		03/21/2012	09/14/2015	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGRAM (PF.SUCROSE-FREE) 50MG/ML	200	ML	VL	IV	ML	500	MG	0.1	03/21/2012	09/14/2015						
44206-0455-10		J1559		10/01/2013	99/99/9999	INJECTION, IMMUNE GLOBULIN (HIZENTRA), 100 MG	HIZENTRA (SINGLE-USE VIAL,PF) 20% GAMMAPLEX (1X50ML,SINGLE USE) 2.5 GM/50ML	50	ML	VL	SC	ML	100	MG	2	10/01/2013	99/99/9999						
64208-8234-05		J1557		07/26/2013	01/31/2015	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMMAPLEX (1X100ML,SINGLE USE) 5 GM/100ML	100	ML	VL	IV	ML	500	MG	0.1	07/26/2013	01/31/2015						
64208-8234-06		J1557		07/26/2013	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	GAMMAPLEX (1X200ML,SINGLE USE) 10 GM/200ML	200	ML	VL	IV	ML	500	MG	0.1	07/26/2013	99/99/9999						
64208-8234-07		J1557		07/26/2013	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	MORPHINE SULFATE (USP, PUMP-JET) 1 MG/ML	30	ML	SR	IJ	ML	10	MG	0.1	11/01/2013	99/99/9999						
76329-1911-01		J2270		11/01/2013	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	PREDNISOLONE (4X60 ML,RED CHERRY) 15 MG/5 ML	60	ML	BO	PO	ML	5	MG	0.6	09/27/2013	02/03/2016						
54569-4827-01		J7510		09/27/2013	02/03/2016	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (DYE-FREE, GRAPE) 15 MG/5 ML	240	ML	BO	PO	ML	5	MG	0.6	01/21/2014	99/99/9999						
54569-5749-00		J7510		01/21/2014	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	CAPECITABINE (USP,FILM-COATED) 150 MG	60	EA	BO	PO	EA	150	MG	1	03/07/2014	99/99/9999						
00093-7473-06		None		03/07/2014	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM-COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	03/07/2014	99/99/9999						
00093-7474-89		None		03/07/2014	99/99/9999	CAPECITABINE, 500 MG, ORAL	MORPHINE SULFATE (ISECURE SINGLE USE) 2 MG/ML	1	ML	SR	IV	ML	10	MG	0.2	01/06/2014	12/31/2014						
00409-1890-11		J2275		01/06/2014	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MORPHINE SULFATE (ISECURE SINGLE USE) 4 MG/ML	1	ML	SR	IV	ML	10	MG	0.4	01/13/2014	12/31/2014						
00409-1891-11		J2275		01/13/2014	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	CYTARABINE (SDV,PF,LATEX-FREE) 100 MG/ML	20	ML	VL	IJ	ML	100	MG	1	02/26/2014	99/99/9999						
67457-0452-20		J9100		02/26/2014	99/99/9999	INJECTION, CYTARABINE, 100 MG	MORPHINE SULFATE (SINGLE USE,PF) 2 MG/ML	1	ML	SR	IJ	ML	10	MG	0.2	04/01/2014	12/31/2014						
76045-0004-10		J2275		04/01/2014	12/31/2014	INJECTION, MORPHINE SULFATE (PRESERVATIVE-FREE STERILE SOLUTION), PER 10 MG	MYCOPHENOLIC ACID 180 MG	120	EA	BO	PO	EA	180	MG	1	03/11/2014	99/99/9999						
60505-2965-07		J7518		03/11/2014	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG																	

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00781-7157-86		J7644		09/11/2009	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	09/11/2009	99/99/9999							
00781-7157-86	KO	J7644	KO	09/11/2009	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (25X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	09/11/2009	99/99/9999							
59762-1001-01		J7520		01/16/2014	99/99/9999	SIROLIMUS, ORAL, 1 MG	SIROLIMUS 0.5 MG	100	EA	BO	PO	EA	1	MG	0.5	01/16/2014	99/99/9999							
68382-0520-01		J7520		01/09/2014	99/99/9999	SIROLIMUS, ORAL, 1 MG	SIROLIMUS (COATED) 0.5 MG	100	EA	BO	PO	EA	1	MG	0.5	01/09/2014	99/99/9999							
65862-0391-10		Q0162		03/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP 3X10) 8 MG	30	EA	BX	PO	EA	1	MG	8	03/01/2012	99/99/9999							
64380-0725-06		J7517		01/06/2014	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 MG	100	EA	BO	PO	EA	250	MG	2	01/06/2014	99/99/9999							
64380-0726-06		J7517		01/06/2014	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (HARD GELATIN) 250 MG	100	EA	BO	PO	EA	250	MG	1	01/06/2014	99/99/9999							
00944-2814-01		J0256		05/01/2014	99/99/9999	INJECTION, ALPHA 1 PROTEINASE INHIBITOR (HUMAN), NOT OTHERWISE SPECIFIED, 10 MG	ARALAST NP (500MG W/DILUENT) 1 MG	1	EA	VL	IV	EA	10	MG	0.1	05/01/2014	99/99/9999							
00944-2815-01		J0256		05/01/2014	99/99/9999	INJECTION, ALPHA 1 PROTEINASE INHIBITOR (HUMAN), NOT OTHERWISE SPECIFIED, 10 MG	ARALAST NP (1000MG W/DILUENT) 1 MG	1	EA	VL	IV	EA	10	MG	0.1	05/01/2014	99/99/9999							
25021-0301-67		J0150		05/01/2014	12/31/2014	INJECTION, ADENOSINE FOR THERAPEUTIC USE, 6 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS, INSTEAD USE A9270)	ADENOSINE (10X2ML,USP,PRF SYRINGE) 3 MG/ML	2	ML	SR	IV	ML	6	MG	0.5	05/01/2014	12/31/2014							
47335-0361-41		J0894		05/01/2014	99/99/9999	INJECTION, DECITABINE, 1 MG	DECITABINE (W/DILUENT,LYOPHILIZED) 50 MG	1	EA	VL	IV	EA	1	MG	50	05/01/2014	99/99/9999							
52118-0002-01		J3095		05/05/2014	11/30/2016	INJECTION, TELEVANCIN, 10 MG	VIBATIV (SDV,PF,LYOPHILIZED) 250 MG	10	EA	VL	IV	EA	10	MG	25	05/05/2014	11/30/2016							
67457-0424-10		J9060		05/23/2014	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (MDV) 1 MG/ML	100	ML	VL	IV	ML	10	MG	0.1	05/23/2014	99/99/9999							
67457-0425-51		J9060		05/23/2014	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN 1 MG/ML	50	ML	VL	IV	ML	10	MG	0.1	05/23/2014	99/99/9999							
47335-0890-21	None			02/13/2014	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 5 MG	14	EA	BO	PO	EA	5	MG	1	02/13/2014	99/99/9999							
47335-0890-80	None			02/13/2014	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 5 MG	5	EA	BO	PO	EA	5	MG	1	02/13/2014	99/99/9999							
47335-0891-21	None			02/13/2014	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 20 MG	14	EA	BO	PO	EA	20	MG	1	02/13/2014	99/99/9999							
47335-0891-80	None			02/13/2014	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 20 MG	5	EA	BO	PO	EA	20	MG	1	02/13/2014	99/99/9999							
47335-0892-21	None			02/13/2014	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 100 MG	14	EA	BO	PO	EA	100	MG	1	02/13/2014	99/99/9999							
47335-0892-80	None			02/13/2014	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 100 MG	5	EA	BO	PO	EA	100	MG	1	02/13/2014	99/99/9999							
47335-0893-80	None			02/13/2014	99/99/9999	TEMODAR, 250 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 250 MG	5	EA	BO	PO	EA	250	MG	1	02/13/2014	99/99/9999							
54569-1818-09	None			05/13/2008	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	36	EA	BO	PO	EA	2.5	MG	1	05/13/2008	99/99/9999							
47335-0929-21	None			02/13/2014	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 140 MG	14	EA	BO	PO	EA	20	MG	7	02/13/2014	99/99/9999							
47335-0929-80	None			02/13/2014	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 140 MG	5	EA	BO	PO	EA	20	MG	7	02/13/2014	99/99/9999							
47335-0930-21	None			02/13/2014	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 180 MG	14	EA	BO	PO	EA	20	MG	9	02/13/2014	99/99/9999							
47335-0930-80	None			02/13/2014	99/99/9999	TEMODAR, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 180 MG	5	EA	BO	PO	EA	20	MG	9	02/13/2014	99/99/9999							
51079-0508-20		J7518		02/12/2014	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (FILM-COATED) 180 MG	100	EA	BX	PO	EA	180	MG	1	02/12/2014	99/99/9999							
23155-0196-43		J2405		06/12/2014	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON 2 MG/ML	2	ML	VL	IJ	ML	1	MG	2	06/12/2014	99/99/9999							
25021-0230-02		J9206		07/01/2014	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X2ML SINGLE DOSE PF) 20 MG/ML	2	ML	VL	IV	ML	20	MG	1	07/01/2014	99/99/9999							
25021-0230-05		J9206		07/01/2014	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML SINGLE DOSE PF) 20 MG/ML	5	ML	VL	IV	ML	20	MG	1	07/01/2014	99/99/9999							
58468-0127-01		J1270		06/11/2014	99/99/9999	INJECTION, DOXERCALCIFEROL, 1 MCG	HECTOROL (50X2ML,MDV) 2 MCG/ML	2	ML	VL	IV	ML	1	MCG	2	06/11/2014	99/99/9999							
67457-0450-10		J9065		06/12/2014	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	CLADRIBINE (1X10ML,SDV,PF) 1 MG/ML	10	ML	VL	IV	ML	1	MG	1	06/12/2014	99/99/9999							
00574-0866-10		J7516		12/12/2012	99/99/9999	CYCLOSPORIN, PARENTERAL, 250 MG	CYCLOSPORINE 50 MG/ML	5	ML	AM	IV	ML	250	MG	0.2	12/12/2012	99/99/9999							
00703-2856-04		J3490		03/25/2013	01/08/2016	UNCLASSIFIED DRUGS	PROPOFOL (SDV,25X20ML) 10 MG/ML	20	ML	VL	IV	ML	1	EA	1	03/25/2013	01/08/2016							
00703-2858-09		J3490		01/02/2014	99/99/9999	UNCLASSIFIED DRUGS	PROPOFOL (SDV,20X50ML) 10 MG/ML	50	ML	VL	IV	ML	1	EA	1	01/02/2014	99/99/9999							
00703-2859-03		J3490		05/01/2013	05/24/2016	UNCLASSIFIED DRUGS	PROPOFOL (SDV,10X100ML) 10 MG/ML	100	ML	VL	IV	ML	1	EA	1	05/01/2013	05/24/2016							
00703-4502-04		J2765		12/20/2013	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HYDROCHLORIDE (S.D.V.) 5 MG/ML	2	ML	VL	IJ	ML	10	MG	0.5	12/20/2013	99/99/9999							
38779-0632-04		J7699		05/15/2014	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	25	GM	BO	NA	GM	1	MG	1	05/15/2014	99/99/9999							
38779-0632-05		J7699		05/15/2014	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	100	GM	BO	NA	GM	1	MG	1	05/15/2014	99/99/9999							
38779-0632-08		J7699		05/15/2014	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	500	GM	BO	NA	GM	1	MG	1	05/15/2014	99/99/9999							
38779-0632-09		J7699		05/15/2014	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	GENTAMICIN SULFATE (U.S.P.)	1000	GM	BO	NA	GM	1	MG	1	05/15/2014	99/99/9999							
50111-0788-67		Q0144		02/26/2014	02/03/2016	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (3X3,FILM-COATED) 500 MG	9	EA	BP	PO	EA	1000	MG	0.5	02/26/2014	02/03/2016							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54569-3701-00		J1050		01/01/2013	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	DEPO-PROVERA CONTRACEPTIVE (VIAL) 150 MG/ML	1	ML	VL	IM	ML	1	MG	150	01/01/2013	99/99/9999						
54569-4904-00		J1050		01/01/2013	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	DEPO-PROVERA CONTRACEPTIVE (SRN, PREFILLED) 150 MG/ML	1	ML	SR	IM	ML	1	MG	150	01/01/2013	99/99/9999						
62991-2002-01		J0278		10/31/2011	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (U.S.P.)	5	GM	BO	NA	GM	100	MG	10	10/31/2011	99/99/9999						
62991-2002-02		J0278		10/31/2011	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (U.S.P.)	25	GM	BO	NA	GM	100	MG	10	10/31/2011	99/99/9999						
65862-0390-10		Q0162		03/01/2012	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP 3X10) 4 MG	30	EA	BX	PO	EA	1	MG	4	03/01/2012	99/99/9999						
00574-0827-10		J1080		06/19/2014	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG	TESTOSTERONE CYPIONATE (USP, MDV) 200 MG/ML	10	ML	VL	IM	ML	200	MG	1	06/19/2014	12/31/2014						
00591-2738-23		J7614		07/01/2014	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	07/01/2014	99/99/9999						
00591-2738-23	KO	J7614	KO	07/01/2014	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 CC, 200 MG COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL HCL (24X3ML,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	07/01/2014	99/99/9999						
00781-7171-56		J7682		07/08/2014	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (PF) 300 MG/5 ML	5	ML	PC	IH	ML	300	MG	0.2	07/08/2014	99/99/9999						
00781-7171-56	KO	J7682	KO	07/08/2014	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (PF) 300 MG/5 ML	5	ML	PC	IH	ML	300	MG	0.2	07/08/2014	99/99/9999						
36000-0282-25		J1940		07/01/2014	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/ML	2	ML	VL	U	ML	20	MG	0.5	07/01/2014	99/99/9999						
36000-0283-25		J1940		07/01/2014	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/ML	4	ML	VL	U	ML	20	MG	0.5	07/01/2014	99/99/9999						
36000-0284-25		J1940		07/01/2014	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/ML	10	ML	VL	U	ML	20	MG	0.5	07/01/2014	99/99/9999						
42023-0129-01		J2680		07/09/2014	99/99/9999	INJECTION, FLUPHENAZINE DECANOATE, UP TO 25 MG	FLUPHENAZINE DECANOATE (LATEX-FREE) 25 MG/ML	5	ML	VL	U	ML	25	MG	1	07/09/2014	99/99/9999						
63739-0900-26		J1644		06/13/2014	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X10ML,LATEX-FREE) 1000 U/ML	2	ML	VL	U	ML	1000	U	1	06/13/2014	99/99/9999						
63739-0901-28		J1644		06/13/2014	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X10ML,LATEX-FREE) 5000 U/ML	10	ML	VL	U	ML	1000	U	5	06/13/2014	99/99/9999						
63739-0920-25		J1644		06/13/2014	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X1ML,LATEX-FREE) 1000 U/ML	1	ML	VL	U	ML	1000	U	1	06/13/2014	99/99/9999						
63739-0953-25		J1644		06/13/2014	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X10ML,LATEX-FREE) 5000 U/ML	1	ML	VL	U	ML	1000	U	5	06/13/2014	99/99/9999						
63739-0964-25		J1644		06/13/2014	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X1ML,LATEX-FREE) 10000 U/ML	1	ML	VL	U	ML	1000	U	10	06/13/2014	99/99/9999						
63739-0986-25		J1644		06/13/2014	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X1ML,LATEX-FREE) 20000 U/ML	1	ML	VL	U	ML	1000	U	20	06/13/2014	99/99/9999						
00093-4148-64		J7614		04/29/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X4,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	04/29/2013	99/99/9999						
00093-4148-64	KO	J7614	KO	04/29/2013	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X4,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	04/29/2013	99/99/9999						
00085-1136-02		J1327		08/18/2014	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	INTEGRILIN 0.75 MG/ML	100	ML	VL	IV	ML	5	MG	0.15	08/18/2014	99/99/9999						
67457-0263-30		J1205		08/04/2014	99/99/9999	INJECTION, CHLOROTHIAZIDE SODIUM, PER 500 MG	CHLOROTHIAZIDE SODIUM (USP, SDV,LYOPHILIZED) 0.5 GM	1	EA	VL	IV	EA	500	MG	1	08/04/2014	99/99/9999						
67457-0434-51		J9265		08/07/2014	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (MDV) 6 MG/ML	50	ML	VL	IV	ML	30	MG	0.2	08/07/2014	12/31/2014						
67457-0449-17		J9265		08/07/2014	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (MDV) 6 MG/ML	16.7	ML	VL	IV	ML	30	MG	0.2	08/07/2014	12/31/2014						
67457-0471-52		J9265		08/07/2014	12/31/2014	INJECTION, PACLITAXEL, 30 MG	PACLITAXEL (MDV) 6 MG/ML	5	ML	VL	IV	ML	30	MG	0.2	08/07/2014	12/31/2014						
00074-2108-03		J1950		08/03/2009	99/99/9999	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT-PED (LYOPHILIZED) 7.5 MG	1	EA	BX	IM	EA	3.75	MG	2	08/03/2009	99/99/9999						
00074-2282-03		J1950		04/03/2009	99/99/9999	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT-PED (LYOPHILIZED) 11.25 MG	1	EA	BX	IM	EA	3.75	MG	3	04/03/2009	99/99/9999						
00074-2440-03		J1950		04/17/2009	99/99/9999	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT-PED (LYOPHILIZED) 15 MG	1	EA	BX	IM	EA	3.75	MG	4	04/17/2009	99/99/9999						
00074-3346-03		J9217		04/02/2009	99/99/9999	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	LUPRON DEPOT (STERILE,1X22.5MG) 22.5 MG	1	EA	BX	IM	EA	7.5	MG	3	04/02/2009	99/99/9999						
00074-3473-03		J9217		06/17/2011	99/99/9999	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	LUPRON DEPOT (LYOPHILIZED) 45 MG	1	EA	BX	IM	EA	7.5	MG	6	06/17/2011	99/99/9999						
00074-3641-03		J1950		04/13/2009	99/99/9999	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT 3.75 MG	1	EA	BX	IM	EA	3.75	MG	1	04/13/2009	99/99/9999						
00074-3642-03		J9217		03/25/2009	99/99/9999	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	LUPRON DEPOT (STERILE,1X7.5MG) 7.5 MG	1	EA	BX	IM	EA	7.5	MG	1	03/25/2009	99/99/9999						
00074-3663-03		J1950		05/21/2009	99/99/9999	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT (DUAL-CHAMBER SYRINGE) 11.25 MG	1	EA	BX	IM	EA	3.75	MG	3	05/21/2009	99/99/9999						
00074-3683-03		J9217		04/17/2009	99/99/9999	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	LUPRON DEPOT (LYOPHILIZED) 30 MG	1	EA	BX	IM	EA	7.5	MG	4	04/17/2009	99/99/9999						
00074-3779-03		J1950		08/15/2011	99/99/9999	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT-PED (SINGLE DOSE) 11.25 MG	1	EA	BX	IM	EA	3.75	MG	3	08/15/2011	99/99/9999						
00074-9694-03		J1950		08/15/2011	99/99/9999	INJECTION, LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), PER 3.75 MG	LUPRON DEPOT-PED (SINGLE DOSE) 30 MG	1	EA	BX	IM	EA	3.75	MG	8	08/15/2011	99/99/9999						
16714-0671-01		Q0162		10/15/2009	10/31/2016	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,1X60ML,STRAWBERRY) 4 MG/5ML	60	ML	BO	PO	ML	1	MG	0.8	10/15/2009	10/31/2016						
51079-0620-06		J7500		07/23/2010	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE (5X10,USP) 50 MG	50	EA	BX	PO	EA	50	MG	1	07/23/2010	99/99/9999						
00378-2511-91	None			08/08/2014	99/99/9999	CAPECITABINE, 150 MG	CAPECITABINE (USP,FILM COATED) 150 MG	60	EA	BO	PO	EA	150	MG	1	08/08/2014	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00378-2512-78		None		08/08/2014	99/99/9999	CAPECITABINE, 500 MG	CAPECITABINE (USP,FILM COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	08/08/2014	99/99/9999							
17478-0340-38		J7682		09/11/2014	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5	ML	PC	IH	ML	300	MG	0.2	09/11/2014	99/99/9999							
17478-0340-38	KO	J7682	KO	09/11/2014	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5	ML	PC	IH	ML	300	MG	0.2	09/11/2014	99/99/9999							
25021-0700-01		J1885		09/01/2014	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV,25X1ML,PF) 15 MG/ML	1	ML	VL	IJ	ML	15	MG	1	09/01/2014	99/99/9999							
25021-0701-01		J1885		09/01/2014	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV,25X1ML,PF) 30 MG/ML	1	ML	VL	IJ	ML	15	MG	2	09/01/2014	99/99/9999							
25021-0701-02		J1885		09/01/2014	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV,25X2ML,PF) 30 MG/ML	2	ML	VL	IM	ML	15	MG	2	09/01/2014	99/99/9999							
25021-0827-61		J1740		09/02/2014	99/99/9999	INJECTION, IBANDRONATE SODIUM, 1 MG	IBANDRONATE SODIUM (PREFILLED, SINGLE-USE) 1 MG/ML	3	ML	SR	IV	ML	1	MG	1	09/02/2014	99/99/9999							
64380-0720-06		J7507		09/10/2014	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS 0.5 MG	100	EA	BO	PO	EA	1	MG	0.5	09/10/2014	99/99/9999							
64380-0721-06		J7507		09/10/2014	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS 1 MG	100	EA	BO	PO	EA	1	MG	1	09/10/2014	99/99/9999							
64380-0722-06		J7507		09/10/2014	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS 5 MG	100	EA	BO	PO	EA	1	MG	5	09/10/2014	99/99/9999							
67457-0429-20		J9208		09/04/2014	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (1X20ML) 1 GM/20 ML	20	ML	VL	IV	ML	1	GM	0.05	09/04/2014	99/99/9999							
67457-0474-04		J9351		09/04/2014	99/99/9999	INJECTION, TOPOTECAN, 0.1 MG	TOPOTECAN HYDROCHLORIDE (SINGLE-DOSE,LYOPHILIZED) 4 MG	1	EA	VL	IV	EA	0.1	MG	40	09/04/2014	99/99/9999							
67457-0476-10		J9263		09/04/2014	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF,LYOPHILIZED) 100 MG	1	EA	VL	IV	EA	0.5	MG	200	09/04/2014	99/99/9999							
67457-0524-33		J1740		09/02/2014	99/99/9999	INJECTION, IBANDRONATE SODIUM, 1 MG	IBANDRONATE SODIUM 1 MG/ML	5	ML	SR	IV	ML	1	MG	1	09/02/2014	99/99/9999							
68982-0850-01		J1568		09/05/2014	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM 10% (PF,LATEX-FREE) 100 MG/ML	20	ML	VL	IV	ML	500	MG	0.2	09/05/2014	99/99/9999							
68982-0850-02		J1568		09/05/2014	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM 10% (PF,LATEX-FREE) 100 MG/ML	50	ML	VL	IV	ML	500	MG	0.2	09/05/2014	99/99/9999							
68982-0850-03		J1568		09/05/2014	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM 10% (PF,LATEX-FREE) 100 MG/ML	100	ML	VL	IV	ML	500	MG	0.2	09/05/2014	99/99/9999							
68982-0850-04		J1568		09/05/2014	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM 10% (PF,LATEX-FREE) 100 MG/ML	200	ML	VL	IV	ML	500	MG	0.2	09/05/2014	99/99/9999							
00054-0383-25		None		06/23/2014	99/99/9999	CYCLOPHOSPHAMIDE; ORAL, 50 MG	CYCLOPHOSPHAMIDE 50 MG	100	EA	BO	PO	EA	50	MG	1	06/23/2014	99/99/9999							
00054-0382-25		None		06/23/2014	99/99/9999	CYCLOPHOSPHAMIDE; ORAL, 25 MG	CYCLOPHOSPHAMIDE 25 MG	100	EA	BO	PO	EA	25	MG	1	06/23/2014	99/99/9999							
50383-0810-16		J8499		09/04/2014	09/01/2017	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR (BANANA) 200 MG/5 ML	473	ML	BO	PO	ML	1	MG	40	09/04/2014	09/01/2017							
13533-0661-06		J2788		11/01/2013	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, MINIDOSE, 50 MICROGRAMS (250 I.U.)	HYPERRHO S/D (MINI-DOSE,SD,PF)	10	EA	SR	IM	EA	50	MCG	1	11/01/2013	99/99/9999							
00051-0022-21		Q0167		01/01/2014	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFT GELATIN) 5 MG	60	EA	BO	PO	EA	2.5	MG	2	01/01/2014	99/99/9999							
00051-0023-21		Q0167		01/01/2014	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	MARINOL (SOFTGEL) 10 MG	60	EA	BO	PO	EA	2.5	MG	4	01/01/2014	99/99/9999							
00074-3799-03		J0135		10/01/2014	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PEDIATRIC,PF) 40 MG/0.8 ML	3	EA	BX	MR	EA	20	MG	2	10/01/2014	99/99/9999							
00074-3799-06		J0135		10/01/2014	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PEDIATRIC,PF) 40 MG/0.8 ML	6	EA	BX	MR	EA	20	MG	2	10/01/2014	99/99/9999							
62935-0302-30		J9217		10/02/2014	05/06/2015	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	ELIGARD (SINGLE-USE) 30 MG	1	EA	BX	SC	EA	7.5	MG	4	10/02/2014	05/06/2015							
67457-0443-60		J9208		10/07/2014	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (1X60ML) 3 GM/60 ML	60	ML	VL	IV	ML	1	GM	0.05	10/07/2014	99/99/9999							
76045-0001-20		J2250		10/01/2014	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (PREFILLED, USP,PF) 1 MG/ML	2	ML	SR	IJ	ML	1	MG	1	10/01/2014	99/99/9999							
76045-0002-10		J2250		10/01/2014	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (PF) 5 MG/ML	1	ML	SR	IJ	ML	1	MG	5	10/01/2014	99/99/9999							
76045-0003-20		J2250		10/01/2014	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (PF) 5 MG/ML	2	ML	SR	IJ	ML	1	MG	5	10/01/2014	99/99/9999							
16729-0042-01		J7507		09/30/2011	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	BO	PO	EA	1	MG	1	09/30/2011	99/99/9999							
54505-0101-01		J0171		11/13/2014	10/03/2015	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE 0.15 MG/0.15 ML	1	EA	SR	IJ	EA	0.1	MG	1.5	11/13/2014	10/03/2015							
63323-0285-61		J2795		11/03/2014	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	NAROPIN (IN FREEFLEX BAG,PF) 2 MG/ML	100	ML	BG	IJ	ML	1	MG	2	11/03/2014	99/99/9999							
63323-0285-63		J2795		11/03/2014	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	NAROPIN (IN FREEFLEX BAG,PF) 2 MG/ML	200	ML	BG	IJ	ML	1	MG	2	11/03/2014	99/99/9999							
67457-0396-10		J9000		11/07/2014	99/99/9999	INJECTION, VINORELBINE HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP-STERILE,MDV) 2 MG/ML	100	ML	VL	IV	ML	10	MG	0.2	11/07/2014	99/99/9999							
67457-0431-11		J9390		11/07/2014	08/31/2016	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE (S.D.V., 1X1ML) 10 MG/ML	1	ML	VL	IV	ML	10	MG	1	11/07/2014	08/31/2016							
62935-0752-75		J9217		09/25/2014	05/06/2015	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	ELIGARD (SINGLE-USE) 7.5 MG	1	EA	BX	SC	EA	7.5	MG	1	09/25/2014	05/06/2015							
00009-0347-02		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	DEPO-TESTOSTERONE (VIAL) 100 MG/ML	10	ML	VL	IM	ML	1	MG	100	01/01/2015	99/99/9999							
00009-0417-01		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML	1	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999							
00009-0417-02		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML	10	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999							
00172-3753-96		J9267		01/01/2015	02/10/2016	INJECTION, PACLITAXEL, 1 MG	NOV-ONXOL (M.D.V.) 6 MG/ML	50	ML	VL	IV	ML	1	MG	6	01/01/2015	02/10/2016							
00172-3754-94		J9267		01/01/2015	02/10/2016	INJECTION, PACLITAXEL, 1 MG	NOV-ONXOL (M.D.V.) 6 MG/ML	5	ML	VL	IV	ML	1	MG	6	01/01/2015	02/10/2016							
00172-3756-95		J9267		01/01/2015	02/10/2016	INJECTION, PACLITAXEL, 1 MG	NOV-ONXOL (M.D.V.) 6 MG/ML	25	ML	VL	IV	ML	1	MG	6	01/01/2015	02/10/2016							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00406-1521-53		J2270		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE	5	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999						
00406-1521-55		J2270		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE	25	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999						
00406-1521-56		J2270		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE	50	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999						
00406-1521-57		J2270		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE	100	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999						
00409-1134-03		J2270		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (VIAL, FLIPTOP) 50 MG/ML	20	ML	VL	IJ	ML	10	MG	5	01/01/2015	99/99/9999						
00409-1134-05		J2270		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (LATEX-FREE) 50 MG/ML	50	ML	VL	IJ	ML	10	MG	5	01/01/2015	99/99/9999						
00409-1135-02		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (HIGH CONCENTRATION,PF) 25 MG/ML	10	ML	VL	IJ	ML	10	MG	2.5	01/01/2015	99/99/9999						
00409-1890-01		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 2 MG/ML	1	ML	SR	IV	ML	10	MG	0.2	01/01/2015	99/99/9999						
00409-1890-11		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (ISECURE SINGLE USE) 2 MG/ML	1	ML	SR	IV	ML	10	MG	0.2	01/01/2015	99/99/9999						
00409-1891-01		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 4 MG/ML	1	ML	SR	IV	ML	10	MG	0.4	01/01/2015	99/99/9999						
63323-0371-10		J0878		04/11/2018	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (PF,LYOPHILIZED) 500 MG	1	EA	VL	IV	EA	1	MG	500	04/11/2018	99/99/9999						
00409-1891-11		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (ISECURE SINGLE USE) 4 MG/ML	1	ML	SR	IV	ML	10	MG	0.4	01/01/2015	99/99/9999						
00409-1893-01		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 10 MG/ML	1	ML	SR	IV	ML	10	MG	1	01/01/2015	99/99/9999						
00409-1894-01		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (CARPUJECT SINGLE-USE) 15 MG/ML	1	ML	SR	IV	ML	10	MG	1.5	01/01/2015	99/99/9999						
00409-3814-12		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (5X10ML,PF,LATEX-FREE) 0.5 MG/ML	10	ML	VL	IJ	ML	10	MG	0.05	01/01/2015	99/99/9999						
00409-4057-12		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (PF,LATEX-FREE) 0.5 MG/ML	10	ML	AM	IJ	ML	10	MG	0.05	01/01/2015	99/99/9999						
00409-6028-04		J2270		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (SDV,30MLX10,PF) 5 MG/ML	30	ML	VL	IV	ML	10	MG	0.5	01/01/2015	99/99/9999						
00463-1069-10		J3490		01/01/2015	07/23/2015	UNCLASSIFIED DRUGS	TESTRO AQ (VIAL) 100 MG/ML	10	ML	VL	IM	ML	1	EA	1	01/01/2015	07/23/2015						
00463-1073-10		J3490		01/01/2015	02/03/2016	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (VIAL) 100 MG/ML	10	ML	VL	IM	ML	1	EA	1	01/01/2015	02/03/2016						
00469-8234-12		J0153		01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOCARD (ANSYR,LUER LOK) 3 MG/ML	2	ML	SR	IV	ML	1	MG	3	01/01/2015	99/99/9999						
00469-8234-14		J0153		01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOCARD (ANSYR,LUER LOK) 3 MG/ML	4	ML	SR	IV	ML	1	MG	3	01/01/2015	99/99/9999						
00574-0820-01		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (1X1ML,USP) 200 MG/ML	1	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999						
00574-0827-10		J1071		01/01/2015	08/31/2017	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (USP, MDV) 200 MG/ML	10	ML	VL	IM	ML	1	MG	200	01/01/2015	08/31/2017						
00591-3221-26		J3121		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE ENANTHATE, 1 MG	TESTOSTERONE ENANTHATE 200 MG/ML	5	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999						
00591-3223-79		J1071		01/01/2015	03/04/2015	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (M.D.V.) 200 MG/ML	10	ML	VL	IM	ML	1	MG	200	01/01/2015	03/04/2015						
00641-6019-10		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	DURAMORPH (10X10ML,PF) 1 MG/ML	10	ML	AM	IJ	ML	10	MG	0.1	01/01/2015	99/99/9999						
00641-6020-10		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	DURAMORPH (10X10ML,PF) 0.5 MG/ML	10	ML	AM	IJ	ML	10	MG	0.05	01/01/2015	99/99/9999						
00641-6039-01		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	INFUMORPH 200 (1X20ML,PF) 10 MG/ML	20	ML	AM	IJ	ML	10	MG	1	01/01/2015	99/99/9999						
00641-6040-01		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	INFUMORPH 500 (1X20ML,PF) 25 MG/ML	20	ML	AM	IJ	ML	10	MG	2.5	01/01/2015	99/99/9999						
00641-6071-25		J2270		01/01/2015	02/28/2017	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE, (S.D.V., 1MLX25) 15MG/ML	1	ML	VL	IJ	ML	10	MG	1.5	01/01/2015	02/28/2017						
00641-6072-01		J2270		01/01/2015	09/16/2015	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (M.D.V.) 15MG/ML	20	ML	VL	IJ	ML	10	MG	1.5	01/01/2015	09/16/2015						
10019-0179-36		J2270		01/01/2015	10/17/2016	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (MDV) 15 MG/ML	20	ML	NA	IJ	ML	10	MG	1.5	01/01/2015	10/17/2016						
25021-0301-67		J0153		01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (10X2ML,USP,PRF SYRINGE) 3 MG/ML	2	ML	SR	IV	ML	1	MG	3	01/01/2015	99/99/9999						
35356-0058-10		J1071		01/01/2015	01/01/2015	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	DEPO-TESTOSTERONE 100 MG/ML	10	ML	VL	IM	ML	1	MG	100	01/01/2015	01/01/2015						
38779-0164-03		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.)	5	GM	BO	NA	GM	1	MG	1000	01/01/2015	99/99/9999						
38779-0164-04		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.)	25	GM	BO	NA	GM	1	MG	1000	01/01/2015	99/99/9999						
38779-0164-05		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE (U.S.P.)	100	GM	BO	NA	GM	1	MG	1000	01/01/2015	99/99/9999						
38779-0164-08		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.)	500	GM	BO	NA	GM	1	MG	1000	01/01/2015	99/99/9999						
38779-0164-09		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1000	GM	JR	NA	GM	1	MG	1000	01/01/2015	99/99/9999						
38779-0165-03		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (USP,MICRONIZED)	5	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999						
38779-0165-04		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (USP,MICRONIZED)	25	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999						
38779-0165-05		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	100	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999						
38779-0165-08		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	500	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
38779-0673-03	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	5	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999							
38779-0673-04	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	25	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999							
38779-0673-05	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	100	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999							
38779-0673-07	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	250	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999							
38779-0855-04	J3121			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE ENANTHATE, 1 MG	TESTOSTERONE ENANTHATE	25	GM	NA	NA	GM	1	MG	1000	01/01/2015	99/99/9999							
51552-0029-01	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.)	1	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0029-02	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED (U.S.P.)	5	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0030-01	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.)	1	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0030-02	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.)	5	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0030-04	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.)	25	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0030-05	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.)	100	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0030-08	J3490			01/01/2015	01/01/2015	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P., MICRONIZED)	0.3	GM	BO	NA	GM	1	EA	1	01/01/2015	01/01/2015							
51552-0030-09	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P., MICRONIZED)	0.6	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0564-04	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.)	25	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0564-05	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P., MICRONIZED)	100	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0564-07	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.)	1000	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51552-0678-02	J2270			01/01/2015	01/01/2015	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (1X5GM USP)	5	GM	NA	NA	GM	10	MG	100	01/01/2015	01/01/2015							
51552-0678-04	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (1X25GM USP)	25	GM	JR	NA	GM	10	MG	100	01/01/2015	99/99/9999							
51552-0678-06	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (1X100GM USP)	100	GM	JR	NA	GM	10	MG	100	01/01/2015	99/99/9999							
51927-1000-00	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.; CII)	1	GM	JR	NA	GM	10	MG	100	01/01/2015	99/99/9999							
51927-1026-00	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (USP; NON MICRONIZED; SOY)	1	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51927-1027-00	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED (U.S.P.; SOY; CII)	1	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51927-1029-00	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE MICRONIZED (MICRONIZED, CII)	1	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999							
51927-2706-00	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.; CIII)	1	GM	JR	NA	GM	1	MG	1000	01/01/2015	99/99/9999							
54569-5610-00	J0153			01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (PF) 3 MG/ML	2	ML	NA	IV	ML	1	MG	3	01/01/2015	99/99/9999							
54868-0216-00	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML	10	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999							
54868-0796-00	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1MG	DEPO-TESTOSTERONE 100 MG/ML	10	ML	VL	IM	ML	100	MG	100	01/01/2015	99/99/9999							
54868-3618-01	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE 200 MG/ML	1	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999							
54868-4050-00	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE	25	GM	JR	NA	GM	10	MG	100	01/01/2015	99/99/9999							
54868-5016-00	J3121			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE ENANTHATE, 1 MG	DELATESTRYL 200 MG/ML	5	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999							
54868-5551-00	J0153			01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE 3 MG/ML	2	ML	VL	IV	ML	6	MG	3	01/01/2015	99/99/9999							
55390-0067-10	J0153			01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (S.D.V.,PF) 3 MG/ML	2	ML	VL	IV	ML	1	MG	3	01/01/2015	99/99/9999							
63323-0942-05	J2469			03/27/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL (LATEX-FREE) 0.05 MG/1 ML	5	ML	VL	IV	ML	25	MCG	2	03/27/2018	99/99/9999							
60977-0016-73	J2274			01/01/2015	02/28/2015	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	DURAMORPH (PF) 0.5 MG/ML	10	ML	AM	IJ	ML	10	MG	0.05	01/01/2015	02/28/2015							
60977-0114-74	J2274			01/01/2015	02/03/2016	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	INFUMORPH 200 (PF) 10 MG/ML	1	ML	NA	IJ	ML	10	MG	1	01/01/2015	02/03/2016							
60977-0115-74	J2274			01/01/2015	02/03/2016	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	INFUMORPH 500 (PF) 25 MG/ML	1	ML	NA	IJ	ML	10	MG	2.5	01/01/2015	02/03/2016							
61553-0649-75	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (5X50ML,LATEX-FREE) 50 MG/ML	50	ML	EA	IJ	ML	10	MG	5	01/01/2015	99/99/9999							
61553-0651-76	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE-SODIUM CHLORIDE (5X55ML,LATEX-FREE) 1 MG/ML-0.9%	55	ML	EA	IJ	ML	10	MG	0.1	01/01/2015	99/99/9999							
61703-0342-09	J9267			01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (M.D.V.) 6 MG/ML	5	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999							
61703-0342-22	J9267			01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (M.D.V.) 6 MG/ML	16.7	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999							
61703-0342-50	J9267			01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (M.D.V.) 6 MG/ML	50	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999							
62991-1707-01	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.)	5	GM	BO	NA	GM	1	MG	1000	01/01/2015	99/99/9999							
62991-1707-02	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE (U.S.P.)	25	GM	BO	NA	GM	1	MG	1000	01/01/2015	99/99/9999							
62991-1707-03	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.)	100	GM	BO	NA	GM	1	MG	1000	01/01/2015	99/99/9999							
62991-2150-01	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED (U.S.P.)	5	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
62991-2150-02	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED (U.S.P.)	25	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
62991-2150-03	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED (U.S.P.)	100	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
62991-2150-04	J3490			01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED (U.S.P.)	500	GM	BO	NA	GM	1	EA	1	01/01/2015	99/99/9999							
63275-1025-04	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	25	GM	BO	NA	GM	10	MG	100	01/01/2015	99/99/9999							
63275-1100-05	J2270			01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	100	GM	BO	NA	GM	10											

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
63275-9983-04		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED	25	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999						
63275-9983-05		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED	100	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999						
63275-9983-08		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED	500	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999						
63275-9983-09		J3490		01/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE MICRONIZED	1000	GM	JR	NA	GM	1	EA	1	01/01/2015	99/99/9999						
63323-0651-02		J0153		01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (PF) 3 MG/ML	2	ML	VL	IV	ML	1	MG	3	01/01/2015	99/99/9999						
63323-0651-04		J0153		01/01/2015	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (PF) 3 MG/ML	4	ML	VL	IV	ML	1	MG	3	01/01/2015	99/99/9999						
66758-0043-01		J9267		01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (USP,1X5ML,MULTI-DOSE) 6 MG/ML	5	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999						
66758-0043-02		J9267		01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (USP,1X16.7ML,MULTI-DOSE) 6 MG/ML	16.7	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999						
66758-0043-03		J9267		01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (USP,1X50ML,MULTI-DOSE) 6 MG/ML	50	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999						
67457-0434-51		J9267		01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV) 6 MG/ML	50	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999						
67457-0448-17		J9267		01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV) 6 MG/ML	16.7	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999						
67457-0471-52		J9267		01/01/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV) 6 MG/ML	5	ML	VL	IV	ML	1	MG	6	01/01/2015	99/99/9999						
76045-0004-10		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (SINGLE USE,PF) 2 MG/ML	1	ML	SR	U	ML	10	MG	0.2	01/01/2015	99/99/9999						
51079-0510-01	None			08/25/2014	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE,(USP,FILM COATED) 500MG	1	EA	BP	PO	EA	500	MG	1	08/25/2014	99/99/9999						
51079-0510-05	None			08/25/2014	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE,(USP,FILM COATED) 500MG	20	EA	BX	PO	EA	500	MG	1	08/25/2014	99/99/9999						
54569-1411-00		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML TESTOSTERONE CYPIONATE (M.D.V.) 200 MG/ML	10	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999						
54868-3818-00		J1071		01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	DEPO-TESTOSTERONE (VIAL) 200 MG/ML TESTOSTERONE CYPIONATE (M.D.V.) 200 MG/ML	10	ML	VL	IM	ML	1	MG	200	01/01/2015	99/99/9999						
55111-0653-01		J7520		10/27/2014	99/99/9999	SIROLIMUS, ORAL, 1 MG	SIROLIMUS 1 MG	100	EA	BO	PO	EA	1	MG	1	10/27/2014	99/99/9999						
00703-8510-21		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 150 MG/ML	1	ML	SR	U	ML	10	MG	15	11/19/2014	99/99/9999						
00703-8510-23		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 150 MG/ML	1	ML	SR	U	ML	10	MG	15	11/19/2014	99/99/9999						
00703-8530-21		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 30 MG/0.3 ML	0.3	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8530-23		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 30 MG/0.3 ML	0.3	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8540-21		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 40 MG/0.4 ML	0.4	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8540-23		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 40 MG/0.4 ML	0.4	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8560-21		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 60 MG/0.6 ML	0.6	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8560-23		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 60 MG/0.6 ML	0.6	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8580-21		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 100 MG/ML	1	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8580-23		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 100 MG/ML	1	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8610-21		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 120 MG/0.8 ML	0.8	ML	SR	U	ML	10	MG	15	11/19/2014	99/99/9999						
00703-8610-23		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 120 MG/0.8 ML	0.8	ML	SR	U	ML	10	MG	15	11/19/2014	99/99/9999						
00703-8680-21		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 80 MG/0.8 ML	0.8	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
00703-8680-23		J1650		11/19/2014	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 80 MG/0.8 ML	0.8	ML	SR	U	ML	10	MG	10	11/19/2014	99/99/9999						
47335-0150-40		J9045		11/17/2014	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/ML	5	ML	VL	IV	ML	50	MG	0.2	11/17/2014	99/99/9999						
47335-0151-40		J9045		11/17/2014	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/ML	15	ML	VL	IV	ML	50	MG	0.2	11/17/2014	99/99/9999						
47335-0284-40		J9045		11/17/2014	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/ML	60	ML	VL	IV	ML	50	MG	0.2	11/17/2014	99/99/9999						
47335-0300-40		J9045		11/17/2014	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/ML	45	ML	VL	IV	ML	50	MG	0.2	11/17/2014	99/99/9999						
67457-0479-53		J9390		09/04/2014	08/31/2016	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE (S.D.V.) 10 MG/ML	5	ML	VL	IV	ML	10	MG	1	09/04/2014	08/31/2016						
63459-0177-14		J9262		11/12/2012	99/99/9999	INJECTION, OMACETAXINE MEPEUSUCCINATE, 0.01 MG	SYNRIBO (PF,LYOPHILIZED) 3.5MG	1	EA	VL	SC	EA	0.01	MG	350	11/12/2012	99/99/9999						
59762-1002-01		J7520		10/27/2014	99/99/9999	SIROLIMUS, ORAL, 1 MG	SIROLIMUS 1 MG	100	EA	BO	PO	EA	1	MG	1	10/27/2014	99/99/9999						
59762-1003-01		J7520		10/27/2014	99/99/9999	SIROLIMUS, ORAL, 1 MG	SIROLIMUS 2 MG	100	EA	BO	PO	EA	1	MG	2	10/27/2014	99/99/9999						
38779-0310-09		J2675		09/26/2008	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (MILLED, U.S.P.)	1000	GM	BO	NA	GM	50	MG	20	09/26/2008	99/99/9999						
00024-5860-01		J9027		12/15/2014	99/99/9999	INJECTION, CLOFARABINE, 1 MG	CLOLAR (SINGLE-USE VIAL,PF) 1 MG/ML	20	ML	VL	IV	ML	1	MG	1	12/15/2014	99/99/9999						
00093-4147-19		J7614		12/11/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (INNER PACK,PF) 1.25 MG/0.5 ML	1	EA	PC	IH	EA	0.5	MG	2.5	12/11/2014	99/99/9999						
00093-4147-19	KO	J7614	KO	12/11/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (INNER PACK,PF) 1.25 MG/0.5 ML	1	EA	PC	IH	EA	0.5	MG	2.5	12/11/2014	99/99/9999						
00093-4147-56		J7614		12/11/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (USP,PF) 1.25 MG/0.5 ML	30	EA	PC	IH	EA	0.5	MG	2.5	12/11/2014	99/99/9999						
00093-4147-56	KO	J7614	KO	12/11/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (USP,PF) 1.25 MG/0.5 ML SOMATULINE DEPOT (1X0.2ML, SINGLE USE) 60 MG/0.2 ML	30	EA	PC	IH	EA	0.5	MG	2.5	12/11/2014	99/99/9999						
15054-1060-03		J1930		01/02/2015	99/99/9999	INJECTION, LANREOTIDE, 1 MG	LANREOTIDE (SINGLE USE) 1 MG/ML	0.2	ML	SR	SC	ML	1	MG	300	01/02/2015	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
15054-1090-03		J1930		01/02/2015	99/99/9999	INJECTION, LANREOTIDE, 1 MG	SOMATULINE DEPOT (1X0.3ML, SINGLE USE) 90 MG/0.3 ML	0.3	ML	SR	SC	ML	1	MG	300	01/02/2015	99/99/9999							
15054-1120-03		J1930		01/02/2015	99/99/9999	INJECTION, LANREOTIDE, 1 MG	SOMATULINE DEPOT (1X0.5ML, SINGLE USE) 120 MG/0.5 ML	0.5	ML	SR	SC	ML	1	MG	240	01/02/2015	99/99/9999							
24987-0362-10		J2780		12/01/2014	01/10/2017	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC 25 MG/ML	2	ML	VL	IJ	ML	25	MG	1	12/01/2014	01/10/2017							
25021-0159-10		J0770		12/15/2014	99/99/9999	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE (USP,LYOPHILIZED) 150 MG	1	EA	VL	IJ	EA	150	MG	1	12/15/2014	99/99/9999							
25021-0234-10		J9201		01/01/2015	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE HCL (SDV,USP,PF,LYOPHILIZED) 200 MG	1	EA	VL	IV	EA	200	MG	1	01/01/2015	99/99/9999							
25021-0235-50		J9201		01/01/2015	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE HCL (SDV,USP,PF,LYOPHILIZED) 1 GM	1	EA	VL	IV	EA	200	MG	5	01/01/2015	99/99/9999							
52118-0001-01		J3095		01/02/2015	09/30/2016	INJECTION, TELEVANCIN, 10 MG	VIBATIV (SDV,PF,LYOPHILIZED) 750 MG	10	EA	VL	IV	EA	10	MG	75	01/02/2015	09/30/2016							
63323-0404-00		J0290		12/12/2014	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (BULK PACKAGE,LATEX-FREE) 10 GM	1	EA	VL	IV	EA	500	MG	20	12/12/2014	99/99/9999							
67457-0273-10		J2800		12/05/2014	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (25X10ML, SDV) 100 MG/ML	10	ML	VL	IJ	ML	10	ML	0.1	12/05/2014	99/99/9999							
67457-0395-25		J9000		12/16/2014	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,STERILE,SDV) 2 MG/ML	25	ML	VL	IV	ML	10	MG	0.2	12/16/2014	99/99/9999							
00591-2737-23		J7614		08/07/2014	99/99/9999	COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL HCL (24X3ML,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	08/07/2014	99/99/9999							
00591-2737-23	KO	J7614	KO	08/07/2014	99/99/9999	COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL HCL (24X3ML,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	08/07/2014	99/99/9999							
60505-2966-07		J7518		08/20/2014	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (ENTERIC COATED) 360 MG	120	EA	BO	PO	EA	180	MG	2	08/20/2014	99/99/9999							
00409-1008-01		J2501		11/01/2014	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL 0.005 MG/ML	1	ML	VL	IV	ML	1	MCG	5	11/01/2014	99/99/9999							
00409-1008-02		J2501		11/01/2014	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL 0.005 MG/ML	2	ML	VL	IV	ML	1	MCG	5	11/01/2014	99/99/9999							
12496-0757-05		J0592		01/19/2015	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENEX 0.3 MG/ML	1	ML	AM	IJ	ML	0.1	MG	3	01/19/2015	99/99/9999							
23155-0473-41		J1940		12/08/2014	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/ML	2	ML	VL	IJ	ML	20	MG	0.5	12/08/2014	99/99/9999							
23155-0473-42		J1940		12/08/2014	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/ML	4	ML	VL	IJ	ML	20	MG	0.5	12/08/2014	99/99/9999							
23155-0473-44		J1940		12/08/2014	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/ML	10	ML	VL	IJ	ML	20	MG	0.5	12/08/2014	99/99/9999							
25021-0236-04		J9351		01/01/2015	99/99/9999	INJECTION, TOPOTECAN, 0.1 MG	TOPOTECAN HCL (1X4ML,PF) 1 MG/ML	4	ML	VL	IV	ML	0.1	MG	10	01/01/2015	99/99/9999							
45963-0608-60		J9178		01/13/2015	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HCL (SDV,PF) 2 MG/ML	100	ML	VL	IV	ML	2	MG	1	01/13/2015	99/99/9999							
45963-0608-68		J9178		02/02/2015	99/99/9999	INJECTION, EPIRUBICIN HCL, 2 MG	EPIRUBICIN HCL (SDV,PF) 2 MG/ML	25	ML	VL	IV	ML	2	MG	1	02/02/2015	99/99/9999							
45963-0609-55		J9185		01/13/2015	99/99/9999	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE (USP,SDV,PF,LYOPHILIZED) 50 MG	1	EA	VL	IV	EA	50	MG	1	01/13/2015	99/99/9999							
45963-0611-53		J9263		01/13/2015	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (SDV,PF,LYOPHILIZED) 50 MG	1	EA	VL	IV	EA	0.5	MG	100	01/13/2015	99/99/9999							
45963-0611-59		J9263		01/13/2015	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (SDV,PF,LYOPHILIZED) 100 MG	1	EA	VL	IV	EA	0.5	MG	200	01/13/2015	99/99/9999							
45963-0612-57		J9201		01/13/2015	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (SDV,USP,PF,LYOPHILIZED) 200 MG	1	EA	VL	IV	EA	200	MG	1	01/13/2015	99/99/9999							
45963-0614-51		J9206		01/13/2015	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV,USP,PF) 20 MG/ML	2	ML	VL	IV	ML	20	MG	1	01/13/2015	99/99/9999							
45963-0614-55		J9206		01/13/2015	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV,USP,PF) 20 MG/ML	5	ML	VL	IV	ML	20	MG	1	01/13/2015	99/99/9999							
45963-0615-56		J9351		01/13/2015	99/99/9999	INJECTION, TOPOTECAN, 0.1 MG	TOPOTECAN HCL (SDV,PF) 4 MG	1	EA	VL	IV	EA	0.1	MG	40	01/13/2015	99/99/9999							
45963-0619-59		J9201		01/13/2015	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (SDV,USP,PF,LYOPHILIZED) 1 GM	1	EA	VL	IV	EA	200	MG	5	01/13/2015	99/99/9999							
45963-0733-55		J9000		01/13/2015	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,SDV,PF) 2 MG/ML	5	ML	VL	IV	ML	10	MG	0.2	01/13/2015	99/99/9999							
45963-0733-57		J9000		01/13/2015	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,SDV,PF) 2 MG/ML	10	ML	VL	IV	ML	10	MG	0.2	01/13/2015	99/99/9999							
45963-0733-60		J9000		01/13/2015	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,MDV,PF) 2 MG/ML	100	ML	VL	IV	ML	10	MG	0.2	01/13/2015	99/99/9999							
45963-0733-68		J9000		01/13/2015	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,SDV,PF) 2 MG/ML	25	ML	VL	IV	ML	10	MG	0.2	01/13/2015	99/99/9999							
45963-0734-52		J9171		01/13/2015	12/21/2016	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SINGLE-USE VIAL,PF) 20 MG/ML	4	ML	VL	IV	ML	1	MG	20	01/13/2015	12/21/2016							
45963-0734-54		J9171		01/13/2015	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SINGLE-USE VIAL,PF) 20 MG/ML	1	ML	VL	IV	ML	1	MG	20	01/13/2015	99/99/9999							
45963-0734-74		J9171		01/13/2015	05/31/2016	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SINGLE-USE VIAL,PF) 20 MG/ML	7	ML	VL	IV	ML	1	MG	20	01/13/2015	05/31/2016							
55513-0192-01		J2505		02/02/2015	99/99/9999	INJECTION, PEGFILGRASTIM, 6 MG	NEULASTA (DELIVERY KIT,PF) 6 MG/0.6 ML	0.6	ML	SR	SC	ML	6	MG	1.66667	02/02/2015	99/99/9999							
67457-0440-22		J2405		12/22/2014	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HCL (25X2ML; SDV,USP,PF) 2 MG/ML	2	ML	VL	IJ	ML	1	MG	2	12/22/2014	99/99/9999							
67457-0441-20		J2405		12/22/2014	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HCL (1X20ML,MDV,USP,PF) 2 MG/ML	20	ML	VL	IJ	ML	1	MG	2	12/22/2014	99/99/9999							
67457-0582-10		J1652		01/01/2015	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PREFILLED,PF) 2.5 MG/0.5 ML	0.5	ML	SR	SC	ML	0.5	MG	10	01/01/2015	99/99/9999							
67457-0583-04		J1652		01/01/2015	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PFS,PF) 5 MG/0.4 ML	0.4	ML	SR	SC	ML	0.5	MG	25	01/01/2015	99/99/9999							
67457-0584-06		J1652		01/01/2015	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PREFILLED,PF) 7.5 MG/0.6 ML	0.6	ML	SR	SC	ML	0.5	MG	25	01/01/2015	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
67457-0585-08	J1652			01/01/2015	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PREFILLED,PF) 10 MG/0.8 ML	0.8 ML	SR	SC	ML		0.5 MG		25	01/01/2015	99/99/9999						
68001-0265-25	J9181			02/05/2015	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (USP, MDV) 20 MG/ML	5 ML	VL	IV	ML		10 MG		2	02/05/2015	99/99/9999						
68001-0265-26	J9181			02/05/2015	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (USP, MDV) 20 MG/ML	25 ML	VL	IV	ML		10 MG		2	02/05/2015	99/99/9999						
68001-0265-27	J9181			02/05/2015	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (USP, MDV) 20 MG/ML	50 ML	VL	IV	ML		10 MG		2	02/05/2015	99/99/9999						
55111-0654-01	J7520			10/27/2014	99/99/9999	SIROLIMUS, ORAL, 1 MG	SIROLIMUS 2 MG	100 EA	BO	PO	EA		1 MG		2	10/27/2014	99/99/9999						
00003-0293-28	J3301			07/01/1989	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG-40 (VIAL) 40 MG/ML	10 ML	VL	IJ	ML		10 MG		4	07/01/1989	99/99/9999						
00378-6986-01	A4216			10/08/2009	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (100X5ML,PF) 0.9%	5 ML	PC	IH	ML		10 ML		0.1	10/08/2009	99/99/9999						
00487-9007-60	A4216			07/05/2012	03/12/2017	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (PF) 0.7%	4 ML	PC	IH	ML		10 ML		0.1	07/05/2012	03/12/2017						
59762-4537-01	J1050			09/27/2004	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	MEDROXYPROGESTERONE ACETATE 150 MG/ML	1 ML	VL	IM	ML		1 MG		150	09/27/2004	99/99/9999						
42291-0594-01	None			12/04/2014	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM (USP) 2.5 MG VINORELBINE (USP;SINGLE-USE VIAL,PF) 10 MG/ML	100 EA	BO	PO	EA		2.5 MG		1	12/04/2014	99/99/9999						
45963-0607-55	J9390			02/26/2015	99/99/9999	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE (USP;SINGLE-USE VIAL,PF) 10 MG/ML	1 ML	VL	IV	ML		10 MG		1	02/26/2015	99/99/9999						
45963-0607-56	J9390			02/26/2015	99/99/9999	INJECTION, VINORELBINE TARTRATE, 10 MG	VINORELBINE (USP;SINGLE-USE VIAL,PF) 10 MG/ML	5 ML	VL	IV	ML		10 MG		1	02/26/2015	99/99/9999						
47335-0936-40	J9218			03/02/2015	99/99/9999	LEUPROLIDE ACETATE, PER 1 MG	LEUPROLIDE ACETATE (MDV) 5 MG/ML ARANESP (SINGLE USE,PF) 0.01 MG/0.4 ML	1 EA	BX	SC	EA		1 MG		5	03/02/2015	99/99/9999						
5513-0098-04	J0881			03/16/2015	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MICROGRAM (NON-ESRD USE)	NEOSTIGMINE METHYLSULFATE (MDV, USP) 0.5 MG/ML	0.4 ML	SR	IJ	ML		1 MCG		25	03/16/2015	99/99/9999						
63323-0413-10	J2710			02/18/2015	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (MDV, USP) 1 MG/ML	10 ML	VL	IV	ML		0.5 MG		1	02/18/2015	99/99/9999						
63323-0415-10	J2710			02/18/2015	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (MDV, USP) 1 MG/ML	10 ML	VL	IV	ML		0.5 MG		2	02/18/2015	99/99/9999						
63323-0565-86	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (MDV;RED LABEL) 100 MG/ML	3 ML	VL	IJ	ML		10 MG		10	04/01/2015	99/99/9999						
63323-0568-83	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (MED BLUE LABEL,PF) 30 MG/0.3 ML	0.3 ML	SR	SC	ML		10 MG		10	04/01/2015	99/99/9999						
63323-0568-84	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (BLACK LABEL,PF) 100 MG/ML	1 ML	SR	SC	ML		10 MG		10	04/01/2015	99/99/9999						
63323-0568-87	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (YELLOW LABEL,PF) 40 MG/0.4 ML	0.4 ML	SR	SC	ML		10 MG		10	04/01/2015	99/99/9999						
63323-0568-88	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (ORANGE LABEL,PF) 60 MG/0.6 ML	0.6 ML	SR	SC	ML		10 MG		10	04/01/2015	99/99/9999						
63323-0568-90	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 80 MG/0.8 ML	0.8 ML	SR	SC	ML		10 MG		10	04/01/2015	99/99/9999						
63323-0569-84	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (NAVY BLUE LABEL,PF) 150 MG/ML	1 ML	SR	SC	ML		10 MG		15	04/01/2015	99/99/9999						
63323-0569-90	J1650			04/01/2015	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PURPLE LABEL,PF) 120 MG/0.8 ML	0.8 ML	SR	SC	ML		10 MG		15	04/01/2015	99/99/9999						
67457-0211-02	J1451			09/30/2009	99/99/9999	INJECTION, FOMEPIZOLE, 15 MG	FOMEPIZOLE (1X1.5ML,PF) 1 GMMML	1.5 ML	VL	IV	ML		15 MG		66.666666	09/30/2009	99/99/9999						
67253-0102-10	J8499			03/03/2015	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	100 EA	BO	PO	EA		1 MG		1	03/03/2015	99/99/9999						
67253-0102-50	J8499			03/03/2015	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 800 MG	500 EA	BO	PO	EA		1 MG		1	03/03/2015	99/99/9999						
69238-1076-01	J7500			01/29/2015	04/28/2017	AZATHIOPRINE, ORAL, 50MG	AZATHIOPRINE (USP)50 MG	1 EA	BO	PO	EA		50 MG		1	01/29/2015	04/28/2017						
00009-5137-04	J2020			04/06/2015	99/99/9999	INJECTION, LINEZOLID, 200MG	ZYVOX (FREEFLEX BAGS) 2 MG/ML	100 ML	FC	IV	ML		200 MG		0.01	04/06/2015	99/99/9999						
00009-5140-04	J2020			04/06/2015	99/99/9999	INJECTION, LINEZOLID, 200MG	ZYVOX (FREEFLEX BAG,LATEX-FREE) 2 MG/ML	300 ML	FC	IV	ML		200 MG		0.01	04/06/2015	99/99/9999						
00069-0195-02	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED SYRINGE,PF) 2500 IU/0.2 ML	0.2 ML	SR	SC	ML		2500 IU		5	03/18/2015	99/99/9999						
00069-0196-02	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED SYRINGE,PF) 5000 IU/0.2 ML	0.2 ML	SR	SC	ML		2500 IU		10	03/18/2015	99/99/9999						
00069-0206-02	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED SYRINGE,PF) 7500 IU/0.3 ML	0.3 ML	SR	SC	ML		2500 IU		10	03/18/2015	99/99/9999						
00069-0217-02	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED SYRINGE,PF) 10000 IU/ML	1 ML	SR	SC	ML		2500 IU		4	03/18/2015	99/99/9999						
00069-0220-02	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED SYRINGE,PF) 12500 IU/0.5 ML	0.5 ML	SR	SC	ML		2500 IU		10	03/18/2015	99/99/9999						
00069-0223-02	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED SYRINGE,PF) 15000 IU/0.6 ML	0.6 ML	SR	SC	ML		2500 IU		10	03/18/2015	99/99/9999						
00069-0228-02	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (PREFILLED SYRINGE,PF) 18000 IU/0.72 ML	0.72 ML	SR	SC	ML		2500 IU		10	03/18/2015	99/99/9999						
00069-0232-01	J1645			03/18/2015	99/99/9999	INJECTION, DALTEPARIN SODIUM, PER 2500 IU	FRAGMIN (MDV) 25000 IU/ML	3.8 ML	VL	SC	ML		2500 IU		10	03/18/2015	99/99/9999						
00078-0646-81	J2353			04/10/2015	05/09/2017	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1 1/2"X20G) 10 MG	1 EA	BX	IM	EA		1 MG		10	04/10/2015	05/09/2017						
00078-0647-81	J2353			04/10/2015	12/07/2016	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1 1/2"X20G) 20 MG	1 EA	BX	IM	EA		1 MG		20	04/10/2015	12/07/2016						
00078-0648-81	J2353			04/10/2015	12/05/2016	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1 1/2"X20G) 30 MG	1 EA	BX	IM	EA		1 MG		30	04/10/2015	12/05/2016						
00169-7703-21	J2941			03/23/2015	99/99/9999	INJECTION, SOMATROPIN, 1 MG	NORDITROPIN FLEXPRO (PREFILLED PURPLE PEN) 30 MG/3 ML	3 ML	SR	SC	ML		1 MG		10	03/23/2015	99/99/9999						
00409-0212-01	J2260			04/06/2015	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (SDV,PF) 1 MG/ML	10 ML	VL	IV	ML		5 MG		0.2	04/06/2015	99/99/9999						
00409-0212-02	J2260			04/06/2015	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (SDV,PF) 1 MG/ML	20 ML	VL	IV	ML		5 MG		0.2	04/06/2015	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00409-0212-03		J2260		04/06/2015	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	MILRINONE LACTATE (SDV,PF) 1 MG/ML	50	ML	VL	IV	ML	5 MG		0.2	04/06/2015	99/99/9999						
00517-1820-01	J1205			04/01/2015	99/99/9999	INJECTION, CHLOROTHIAZIDE SODIUM, PER 500 MG	CHLOROTHIAZIDE SODIUM (USP, SDV,LYOPHILIZED) 0.5 GM	1	EA	VL	IV	EA	500 MG		1	04/01/2015	99/99/9999						
55566-2200-00	J2597			04/15/2015	99/99/9999	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DDAVP 4 MCG/ML	1	ML	AM	IJ	ML	1 MCG		4	04/15/2015	99/99/9999						
59627-0111-03	J1826			04/01/2015	99/99/9999	INJECTION, INTERFERON BETA-1A, 30 MCG	AVONEX (4 DOSE PACKS; S.D.V.) 30 MCG	4	EA	BX	IM	EA	30 MCG		1	04/01/2015	99/99/9999						
59627-0222-05	J1826			04/01/2015	99/99/9999	INJECTION, INTERFERON BETA-1A, 30 MCG	AVONEX (4 DOSE PACKS) 30 MCG/0.5 ML	1	EA	BX	MR	EA	30 MCG		1	04/01/2015	99/99/9999						
59627-0333-04	J1826			04/01/2015	99/99/9999	INJECTION, INTERFERON BETA-1A, 30 MCG	AVONEX PEN (SINGLE USE,25G,5/8") 30 MCG/0.5 ML	1	EA	BX	MR	EA	30 MCG		1	04/01/2015	99/99/9999						
63323-0311-19	J0610			03/26/2015	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (SDV,PF,LATEX-FREE) 100 MG/ML	10	ML	VL	IV	ML	10 ML		0.1	03/26/2015	99/99/9999						
63323-0311-59	J0610			03/26/2015	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (SDV,PF,LATEX-FREE) 100 MG/ML	50	ML	VL	IV	ML	10 ML		0.1	03/26/2015	99/99/9999						
63323-0311-66	J0610			03/26/2015	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (PHARMACY BULK, 2X20,PF) 100 MG/ML	100	ML	VL	IV	ML	10 ML		0.1	03/26/2015	99/99/9999						
55566-2300-00	J2597			05/10/2015	99/99/9999	INJECTION, DESMOPRESSIN ACETATE, PER 1 MCG	DDAVP 4 MCG/ML	10	ML	VL	IJ	ML	1 MCG		4	05/10/2015	99/99/9999						
00143-9570-10	J2916			04/21/2015	99/99/9999	INJECTION, SODIUM FERRIC GLUCONATE COMPLEX IN SUCROSE INJECTION, 12.5 MG	SODIUM FERRIC GLUCONATE COMPLEX IN SUCROSE (SDV) 62.5 MG/5 ML	5	ML	VL	IV	ML	12.5 MG		1	04/21/2015	99/99/9999						
00703-4805-01	J9209			04/23/2015	99/99/9999	INJECTION, MESNA, 200 MG	MESNA (M.D.V.) 100 MG/ML	10	ML	VL	IV	ML	200 MG		0.5	04/23/2015	99/99/9999						
00781-3315-70	J9263			04/14/2015	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (1X10ML,SINGLE USE,PF) 5 MG/ML	10	ML	VL	IV	ML	0.5 MG		10	04/14/2015	99/99/9999						
00781-3317-80	J9263			04/14/2015	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (1X20ML,SINGLE USE,PF) 5 MG/ML	20	ML	VL	IV	ML	0.5 MG		10	04/14/2015	99/99/9999						
55566-1801-01	J2941			05/18/2015	99/99/9999	INJECTION, SOMATROPIN, 1 MG	ZOMACTON (VIAL W/DILUENT) 5 MG	1	EA	VL	SC	EA	1 MG		5	05/18/2015	99/99/9999						
55566-1901-01	J2941			05/18/2015	99/99/9999	INJECTION, SOMATROPIN, 1 MG	ZOMACTON (VIAL W/DILUENT) 10 MG	1	EA	VL	SC	EA	1 MG		10	05/18/2015	99/99/9999						
62935-0223-05	J9217			05/07/2015	99/99/9999	LEUPROLIDE ACETATE (FOR DEPOT SUSPENSION), 7.5 MG	ELIGARD (W/SAFETY NEEDLE) 22.5 MG	1	EA	BX	SC	EA	7.5 MG		3	05/07/2015	99/99/9999						
67457-0592-10	J1652			05/06/2015	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (SRN, PREFL,27GX1/2",PF) 2.5 MG/0.5 ML	0.5	ML	SR	SC	ML	0.5 MG		10	05/06/2015	99/99/9999						
42291-0190-60	None			03/24/2015	03/19/2018	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM-COATED) 150 MG	60	EA	BO	PO	EA	150 MG		1	03/24/2015	03/19/2018						
42291-0191-12	None			03/24/2015	03/19/2018	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (USP,FILM-COATED) 500 MG	120	EA	BO	PO	EA	500 MG		1	03/24/2015	03/19/2018						
67877-0230-22	J7517			11/17/2014	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FRUIT) 200 MG/ML	225	ML	BO	PO	ML	250 MG		0.8	11/17/2014	99/99/9999						
62856-0796-01	Q9978			07/01/2015	12/31/2015	NETUPITANT 300 MG AND PALONOSETRON 0.5 MG, ORAL	AKYNZEO (HARD GELATIN) 300 MG-0.5 MG	1	EA	DP	PO	EA	300.5 MG		1	07/01/2015	12/31/2015						
00597-0053-45	J1610			04/09/2015	99/99/9999	INJECTION, GLUCAGON HYDROCHLORIDE, PER 1 MG	GLUCAGEN (VIAL) 1 MG	10	EA	VL	IJ	EA	1 MG		1	04/09/2015	99/99/9999						
00597-0260-10	J1610			04/09/2015	99/99/9999	INJECTION, GLUCAGON HYDROCHLORIDE, PER 1 MG	GLUCAGEN DIAGNOSTIC KIT (VIAL W/STERILE WATER) 1 MG	1	EA	VL	IJ	EA	1 MG		1	04/09/2015	99/99/9999						
00641-6164-10	J0706			05/14/2015	99/99/9999	INJECTION, CAFFEINE CITRATE, 5MG	CAFCIT (SINGLE USE,10X3ML,PF) 20 MG/ML	3	ML	VL	IV	ML	5 MG		4	05/14/2015	99/99/9999						
16729-0072-12	None			06/15/2015	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM COATED) 150 MG	60	EA	BO	PO	EA	150 MG		1	06/15/2015	99/99/9999						
16729-0073-29	None			06/15/2015	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (USP,FILM COATED) 500 MG	120	EA	BO	PO	EA	500 MG		1	06/15/2015	99/99/9999						
44567-0245-25	J0694			05/20/2015	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN SODIUM (USP,LATEX-FREE) 1 GM	25	EA	VL	IV	EA	1 GM		1	05/20/2015	99/99/9999						
44567-0247-10	J0694			05/20/2015	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN SODIUM (BULK PACKAGE,USP) 10 GM	10	EA	VL	IV	EA	1 GM		10	05/20/2015	99/99/9999						
00781-7157-64	J7644			09/09/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA APPROVED FINAL PRODUCT, NON COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MG	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1 MG		0.2	09/09/2011	99/99/9999						
00781-7157-64	KO	J7644	KO	09/09/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA APPROVED FINAL PRODUCT, NON COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MG	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1 MG		0.2	09/09/2011	99/99/9999						
23155-0119-01	J8499			05/28/2013	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	CALCITRIOL 0.5 MCG	100	EA	BO	PO	EA	1 MCG		1	05/28/2013	99/99/9999						
16571-0600-96	J8499			12/12/2011	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	CROMOLYN SODIUM (96X5ML,CONCENTRATE) 100MG/5ML	5	ML	PC	PO	ML	1 MG		1	12/12/2011	99/99/9999						
43598-0412-25	J7614			09/16/2014	99/99/9999	LEVALBUTERAL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5MG	LEVALBUTERAL HYDROCHLORIDE (5X5,PF), 0.31MG/3ML	3	ML	PC	IH	ML	0.5 MG		0.20666	09/16/2014	99/99/9999						
43598-0412-25	KO	J7614	KO	09/16/2014	99/99/9999	LEVALBUTERAL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5MG	LEVALBUTERAL HYDROCHLORIDE (5X5,PF), 0.31MG/3ML	3	ML	PC	IH	ML	0.5 MG		0.20666	09/16/2014	99/99/9999						
00074-4911-34	J0461			01/01/2010	02/03/2016	INJECTION, ATROPINE SULFATE, 0.01 MG	ATROPINE SULFATE (LIFESHIELD, 21GX1-1/2) 0.1 MG/ML	10	ML	SR	IJ	ML	0.01 MG		10	01/01/2010	02/03/2016						
00338-2691-75	J2175			05/02/2011	99/99/9999	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (SRN,PREFILLED,GLASS) 10 MG/ML	50	ML	SR	IJ	ML	100 MG		0.1	05/02/2011	99/99/9999						
17478-0040-01	J2060			09/21/2011	99/99/9999	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (S.D.V.) 2 MG/ML	1	ML	VL	IJ	ML	2 MG		1	09/21/2011	99/99/9999						
38779-0043-01	J2675			10/01/2012	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,MICRONIZED)	10	GM	BO	NA	GM	50 MG		20	10/01/2012	99/99/9999						
38779-0043-04	J2675			10/01/2012	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,MICRONIZED)	25	GM	BO	NA	GM	50 MG		20	10/01/2012	99/99/9999						
38779-0043-05	J2675			10/01/2012	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,MICRONIZED)	100	GM	BO	NA	GM	50 MG		20	10/01/2012	99/99/9999						
38779-0043-08	J2675			10/01/2012	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,MICRONIZED)	500	GM	BO	NA	GM	50 MG		20	10/01/2012	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
38779-0043-09	J2675			10/01/2012	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,MICRONIZED)	1000	GM	BO	NA	GM	50	MG	20	10/01/2012	99/99/9999						
38779-0082-04	J2001			10/01/2012	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	25	GM	BO	NA	GM	10	MG	100	10/01/2012	99/99/9999						
38779-0082-05	J2001			10/01/2012	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	100	GM	BO	NA	GM	10	MG	100	10/01/2012	99/99/9999						
38779-0082-08	J2001			10/01/2012	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL (U.S.P.)	500	GM	BO	NA	GM	10	MG	100	10/01/2012	99/99/9999						
38779-0189-03	J1320			10/01/2012	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	5	GM	BO	NA	GM	20	MG	50	10/01/2012	99/99/9999						
38779-0189-04	J1320			10/01/2012	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	25	GM	BO	NA	GM	20	MG	50	10/01/2012	99/99/9999						
38779-0189-05	J1320			10/01/2012	99/99/9999	INJECTION, AMITRIPTYLINE HCL, UP TO 20 MG	AMITRIPTYLINE HCL (U.S.P.)	100	GM	BO	NA	GM	20	MG	50	10/01/2012	99/99/9999						
38779-0925-05	J3360			04/23/2012	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (U.S.P.)	100	GM	BO	NA	GM	5	MG	200	04/23/2012	99/99/9999						
38779-0925-08	J3360			04/23/2012	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (U.S.P.)	500	GM	BO	NA	GM	5	MG	200	04/23/2012	99/99/9999						
38779-0925-09	J3360			04/23/2012	99/99/9999	INJECTION, DIAZEPAM, UP TO 5 MG	DIAZEPAM (U.S.P.)	1000	GM	BO	NA	GM	5	MG	200	04/23/2012	99/99/9999						
50111-0787-66	Q0144			01/10/2012	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (6X3,FILM-COATED) 250 MG	18	EA	DP	PO	EA	1	GM	0.25	01/10/2012	99/99/9999						
54569-4827-00	J7510			12/02/2011	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (2X120 ML,RED CHERRY) 15 MG/5 ML	120	ML	BO	PO	ML	5	MG	0.6	12/02/2011	99/99/9999						
54868-0753-00	J0561			01/01/2011	99/99/9999	INJECTION, PENICILLIN G BENZATHINE, 100,000 UNITS	BICILLIN L-A (TUBEX) 600000 U/ML	2	ML	SR	IM	ML	100000	UNITS	6	01/01/2011	99/99/9999						
54868-0753-01	J0561			01/01/2011	99/99/9999	INJECTION, PENICILLIN G BENZATHINE, 100,000 UNITS	BICILLIN L-A (TUBEX) 600000 U/ML	2	ML	SR	IM	ML	100000	UNITS	6	01/01/2011	99/99/9999						
54868-3349-00	J0561			01/01/2011	02/03/2016	INJECTION, PENICILLIN G BENZATHINE, 100,000 UNITS	BICILLIN L-A (M.D.V.) 300000 U/ML	10	ML	VL	IM	ML	100000	UNITS	3	01/01/2011	02/03/2016						
59762-4538-01	J1055			07/30/2011	12/31/2012	INJECTION, MEDROXYPROGESTERONE ACETATE FOR CONTRACEPTIVE USE, 150 MG	MEDROXYPROGESTERONE ACETATE (PREFILLED SYRINGE,USP) 150 MG/ML	1	ML	SR	IM	ML	150	MG	1	07/30/2011	12/31/2012						
59762-4538-01	J1050			01/01/2013	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	MEDROXYPROGESTERONE ACETATE (PREFILLED SYRINGE,USP) 150 MG/ML	1	ML	SR	IM	ML	1	MG	150	01/01/2013	99/99/9999						
62991-1707-05	J1070			10/31/2011	12/31/2014	INJECTION, TESTOSTERONE CYPIONATE, UP TO 100 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1	EA	BO	NA	GM	100	MG	10	10/31/2011	12/31/2014						
62991-1707-05	J1071			01/01/2015	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (U.S.P.)	1000	GM	VL	NA	GM	1	MG	1000	01/01/2015	99/99/9999						
66758-0016-03	J2370			03/04/2011	99/99/9999	INJECTION, PHENYLEPHRINE HCL, UP TO 1 ML	PHENYLEPHRINE HCL (USP,PF) 10 MG/ML	5	ML	VL	IJ	ML	1	ML	1	03/04/2011	99/99/9999						
00409-4883-01	J2020			06/22/2015	99/99/9999	INJECTION, LINEZOLID, 200MG	LINEZOLID 2 MG/ML	300	ML	FC	IV	ML	200	MG	0.01	06/22/2015	99/99/9999						
00781-3158-95	J0583			07/06/2015	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (SINGLE-USE,LYOPHILIZED) 250 MG	10	EA	VL	IV	EA	1	MG	250	07/06/2015	99/99/9999						
17478-0171-30	J7612			06/22/2015	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 0.5 MG	XOPENEX (PF) 1.25 MG/0.5 ML ACETYL CYSTEINE (SDV; 4X30ML,PF) 200 MG/ML	30	EA	PC	IH	EA	0.5	MG	5	06/22/2015	99/99/9999						
17478-0660-30	J0132			06/24/2015	99/99/9999	INJECTION, ACETYLCYSTEINE, 100 MG	MG/ML	30	ML	VL	IV	ML	100	MG	2	06/24/2015	99/99/9999						
18657-0117-04	J3473			07/01/2015	99/99/9999	INJECTION, HYALURONIDASE, RECOMBINANT, 1 USP UNIT	HYLENEX (4X1ML SDV) 150 U/ML	1	ML	VL	IJ	ML	1	UNIT	150	07/01/2015	99/99/9999						
23155-0521-41	J1940			08/01/2015	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	PREMIERPRO RX FUROSEMIDE (SDV) 10 MG/ML	2	ML	VL	IJ	ML	20	MG	0.5	08/01/2015	99/99/9999						
23155-0521-42	J1940			08/01/2015	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	PREMIERPRO RX FUROSEMIDE (SDV) 10 MG/ML	4	ML	VL	IJ	ML	20	MG	0.5	08/01/2015	99/99/9999						
23155-0521-44	J1940			08/01/2015	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	PREMIERPRO RX FUROSEMIDE (SDV) 10 MG/ML	10	ML	VL	IJ	ML	20	MG	0.5	08/01/2015	99/99/9999						
44567-0246-25	J0694			06/25/2015	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN SODIUM (LATEX-FREE) 2 GM	25	EA	VL	IV	EA	1	GM	2	06/25/2015	99/99/9999						
51927-3213-00	J3490			01/13/2015	99/99/9999	UNCLASSIFIED DRUGS	SUFENTANIL CITRATE (U.S.P)	1	GM	BO	NA	GM	1	GM	1	01/13/2015	99/99/9999						
38779-0101-08	J3350			10/01/2012	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P)	500	GM	BO	NA	GM	40	GM	0.025	10/01/2012	99/99/9999						
38779-0101-09	J3350			10/01/2012	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P)	1000	GM	BO	NA	GM	40	GM	0.025	10/01/2012	99/99/9999						
38779-0163-03	J3490			10/01/2012	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.,MICRONIZED)	5	GM	BO	NA	GM	1	GM	1	10/01/2012	99/99/9999						
38779-0163-04	J3490			10/01/2012	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.,MICRONIZED)	25	GM	JR	NA	GM	1	GM	1	10/01/2012	99/99/9999						
38779-0163-05	J3490			10/01/2012	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.,MICRONIZED)	100	GM	BO	NA	GM	1	GM	1	10/01/2012	99/99/9999						
38779-0163-08	J3490			10/01/2012	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.,MICRONIZED)	500	GM	JR	NA	GM	1	GM	1	10/01/2012	99/99/9999						
38779-0163-09	J3490			01/31/2011	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE (U.S.P.,MICRONIZED)	1000	GM	JR	NA	GM	1	GM	1	01/31/2011	99/99/9999						
62991-2577-01	J0456			10/31/2011	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN DIHYDRATE (U.S.P.,MICRONIZED)	1000	GM	NA	NA	GM	500	MG	2	10/31/2011	99/99/9999						
00591-3767-30	J7626			04/02/2013	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2ML,SINGLEDOSE) 0.25MG/2ML	2	ML	AM	IH	ML	0.5	MG	0.25	04/02/2013	99/99/9999						
00591-3767-30	KO	J7626	KO	04/02/2013	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2ML,SINGLEDOSE) 0.25MG/2ML	2	ML	AM	IH	ML	0.5	MG	0.25	04/02/2013	99/99/9999						
00591-3768-30	J7626			04/02/2013	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2ML,SINGLEDOSE) 0.5MG/2ML	2	ML	PC	IH	ML	0.5	MG	0.5	04/02/2013	99/99/9999						
00591-3768-30	KO	J7626	KO	04/02/2013	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2ML,SINGLEDOSE) 0.5MG/2ML	2	ML	PC	IH	ML	0.5	MG	0.5	04/02/2013	99/99/9999						
00003-3772-11	J9999			12/23/2014	12/31/2015	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	OPDIVO (PF) 10 MG/ML	4	ML	VL	IV	ML	1	MG	1	12/23/2014	12/31/2015						
00944-2510-02	J7799			10/06/2014	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	HYQVIA (PF,LATEX-FREE) 160 U/ML-10%	26.25	ML	VL	SC	ML	1	ML	1	10/06/2014	12/31/2015						
00944-2511-02	J7799			10/06/2014	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	HYQVIA (PF,LATEX-FREE) 160 U/ML-10%	52.5	ML	VL	SC	ML	1	ML	1	10/06/2014	12/31/2015						
00944-2512-02	J7799			10/06/2014	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	HYQVIA (PF,LATEX-FREE) 160 U/ML-10%	105	ML	VL	SC	ML	1	ML	1	10/06/2014	12/31/2015						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00944-2513-02		J7799		10/06/2014	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	HYQVIA (PF,LATEX-FREE) 160 U/ML-10%	210	ML	VL	SC	ML	1	ML	1	10/06/2014	12/31/2015						
00944-2514-02		J7799		10/06/2014	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	HYQVIA (PF,LATEX-FREE) 160 U/ML-10%	315	ML	VL	SC	ML	1	ML	1	10/06/2014	12/31/2015						
00093-5740-65		J7515		07/06/2015	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE, MODIFIED (SOFT GELATIN) 25 MG	30	EA	BX	PO	EA	25	MG	1	07/06/2015	99/99/9999						
65162-0801-14	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5MG	14	EA	BO	PO	EA	5	MG	1	05/26/2015	99/99/9999						
65162-0801-51	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5MG	5	EA	BO	PO	EA	5	MG	1	05/26/2015	99/99/9999						
65162-0802-14	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20MG	14	EA	BO	PO	EA	20	MG	1	05/26/2015	99/99/9999						
65162-0802-51	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20MG	5	EA	BO	PO	EA	20	MG	1	05/26/2015	99/99/9999						
65162-0803-14	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100MG	14	EA	BO	PO	EA	100	MG	1	05/26/2015	99/99/9999						
65162-0803-51	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100MG	5	EA	BO	PO	EA	100	MG	1	05/26/2015	99/99/9999						
65162-0806-51	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE 250MG	5	EA	BO	PO	EA	250	MG	1	05/26/2015	99/99/9999						
65162-0804-14	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140MG	14	EA	BO	PO	EA	20	MG	7	05/26/2015	99/99/9999						
65162-0804-51	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140MG	5	EA	BO	PO	EA	20	MG	7	05/26/2015	99/99/9999						
65162-0805-14	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180MG	14	EA	BO	PO	EA	20	MG	9	05/26/2015	99/99/9999						
65162-0805-51	None			05/26/2015	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180MG	5	EA	BO	PO	EA	20	MG	9	05/26/2015	99/99/9999						
00002-7712-27	J1815			05/28/2015	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG (2X3ML) 200 U/ML	3	ML	SR	SC	ML	5	U	40	05/28/2015	99/99/9999						
00409-4882-01	J2020			07/07/2015	10/18/2017	INJECTION, LINEZOLID, 200MG	LINEZOLID 2 MG/ML	300	ML	FC	IV	ML	200	MG	0.01	07/07/2015	10/18/2017						
00409-8300-10	J0583			08/03/2015	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (SINGLE-USE,LYOPHILIZED) 250 MG	10	EA	VL	IV	EA	1	MG	250	08/03/2015	99/99/9999						
00944-4177-05	J2724			07/01/2015	99/99/9999	INJECTION, PROTEIN C CONCENTRATE, INTRAVENOUS, HUMAN, 10 IU	CEPROTIN (POTENCY PRINTED ON VIAL) 1 IU	1	EA	VL	IV	EA	10	IU	0.1	07/01/2015	99/99/9999						
00944-4179-10	J2724			07/01/2015	99/99/9999	INJECTION, PROTEIN C CONCENTRATE, INTRAVENOUS, HUMAN, 10 IU	CEPROTIN (POTENCY PRINTED ON VIAL) 1 IU	1	EA	VL	IV	EA	10	IU	0.1	07/01/2015	99/99/9999						
13533-0701-01	J0256			09/01/2015	99/99/9999	INJECTION, ALPHA 1 PROTEINASE INHIBITOR (HUMAN), NOT OTHERWISE SPECIFIED, 10 MG	PROLASTIN-C (1000MG,LYOPHILIZED) 1 MG	1	EA	VL	IV	EA	10	MG	0.1	09/01/2015	99/99/9999						
42023-0179-05	J0592			07/29/2015	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE (5X1ML,SDV) 0.3 MG/ML	1	ML	VL	IJ	ML	0.1	MG	3	07/29/2015	99/99/9999						
63323-0750-10	J9263			07/30/2015	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (SINGLE-USE VIAL; USP,PF) 5 MG/ML	10	ML	VL	IV	ML	0.5	MG	10	07/30/2015	99/99/9999						
63323-0850-74	J2280			07/20/2015	99/99/9999	INJECTION, MOXIFLOXACIN, 100 MG	MOXIFLOXACIN HCL (FREEFLEX,LATEX-FREE) 400 MG/250 ML	250	ML	FC	IV	ML	100	MG	0.016	07/20/2015	99/99/9999						
65162-0914-46	J7682			07/16/2015	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5	ML	PC	IH	ML	300	MG	0.2	07/16/2015	99/99/9999						
65162-0914-46	KO	J7682	KO	07/16/2015	99/99/9999	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (4 AMPULES X 14 POUCHES) 300 MG/5 ML	5	ML	PC	IH	ML	300	MG	0.2	07/16/2015	99/99/9999						
00074-3012-07	J7799			02/03/2015	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DUOPA 4.63 MG/ML-20 MG/ML	100	ML	BX	NA	ML	100	ML	0.01	02/03/2015	12/31/2015						
55513-0150-01	J7799			12/16/2014	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	BLINCYTO (INNER VIAL NDC,PF) 35 MCG	1	EA	VL	IV	EA	1	MCG	1	12/16/2014	12/31/2015						
55513-0160-01	J7799			12/16/2014	12/31/2015	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	BLINCYTO (W/ SOLN STABILIZER,PF) 35 MCG	1	EA	VL	IV	EA	1	MCG	1	12/16/2014	12/31/2015						
00143-8596-25	J2501			08/17/2015	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (MDV) 0.005 MG/1 ML	2	ML	VL	IV	ML	1	MCG	5	08/17/2015	99/99/9999						
00143-9624-25	J2501			08/17/2015	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (SDV) 0.005 MG/1 ML	1	ML	VL	IV	ML	1	MCG	5	08/17/2015	99/99/9999						
00143-9625-25	J2501			08/17/2015	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (SDV) 0.002 MG/1 ML	1	ML	VL	IV	ML	1	MCG	2	08/17/2015	99/99/9999						
54766-0149-23	J0630			08/31/2015	09/15/2016	INJECTION, CALCITONIN SALMON, UP TO 400 UNITS	MIACALCIN 200 IU/1 ML	2	ML	VL	IJ	ML	400	U	0.5	08/31/2015	09/15/2016						
67457-0593-04	J1652			08/07/2015	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (27GX1/2",PF) 5 MG/0.4 ML	0.4	ML	SR	SC	ML	0.5	MG	25	08/07/2015	99/99/9999						
00003-0293-05	J3301			02/01/1989	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG-40 (VIAL) 40 MG/1 ML	1	ML	VL	IJ	ML	10	MG	4	02/01/1989	99/99/9999						
00003-0293-20	J3301			07/01/1989	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	KENALOG-40 (VIAL) 40 MG/1 ML	5	ML	VL	IJ	ML	10	MG	4	07/01/1989	99/99/9999						
25021-0402-01	J1644			07/06/2010	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,LATEX-FREE) 5000 U/ML	1	ML	VL	IJ	ML	1000	U	5	07/06/2010	99/99/9999						
50242-0140-01	J8999			01/31/2012	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	ERIVEDGE 150 MG	28	EA	BO	PO	EA	1	MG	1	01/31/2012	99/99/9999						
00781-7516-87	KO	J7626	KO	08/20/2015	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.5 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	0.5	08/20/2015	99/99/9999						
24208-0002-02	J3471			09/22/2015	99/99/9999	INJECTION, HYALURONIDASE, OVINE, PRESERVATIVE FREE, PER 1 USP UNIT (UP TO 999 USP UNITS)	USP 1 UNIT	1.2	ML	VL	SC	ML	1	UNIT	200	09/22/2015	99/99/9999						
67457-0348-15	J0295			09/04/2015	11/30/2017	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM 1 GM-0.5 GM	1	EA	VL	IJ	EA	1.5	GM	1	09/04/2015	11/30/2017						
67457-0349-03	J0295			09/04/2015	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM 2 GM-1 GM	1	EA	VL	IJ	EA	1.5	GM	2	09/04/2015	99/99/9999						
67457-0649-10	J0295			09/04/2015	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM 10 GM-5 GM	1	EA	VL	IV	EA	1.5	GM	10	09/04/2015	99/99/9999						
39822-0500-04	J0360			09/21/2015	99/99/9999	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HCL (USP) 20 MG/1 ML	1	ML	VL	IJ	ML	20	MG	1	09/21/2015	99/99/9999						
00409-8300-15	J0583			10/05/2015	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (SINGLE-USE ADD-VANTAGE) 250 MG	10	EA	VL	IV	EA	1	MG	250	10/05/2015	99/99/9999						
60505-0761-01	J0694			10/06/2015	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN SODIUM (BULK PACKAGE) 10 GM	1	EA	VL	IV	EA	1	GM	10	10/06/2015	99/99/9999						
00781-7516-87	J7626			08/20/2015	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.5 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	0.5	08/20/2015	99/99/9999						
60505-0760-01	J0694			10/06/2015	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN SODIUM 2 GM	1	EA	VL	IV	EA	1	GM	2	10/06/2015	99/99/9999						
60505-0759-01	J0694			10/06/2015	99/99/9999	INJECTION, CEFOXITIN SODIUM, 1 GM	CEFOXITIN SODIUM 1 GM	1	EA	VL	IV	EA	1	GM	1	10/06/2015	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
25021-0305-20	J1205			10/15/2015	99/99/9999	INJECTION, CHLOROTHIAZIDE SODIUM, PER 500 MG	CHLOROTHIAZIDE SODIUM (USP, SDV,PF LATEX-FREE) 0.5 GM	1 EA	VL	IV	EA		500 MG		1	10/15/2015	99/99/9999						
17478-0987-12	J1270			09/21/2015	10/21/2016	INJECTION, DOXERCALCIFEROL, 1 MCG	DOXERCALCIFEROL (2MLX10, SDV) 2 MCG/1 ML	2 ML	VL	IV	ML		1 MCG		2	09/21/2015	10/21/2016						
68982-0840-04	J1568			09/15/2015	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (10GM/VIAL,S/D TREATED) 50 MG/1 ML	200 ML	VL	IV	ML		500 MG		0.1	09/15/2015	99/99/9999						
68982-0840-01	J1568			09/15/2015	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (1GM/VIAL,S/D TREATED) 50 MG/1 ML	20 ML	VL	IV	ML		500 MG		0.1	09/15/2015	99/99/9999						
68982-0840-02	J1568			09/15/2015	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (2.5GM/VIAL,S/D TREATED) 50 MG/1 ML	50 ML	VL	IV	ML		500 MG		0.1	09/15/2015	99/99/9999						
68982-0840-03	J1568			09/15/2015	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (5GM/VIAL,S/D TREATED) 50 MG/1 ML	100 ML	VL	IV	ML		500 MG		0.1	09/15/2015	99/99/9999						
68982-0840-05	J1568			09/15/2015	99/99/9999	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NON-LYOPHILIZED (E.G. LIQUID), 500 MG	OCTAGAM (LATEX-FREE) 50 MG/1 ML	500 ML	VL	IV	ML		500 MG		0.1	09/15/2015	99/99/9999						
55150-0191-83	J1740			09/08/2015	99/99/9999	INJECTION, IBANDRONATE SODIUM, 1 MG	IBANDRONATE SODIUM 1 MG/1 ML	3 ML	SR	IV	ML		1 MG		1	09/08/2015	99/99/9999						
00264-3183-11	J2185			09/15/2015	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM 500 MG	24 EA	FC	IV	EA		100 MG		5	09/15/2015	99/99/9999						
00264-3185-11	J2185			09/15/2015	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM 1 GM	24 EA	FC	IV	EA		100 MG		10	09/15/2015	99/99/9999						
60505-0686-01	J2543			10/06/2015	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SDV) 2 GM-0.25 GM	1 EA	VL	IV	EA		1.125 GM		2	10/06/2015	99/99/9999						
60505-0687-01	J2543			10/06/2015	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SDV) 3 GM-0.375 GM	1 EA	VL	IV	EA		1.125 GM		3	10/06/2015	99/99/9999						
60505-0688-01	J2543			10/06/2015	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SDV) 4 GM-0.5 GM	1 EA	VL	IV	EA		1.125 GM		4	10/06/2015	99/99/9999						
00143-9565-01	J8340			08/31/2015	99/99/9999	INJECTION, THIOTEPA, 15 MG	THIOTEPA (LYOPHILIZED) 15 MG	1 EA	VL	IJ	EA		15 MG		1	08/31/2015	99/99/9999						
00548-9090-10	J3470			10/05/2015	99/99/9999	INJECTION, HYALURONIDASE, UP TO 150 UNITS	AMPHADASE 150 U/1 ML	10 EA	VL	SC	EA		150 UNITS		1	10/05/2015	99/99/9999						
00574-0820-10	J1071			12/12/2014	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (1x10 ML,USP) 200 MG/1 ML	10 ML	VL	IM	ML		1 MG		200	12/12/2014	99/99/9999						
49452-0001-03	J0133			06/01/2015	10/17/2016	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	25 GM	BO	NA	GM		5 MG		200	06/01/2015	10/17/2016						
49452-0027-02	J0745			06/01/2015	10/17/2016	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	25 GM	BO	NA	GM		30 MG	33.33333		06/01/2015	10/17/2016						
49452-0027-03	J0745			06/01/2015	10/17/2016	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	5 GM	JR	NA	GM		30 MG	33.33333		06/01/2015	10/17/2016						
49452-0430-01	J0280			06/01/2015	99/99/9999	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	1000 GM	BO	NA	GM		250 MG		4	06/01/2015	99/99/9999						
49452-0430-02	J0280			06/01/2015	10/17/2016	INJECTION, AMINOPHYLLIN, UP TO 250 MG	AMINOPHYLLINE ANHYDROUS (U.S.P.)	500 GM	BO	NA	GM		250 MG		4	06/01/2015	10/17/2016						
49452-0735-01	J9017			06/01/2015	10/17/2016	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (A.C.S.,REAGENT)	125 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-0735-02	J9017			06/01/2015	10/17/2016	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (A.C.S.,REAGENT)	500 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-0783-02	J7501			06/01/2015	10/17/2016	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (U.S.P.)	5 GM	BO	NA	GM		100 MG		10	06/01/2015	10/17/2016						
49452-0970-01	J3490			06/01/2015	10/17/2016	UNCLASSIFIED DRUGS	BENZOCAINE (U.S.P.)	125 GM	BO	NA	GM		1 EA		1	06/01/2015	10/17/2016						
49452-0970-02	J3490			06/01/2015	10/17/2016	UNCLASSIFIED DRUGS	BENZOCAINE (U.S.P.)	500 GM	BO	NA	GM		1 EA		1	06/01/2015	10/17/2016						
49452-0970-03	J3490			06/01/2015	10/17/2016	UNCLASSIFIED DRUGS	BENZOCAINE (U.S.P.)	2500 GM	BO	NA	GM		1 EA		1	06/01/2015	10/17/2016						
49452-1072-02	J3490			06/01/2015	99/99/9999	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	1 GM	BO	NA	GM		1 EA		1	06/01/2015	99/99/9999						
49452-1309-01	J0945			06/01/2015	10/17/2016	INJECTION, BROMPHENIRAMINE MALEATE, PER 10 MG	BROMPHENIRAMINE MALEATE (U.S.P.)	25 GM	BO	NA	GM		10 MG		100	06/01/2015	10/17/2016						
49452-1317-01	J0595			06/01/2015	10/17/2016	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (U.S.P.)	1 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-1317-02	J0595			06/01/2015	10/17/2016	INJECTION, BUTORPHANOL TARTRATE, 1 MG	BUTORPHANOL TARTRATE (U.S.P.)	1 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-2147-02	J0735			06/01/2015	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	1 GM	BO	NA	GM		1 MG		1000	06/01/2015	99/99/9999						
49452-2147-03	J0735			06/01/2015	99/99/9999	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	5 GM	BO	NA	GM		1 MG		1000	06/01/2015	99/99/9999						
49452-2210-02	J0760			06/01/2015	10/17/2016	INJECTION, COLCHICINE, PER 1MG	COLCHICINE (U.S.P.)	1 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-2210-03	J0760			06/01/2015	10/17/2016	INJECTION, COLCHICINE, PER 1MG	COLCHICINE (U.S.P.)	5 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-8253-01	J0592			06/01/2015	10/17/2016	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE (U.S.P.)	0.1 GM	JR	NA	GM		0.1 MG		10000	06/01/2015	10/17/2016						
49452-8253-02	J0592			06/01/2015	10/17/2016	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE (U.S.P.)	0.5 GM	JR	NA	GM		0.1 MG		10000	06/01/2015	10/17/2016						
49452-8253-03	J0592			06/01/2015	10/17/2016	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE (U.S.P.)	1 GM	JR	NA	GM		0.1 MG		10000	06/01/2015	10/17/2016						
00093-5740-19	J7515			07/06/2015	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE, MODIFIED (INNERPACK,SOFT GELATIN) 25 MG	1 EA	BP	PO	EA		25 MG		1	07/06/2015	99/99/9999						
00093-5741-65	J7515			09/28/2015	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE, MODIFIED (USP,SOFTGEL) 50 MG	30 EA	BX	PO	EA		25 MG		2	09/28/2015	99/99/9999						
00093-5742-65	J7502			08/27/2015	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (USP,MODIFIED,SOFTGEL) 100 MG	30 EA	BX	PO	EA		100 MG		1	08/27/2015	99/99/9999						
49999-0028-30	J7512			01/01/2016	12/31/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	30 EA	BO	PO	EA		1 MG		10	01/01/2016	12/31/2016						
51991-0458-01	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.) 1 MG	100 EA	BO	PO	EA		1 MG		1	01/01/2016	99/99/9999						
49999-0028-40	J7512			01/01/2016	06/01/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	40 EA	BO	PO	EA		1 MG		10	01/01/2016	06/01/2017						
67457-0520-40	J9280			03/19/2018	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MITOMYCIN (SDV,PF) 40 MG	1 EA	VL	IV	EA		5 MG		8	03/19/2018	99/99/9999						
00054-0018-20	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (10X10) 20 MG	100 EA	BX	PO	EA		1 MG		20	01/01/2016	99/99/9999						
49999-0110-14	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	14 EA	BO	PO	EA		1 MG		20	01/01/2016	99/99/9999						
49999-0110-00	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	100 EA	BO	PO	EA		1 MG		20	01/01/2016	99/99/9999						
00054-0018-25	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	100 EA	BO	PO	EA		1 MG		20	01/01/2016	99/99/9999						
49999-0110-12	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG																	

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
54569-3043-01	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	12 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
52959-0127-10	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	10 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
54569-3043-05	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	14 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
52959-0126-12	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	12 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
54569-3302-00	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	60 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
67457-0863-01	J1626			03/21/2018	99/99/9999	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X1ML,SDV,PF,LATEX-FREE) 1 MG/1 ML PREDNISONE ANHYDROUS (U.S.P.,MICRONIZED)	1 ML	VL IV ML				100 MCG		10	03/21/2018	99/99/9999						
38779-0154-08	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.,MICRONIZED)	500 GM	BO NA GM	1 MG			1000		1000	01/01/2016	99/99/9999						
38779-0154-05	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.,MICRONIZED)	100 GM	BO NA GM	1 MG			1000		1000	01/01/2016	99/99/9999						
38779-0154-04	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.,MICRONIZED)	25 GM	BO NA GM	1 MG			1000		1000	01/01/2016	99/99/9999						
38779-0154-03	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.,MICRONIZED)	5 GM	BO NA GM	1 MG			1000		1000	01/01/2016	99/99/9999						
54569-3302-01	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	20 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
54569-3413-00	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	21 EA	DP PO EA	1 MG			5		5	01/01/2016	99/99/9999						
54569-3043-02	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	6 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999	01/01/2002	06/10/2003	4			
52959-0126-40	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	40 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
49999-0008-05	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	5 EA	NA PO EA	1 MG			5		5	01/01/2016	99/99/9999						
52959-0126-60	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	60 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-50	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	50 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-45	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	45 EA	NA PO EA	1 MG			10		10	01/01/2016	99/99/9999						
54569-0332-05	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	100 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
54569-0332-09	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	18 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
54569-0333-00	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 50 MG	8 EA	BO PO EA	1 MG			50		50	01/01/2016	99/99/9999						
52959-0126-07	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	7 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-42	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	42 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-10	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	10 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-37	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	37 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-30	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	30 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-25	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	25 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-21	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	21 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-20	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	20 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-18	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	18 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0126-15	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	15 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
52959-0127-07	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	7 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
52959-0126-44	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	44 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
00591-5443-01	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	100 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
00603-5338-21	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	100 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
00603-5338-15	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (DOSE PACK) 10 MG	21 EA	DP PO EA	1 MG			10		10	01/01/2016	99/99/9999						
00603-5337-32	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	1000 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999						
00603-5337-31	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (DOSE PACK) 5 MG	48 EA	DP PO EA	1 MG			5		5	01/01/2016	99/99/9999						
00603-5337-15	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (DOSE PACK) 5 MG	21 EA	DP PO EA	1 MG			5		5	01/01/2016	99/99/9999						
00603-5335-32	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 1 MG	1000 EA	BO PO EA	1 MG			1		1	01/01/2016	99/99/9999						
00603-5335-21	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 1 MG	100 EA	BO PO EA	1 MG			1		1	01/01/2016	99/99/9999						
00463-6140-10	J7512			01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNICOT 10 MG	1000 EA	NA PO EA	1 MG			10		10	01/01/2016	02/03/2016						
00591-5443-05	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	500 EA	BO PO EA	1 MG			20		20	01/01/2016	99/99/9999						
00603-5338-32	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	1000 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
00591-5442-10	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	1000 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
00591-5442-05	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	500 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
00591-5442-01	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	100 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999						
00591-5052-10	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	1000 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999						
00591-5052-01	J7512			01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	100 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999						
00463-6155-10	J7512			01/01/2016	01/01/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNICOT 5 MG	1000 EA	NA PO EA	1 MG			5		5	01/01/2016	01/01/2016						
00463-6141-10	J7512			01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNICOT 20 MG	1000 EA	NA PO EA	1 MG			20		20	01/01/2016	02						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
63874-0327-24		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	24 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
63874-0392-40		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	40 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
63874-0327-21		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	21 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
63874-0373-33		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	33 EA	BO PO EA	1 MG			5		5	01/01/2016	02/03/2016							
63874-0392-28		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	28 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
63874-0373-30		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	30 EA	BO PO EA	1 MG			5		5	01/01/2016	02/03/2016							
63874-0392-24		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	24 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
63874-0392-21		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	21 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
63874-0327-50		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	50 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
63874-0327-60		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	60 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
63874-0373-01		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	100 EA	BO PO EA	1 MG			5		5	01/01/2016	02/03/2016							
66267-0171-42		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	42 EA	BO PO EA	1 MG			10		10	01/01/2016	99/99/9999							
63874-0373-21		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	21 EA	BO PO EA	1 MG			5		5	01/01/2016	02/03/2016							
55289-0373-30		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	30 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999							
63874-0373-10		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	10 EA	BO PO EA	1 MG			5		5	01/01/2016	02/03/2016							
63874-0373-15		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	15 EA	BO PO EA	1 MG			5		5	01/01/2016	02/03/2016							
63874-0327-42		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	42 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
63874-0327-14		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	14 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
55289-0373-36		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	36 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999							
55289-0373-42		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	42 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999							
55289-0373-46		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	46 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999							
63874-0392-10		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	10 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
63874-0392-06		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	60 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
63874-0327-02		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	1000 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
63874-0392-15		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	15 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
55289-0373-55		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	55 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999							
63874-0392-02		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	1000 EA	BO PO EA	1 MG			20		20	01/01/2016	02/03/2016							
63874-0327-10		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	10 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
63874-0327-12		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	12 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
55289-0373-60		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	60 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999							
63874-0327-25		J7512		01/01/2016	10/17/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	25 EA	BO PO EA	1 MG			10		10	01/01/2016	10/17/2016							
55289-0438-20		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	20 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
55289-0438-40		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	40 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
63874-0327-20		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	20 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
55289-0438-60		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	60 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
55289-0438-50		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	50 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
55289-0438-42		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (USP) 10 MG	42 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
63874-0327-19		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	19 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
55289-0373-72		J7512		01/01/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	72 EA	BO PO EA	1 MG			5		5	01/01/2016	99/99/9999							
63874-0327-15		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	15 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
55289-0438-38		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	38 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
55289-0438-36		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	36 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
55289-0438-30		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	30 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
55289-0438-21		J7512		01/01/2016	03/08/2017	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	21 EA	BO PO EA	1 MG			10		10	01/01/2016	03/08/2017							
63874-0327-18		J7512		01/01/2016	02/03/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	18 EA	BO PO EA	1 MG			10		10	01/01/2016	02/03/2016							
62856-0796-01		J8655		01/01/2016	03/31/2017	Netupitant 300 mg and palonosetron 0.5 mg, oral	AKYNZEO (HARD GELATIN) 300 MG-0.5 MG	1 EA	DP PO EA	300.5 MG			1		1	01/01/2016	03/31/2017							
55513-0150-01		J9039		01/01/2016	99/99/9999	INJECTION, BLINATUMOMAB, 1 MICROGRAM	BLINCYTO (INNER VIAL NDC.PF) 35 MCG	1 EA	VL IV EA	1 MCG			35		35	01/01/2016	99/99/9999							
55513-0160-01		J9039		01/01/2016	99/99/9999	INJECTION, BLINATUMOMAB, 1 MICROGRAM	BLINCYTO (W/ SOLN STABILIZER.PF) 35 MCG	1 EA	VL IV EA	1 MCG			35		35	01/01/2016	99/99/9999							
00074-3012-07		J7340		01/01/2016	99/99/9999	CARBIDOPA 5 MG/LEVODOPA 20 MG ENTERAL SUSPENSION, 100 ML	DUOPA 4.63 MG/ML-20 MG/ML	100 ML	BX NA ML	25 MG			1		1	01/01/2016	99/99/9999							
00944-2514-02		J1575		01/01/2016	99/99/9999	INJECTION, IMMUNE GLOBULIN/HYALURONIDASE, (HYQVIA), 100 MG IMMUNEGLOBULIN	HYQVIA (PF,LATEX-FREE) 160 U/ML-10%	315 ML	VL SC ML	100 MG			1		1	01/01/2016	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
60505-0752-03	J0696			11/02/2015	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (SDV, USP, CRYSTALLINE) 1 GM	1 EA	VL	IJ	EA		250 MG		4	11/02/2015	99/99/9999						
60505-0753-03	J0696			11/02/2015	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (SDV, USP, CRYSTALLINE) 2 GM	1 EA	VL	IJ	EA		250 MG		8	11/02/2015	99/99/9999						
23155-0547-41	J2405			11/01/2015	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (SDV, PF) 2 MG/1 ML	2 ML	VL	IJ	ML		1 MG		2	11/01/2015	99/99/9999						
23155-0549-31	J2405			11/01/2015	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV) 2 MG/1 ML	20 ML	VL	IJ	ML		1 MG		2	11/01/2015	99/99/9999						
23155-0547-42	J2405			11/01/2015	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (SDV, PF) 2 MG/1 ML	2 ML	VL	IJ	ML		1 MG		2	11/01/2015	99/99/9999						
17478-0174-24	J7614			10/20/2015	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	XOPENEX (PF) 1.25 MG/3 ML	3 ML	PC	IH	ML		0.5 MG		0.83333	10/20/2015	99/99/9999						
24492-0899-99	J7682			11/01/2015	02/16/2016	MILLIGRAMS	TOBRAMYCIN (PAK PF) 300 MG/5 ML	5 ML	PC	IH	ML		300 MG		0.2	11/01/2015	02/16/2016						
49452-0028-01	J2270			06/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	5 GM	JR	NA	GM		10 MG		100	06/01/2015	99/99/9999						
49452-0028-02	J2270			06/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE (U.S.P.)	25 GM	JR	NA	GM		10 MG		100	06/01/2015	99/99/9999						
49452-0028-03	J2270			06/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, UP TO 10 MG	MORPHINE SULFATE	100 GM	JR	NA	GM		10 MG		100	06/01/2015	99/99/9999						
49452-0029-01	J1170			06/01/2015	10/17/2016	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	1 GM	BO	NA	GM		4 MG		250	06/01/2015	10/17/2016						
49452-0029-02	J1170			06/01/2015	10/17/2016	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	5 GM	JR	NA	GM		4 MG		250	06/01/2015	10/17/2016						
49452-0029-04	J1170			06/01/2015	10/17/2016	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	25 GM	JR	NA	GM		4 MG		250	06/01/2015	10/17/2016						
49452-0031-03	J2175			06/01/2015	10/17/2016	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	5 GM	BO	NA	GM		100 MG		10	06/01/2015	10/17/2016						
49452-0032-01	J3010			06/01/2015	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	1 GM	BO	NA	GM		0.1 MG		10000	06/01/2015	99/99/9999						
49452-0032-02	J3010			06/01/2015	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (U.S.P.)	0.1 GM	JR	NA	GM		0.1 MG		10000	06/01/2015	99/99/9999						
49452-1775-01	J1955			06/01/2015	10/17/2016	INJECTION, LEVOCARNITINE, PER 1 GM	L-CARNITINE FREE BASE	25 GM	BO	NA	GM		1 GM		1	06/01/2015	10/17/2016						
49452-1775-02	J1955			06/01/2015	10/17/2016	INJECTION, LEVOCARNITINE, PER 1 GM	L-CARNITINE FREE BASE	100 GM	BO	NA	GM		1 GM		1	06/01/2015	10/17/2016						
49452-2400-02	J3420			06/01/2015	10/17/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	1 GM	BO	NA	GM		1000 MCG		1000	06/01/2015	10/17/2016						
49452-2400-03	J3420			06/01/2015	10/17/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	5 GM	BO	NA	GM		1000 MCG		1000	06/01/2015	10/17/2016						
49452-2460-01	J1094			06/01/2015	10/17/2016	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE ANHYDROUS (U.S.P. MICRONIZED)	5 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-2460-02	J1094			06/01/2015	99/99/9999	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE ANHYDROUS (U.S.P. MICRONIZED)	25 GM	BO	NA	GM		1 MG		1000	06/01/2015	99/99/9999						
49452-2460-03	J1094			06/01/2015	10/17/2016	INJECTION, DEXAMETHASONE ACETATE, 1 MG	DEXAMETHASONE ACETATE ANHYDROUS (U.S.P. MICRONIZED)	1 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-2588-01	J1212			06/01/2015	10/17/2016	INJECTION, DIMETHYL SULFOXIDE, 50%, 50 ML	DIMETHYL SULFOXIDE (U.S.P.)	500 ML	BO	NA	ML		50 ML		0.02	06/01/2015	10/17/2016						
49452-2612-02	J1160			06/01/2015	10/17/2016	INJECTION, DIGOXIN, UP TO 0.5 MG	DIGOXIN (U.S.P.)	1 GM	BO	NA	GM		0.5 MG		200	06/01/2015	10/17/2016						
49452-2640-01	J1200			06/01/2015	10/17/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (U.S.P.)	100 GM	BO	NA	GM		50 MG		20	06/01/2015	10/17/2016						
49452-2640-02	J1200			06/01/2015	10/17/2016	INJECTION, DIPHENHYDRAMINE HCL, UP TO 50 MG	DIPHENHYDRAMINE HCL (U.S.P.)	500 GM	BO	NA	GM		50 MG		20	06/01/2015	10/17/2016						
49452-2702-03	J3520			06/01/2015	10/17/2016	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM DIHYDRATE (U.S.P.)	125 GM	BO	NA	GM		150 MG		6.66666	06/01/2015	10/17/2016						
49452-2740-01	J7799			06/01/2015	10/17/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	EPINEPHRINE (U.S.P.)	100 GM	BO	NA	GM		1 GM		1	06/01/2015	10/17/2016						
49452-2791-01	J1380			06/01/2015	10/17/2016	INJECTION, ESTRADIOL VALERATE, UP TO 10 MG	ESTRADIOL VALERATE (U.S.P.)	1 GM	BO	NA	GM		10 MG		100	06/01/2015	10/17/2016						
49452-2791-02	J1380			06/01/2015	10/17/2016	INJECTION, ESTRADIOL VALERATE, UP TO 10 MG	ESTRADIOL VALERATE (U.S.P.)	5 GM	BO	NA	GM		10 MG		100	06/01/2015	10/17/2016						
49452-2795-01	J1435			06/01/2015	10/17/2016	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	1 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-2795-02	J1435			06/01/2015	10/17/2016	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	5 GM	BO	NA	GM		1 MG		1000	06/01/2015	10/17/2016						
49452-2795-04	J1435			09/01/2015	10/17/2016	INJECTION, ESTRONE, PER 1 MG	ESTRONE (U.S.P.)	25 GM	BO	NA	GM		1 MG		1000	09/01/2015	10/17/2016						
49452-3038-04	J3490			09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	FAMOTIDINE (U.S.P.)	100 GM	BO	NA	GM		1 GM		1	09/01/2015	10/17/2016						
49452-3038-05	J3490			09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	FAMOTIDINE (U.S.P.)	500 GM	BO	NA	GM		1 GM		1	09/01/2015	10/17/2016						
49452-3175-01	J9190			06/01/2015	10/17/2016	INJECTION, FLUOROURACIL, 500 MG	5-FLUOROURACIL (U.S.P.)	1 GM	BO	NA	GM		500 MG		2	06/01/2015	10/17/2016						
49452-3175-02	J9190			06/01/2015	10/17/2016	INJECTION, FLUOROURACIL, 500 MG	5-FLUOROURACIL (U.S.P.)	5 GM	BO	NA	GM		500 MG		2	06/01/2015	10/17/2016						
49452-3175-03	J9190			06/01/2015	10/17/2016	INJECTION, FLUOROURACIL, 500 MG	5-FLUOROURACIL (U.S.P.)	25 GM	BO	NA	GM		500 MG		2	06/01/2015	10/17/2016						
49452-3175-04	J9190			06/01/2015	10/17/2016	INJECTION, FLUOROURACIL, 500 MG	5-FLUOROURACIL (U.S.P.)	100 GM	BO	NA	GM		500 MG		2	06/01/2015	10/17/2016						
49452-3222-01	J1940			06/01/2015	10/17/2016	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (U.S.P./N.F.)	25 GM	BO	NA	GM		20 MG		50	06/01/2015	10/17/2016						
49452-3446-01	J1630			06/01/2015	10/17/2016	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	5 GM	BO	NA	GM		5 MG		200	06/01/2015	10/17/2016						
49452-3446-02	J1630			06/01/2015	10/17/2016	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (U.S.P.)	25 GM	BO	NA	GM		5 MG		200	06/01/2015	10/17/2016						
49452-3590-01	J1700			06/01/2015	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P. MICRONIZED)	5 GM	BO	NA	GM		25 MG		40	06/01/2015	99/99/9999						
49452-3590-02	J1700			06/01/2015	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P. MICRONIZED)	25 GM	BO	NA	GM		25 MG		40	06/01/2015	99/99/9999						
49452-3590-03	J1700			06/01/2015	99/99/9999	INJECTION, HYDROCORTISONE ACETATE, UP TO 25 MG	HYDROCORTISONE ACETATE (U.S.P. MICRONIZED)	100 GM	BO	NA	GM		25 MG		40	06/01/2015	99/99/9999						
49452-3652-02	J3410			06/01/2015	99/99/9999	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (U.S.P.)	25 GM	BO	NA	GM		25 MG		40	06/01/2015	99/99/9999						
49452-3659-01	Q0177			06/01/2015	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (U.S.P./N.F.)	25 GM	BO	NA	GM		25 MG		40	06/01/2015	99/99/9999						
49452-3659-02	Q0177			06/01/2015	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE (U.S.P./N.F.)	100 GM	BO	NA	GM		25 MG		40	06/01/2015	99/99/9999						
49452-3845-01	J1835			06/01/2015	10/17/2016	INJECTION, ITRACONAZOLE, 50 MG	ITRACONAZOLE	1 GM	BO	NA	GM		50 MG		20	06/01/2015	10/17						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
49452-4036-04		J0640		09/01/2015	10/17/2016	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (U.S.P.)	0.1	GM	BO	NA	GM	50	MG	20	09/01/2015	10/17/2016							
49452-4050-01	J2001			06/01/2015	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL MONOHYDRATE (U.S.P.)	25	GM	BO	NA	GM	10	MG	100	06/01/2015	99/99/9999							
49452-4050-02	J2001			06/01/2015	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL MONOHYDRATE (U.S.P.)	100	GM	BO	NA	GM	10	MG	100	06/01/2015	99/99/9999							
49452-4050-03	J2001			06/01/2015	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL MONOHYDRATE (U.S.P.)	500	GM	BO	NA	GM	10	MG	100	06/01/2015	99/99/9999							
49452-4140-01	J2060			06/01/2015	10/17/2016	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	5	GM	JR	NA	GM	2	MG	500	06/01/2015	10/17/2016							
49452-4140-02	J2060			06/01/2015	10/17/2016	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	25	GM	JR	NA	GM	2	MG	500	06/01/2015	10/17/2016							
49452-4140-03	J2060			06/01/2015	10/17/2016	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	100	GM	JR	NA	GM	2	MG	500	06/01/2015	10/17/2016							
49452-4140-04	J2060			06/01/2015	10/17/2016	INJECTION, LORAZEPAM, 2 MG	LORAZEPAM (U.S.P.)	500	GM	JR	NA	GM	2	MG	500	06/01/2015	10/17/2016							
49452-4300-01	J3475			06/01/2015	10/17/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE HEPTAHYDRATE (U.S.P., E.P., B.P., J.P.)	500	GM	BO	NA	GM	500	MG	2	06/01/2015	10/17/2016							
49452-4300-02	J3475			06/01/2015	10/17/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE HEPTAHYDRATE (U.S.P., E.P., B.P., J.P.)	2500	GM	BO	NA	GM	500	MG	2	06/01/2015	10/17/2016							
49452-4300-03	J3475			06/01/2015	10/17/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE HEPTAHYDRATE (U.S.P., E.P., B.P., J.P.)	12000	GM	BO	NA	GM	500	MG	2	06/01/2015	10/17/2016							
49452-4380-01	J2150			06/01/2015	10/17/2016	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	500	GM	BO	NA	GM	50	ML	0.8	06/01/2015	10/17/2016							
49452-4380-02	J2150			06/01/2015	10/17/2016	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	2500	GM	BO	NA	GM	50	ML	0.8	06/01/2015	10/17/2016							
49452-4380-03	J2150			06/01/2015	10/17/2016	INJECTION, MANNITOL, 25% IN 50 ML	MANNITOL (U.S.P.)	12000	GM	BO	NA	GM	50	ML	0.8	06/01/2015	10/17/2016							
49452-4410-01	J3430			06/01/2015	10/17/2016	INJECTION, PHYTONADIONE (VITAMIN K), PER 1 MG	MENADIOL (U.S.P.)	25	GM	BO	NA	GM	1	MG	1000	06/01/2015	10/17/2016							
49452-4410-02	J3430			06/01/2015	10/17/2016	INJECTION, PHYTONADIONE (VITAMIN K), PER 1 MG	MENADIOL (U.S.P.)	100	GM	BO	NA	GM	1	MG	1000	06/01/2015	10/17/2016							
49452-4553-01	J1230			06/01/2015	10/17/2016	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	5	GM	BO	NA	GM	10	MG	100	06/01/2015	10/17/2016							
49452-4553-02	J1230			06/01/2015	10/17/2016	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	25	GM	BO	NA	GM	10	MG	100	06/01/2015	10/17/2016							
49452-4553-03	J1230			06/01/2015	10/17/2016	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL (U.S.P.)	100	GM	BO	NA	GM	10	MG	100	06/01/2015	10/17/2016							
49452-4686-01	J7509			06/01/2015	10/17/2016	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (U.S.P., MICRONIZED)	1	GM	BO	NA	GM	4	MG	250	06/01/2015	10/17/2016							
49452-4686-02	J7509			06/01/2015	10/17/2016	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (U.S.P., MICRONIZED)	5	GM	BO	NA	GM	4	MG	250	06/01/2015	10/17/2016							
49452-4686-03	J7509			06/01/2015	10/17/2016	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (U.S.P., MICRONIZED)	25	GM	BO	NA	GM	4	MG	250	06/01/2015	10/17/2016							
49452-4688-01	J1030			06/01/2015	10/17/2016	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (U.S.P., MICRONIZED)	1	GM	BO	NA	GM	40	MG	25	06/01/2015	10/17/2016							
49452-4688-02	J1030			06/01/2015	10/17/2016	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (U.S.P., MICRONIZED)	5	GM	BO	NA	GM	40	MG	25	06/01/2015	10/17/2016							
49452-4688-03	J1030			06/01/2015	10/17/2016	INJECTION, METHYLPREDNISOLONE ACETATE, 40 MG	METHYLPREDNISOLONE ACETATE (U.S.P., MICRONIZED)	25	GM	BO	NA	GM	40	MG	25	06/01/2015	10/17/2016							
49452-4715-01	J2765			06/01/2015	99/99/9999	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL MONOHYDRATE (U.S.P.)	10	GM	BO	NA	GM	10	MG	100	06/01/2015	99/99/9999							
49452-4715-02	J2765			06/01/2015	10/17/2016	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL MONOHYDRATE (U.S.P.)	25	GM	BO	NA	GM	10	MG	100	06/01/2015	10/17/2016							
49452-4715-03	J2765			06/01/2015	10/17/2016	INJECTION, METOCLOPRAMIDE HCL, UP TO 10 MG	METOCLOPRAMIDE HCL MONOHYDRATE (U.S.P.)	100	GM	BO	NA	GM	10	MG	100	06/01/2015	10/17/2016							
49452-4726-01	J3490			06/01/2015	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	25	GM	BO	NA	GM	1	GM	1	06/01/2015	99/99/9999							
49452-4726-02	J3490			06/01/2015	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	100	GM	JR	NA	GM	1	GM	1	06/01/2015	10/17/2016							
49452-4726-03	J3490			06/01/2015	99/99/9999	UNCLASSIFIED DRUGS	METRONIDAZOLE (U.S.P.)	500	GM	BO	NA	GM	1	GM	1	06/01/2015	99/99/9999							
49452-4800-01	J2300			06/01/2015	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL	0.1	GM	BO	NA	GM	10	MG	100	06/01/2015	99/99/9999							
49452-4800-02	J2300			06/01/2015	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL	1	GM	BO	NA	GM	10	MG	100	06/01/2015	99/99/9999							
49452-4800-03	J2300			06/01/2015	99/99/9999	INJECTION, NALBUPHINE HYDROCHLORIDE, PER 10 MG	NALBUPHINE HCL	5	GM	BO	NA	GM	10	MG	100	06/01/2015	99/99/9999							
49452-4836-03	J2310			06/01/2015	10/17/2016	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL DIHYDRATE (U.S.P.)	1	GM	JR	NA	GM	1	MG	1000	06/01/2015	10/17/2016							
49452-5000-01	J2440			06/01/2015	10/17/2016	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HCL (U.S.P.)	5	GM	BO	NA	GM	60	MG	16.66666	06/01/2015	10/17/2016							
49452-5000-02	J2440			06/01/2015	10/17/2016	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HCL (U.S.P.)	25	GM	BO	NA	GM	60	MG	16.66666	06/01/2015	10/17/2016							
49452-5000-03	J2440			06/01/2015	10/17/2016	INJECTION, PAPAVERINE HCL, UP TO 60 MG	PAPAVERINE HCL (U.S.P.)	100	GM	BO	NA	GM	60	MG	16.66666	06/01/2015	10/17/2016							
49452-5200-03	J2560			06/01/2015	10/17/2016	INJECTION, PHENOBARBITAL SODIUM, UP TO 120 MG	PHENOBARBITAL SODIUM (U.S.P.)	25	GM	BO	NA	GM	120	MG	8.33333	06/01/2015	10/17/2016							
49452-5217-01	J2760			06/01/2015	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	0.1	GM	BO	NA	GM	5	MG	200	06/01/2015	99/99/9999							
49452-5217-02	J2760			06/01/2015	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	0.5	GM	BO	NA	GM	5	MG	200	06/01/2015	99/99/9999							
49452-5217-05	J2760			06/01/2015	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	5	GM	BO	NA	GM	5	MG	200	06/01/2015	99/99/9999							
49452-5290-01	J7799			06/01/2015	10/17/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	PHENYLEPHRINE HCL (U.S.P.)	5	GM	BO	NA	GM	1	GM	1	06/01/2015	10/17/2016							
49452-5290-02	J7799			06/01/2015	10/17/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	PHENYLEPHRINE HCL (U.S.P.)	25	GM	BO	NA	GM	1	GM	1	06/01/2015	10/17/2016							
49452-5290-03	J7799			06/01/2015	10/17/2016	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	PHENYLEPHRINE HCL (U.S.P.)	100	GM	BO	NA	GM	1	GM	1	06/01/2015	10/17/2016							
49452-5770-01	J3480			06/01/2015	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	500	GM	BO	NA	GM	2	MEQ	6.71141	06/01/2015	10/17/2016							
49452-5770-02	J3480			06/01/2015	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	2500	GM	BO	NA	GM	2	MEQ	6.71141	06/01/2015	10/17/2016							
49452-5770-03	J3480			06/01/2015	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	12000	GM	BO	NA	GM	2	MEQ	6.71141	06/01/2015	10/17/2016							
49452-5780-01	J3480			06/01/2015	10/17/2016	INJECTION, POTASSIUM CHLORIDE, PER 2 MEQ	POTASSIUM CHLORIDE (U.S.P.)	500	GM	BO	NA	GM	2	MEQ	6.71141	06/0								

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	NDC Label	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
13533-0700-02		J0256		11/01/2012	99/99/9999	INJECTION, ALPHA 1 PROTEINASE INHIBITOR (HUMAN), NOT OTHERWISE SPECIFIED, 10 MG	PROLASTIN-C (1000MG W/20ML DILUENT) 1 MG	1	EA	VL	IV	EA	10	MG	0.1	11/01/2012	99/99/9999						
13533-0800-40		J1561		10/01/2014	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/GAMMAKED), NON-LYOPHILIZED (E. G. LIQUID), 500 MG	GAMUNEX-C (1X400ML SINGLE-USE) 100 MG/ML	400	ML	VL	IJ	ML	500	MG	0.2	10/01/2014	99/99/9999						
59676-0610-01		J9999		10/23/2015	99/99/9999	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	YONDELIS (PF,LYOPHILIZED) 1 MG	1	EA	VL	IV	EA	1	MG	1	10/23/2015	99/99/9999						
17478-0174-24	KO	J7614	KO	10/20/2015	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	10/20/2015	99/99/9999						
24492-0899-99	KO	J7682	KO	11/01/2015	02/16/2016	TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN (PAK PF) 300 MG/5 ML	5	ML	PC	IH	ML	300	MG	0.2	11/01/2015	02/16/2016						
49452-6061-05		J2675		06/01/2015	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,YAM,MICRONIZED)	1000	GM	JR	NA	GM	50	MG	20	06/01/2015	99/99/9999						
49452-6080-02		J2675		06/01/2015	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE/U.S.P.)	25	GM	BO	NA	GM	50	MG	20	06/01/2015	99/99/9999						
49452-6080-03		J2675		06/01/2015	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE/U.S.P.)	100	GM	BO	NA	GM	50	MG	20	06/01/2015	99/99/9999						
49452-6080-06		J2675		09/01/2015	10/17/2016	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE/U.S.P.)	500	GM	BO	NA	GM	50	MG	20	09/01/2015	10/17/2016						
49452-6087-01		J2550		06/01/2015	10/17/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	25	GM	BO	NA	GM	50	MG	20	06/01/2015	10/17/2016						
49452-6087-02		J2550		06/01/2015	10/17/2016	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (U.S.P.)	100	GM	BO	NA	GM	50	MG	20	06/01/2015	10/17/2016						
49452-6089-02		J1800		06/01/2015	10/17/2016	INJECTION, PROPRANLOL HCL, UP TO 1 MG	PROPRANLOL HCL (U.S.P.)	5	GM	BO	NA	GM	1	MG	1000	06/01/2015	10/17/2016						
49452-6089-03		J1800		06/01/2015	99/99/9999	INJECTION, PROPRANLOL HCL, UP TO 1 MG	PROPRANLOL HCL (U.S.P.)	25	GM	BO	NA	GM	1	MG	1000	06/01/2015	99/99/9999						
49452-6140-01		J3415		06/01/2015	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	25	GM	BO	NA	GM	100	MG	10	06/01/2015	99/99/9999						
49452-6140-02		J3415		06/01/2015	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	100	GM	BO	NA	GM	100	MG	10	06/01/2015	99/99/9999						
49452-6140-03		J3415		06/01/2015	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL (U.S.P.)	1000	GM	BO	NA	GM	100	MG	10	06/01/2015	99/99/9999						
49452-7660-01		J1071		06/01/2015	10/17/2016	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE (U.S.P.)	5	GM	BO	NA	GM	1	MG	1000	06/01/2015	10/17/2016						
49452-7660-02		J1071		06/01/2015	10/17/2016	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE (U.S.P.)	25	GM	BO	NA	GM	1	MG	1000	06/01/2015	10/17/2016						
49452-7660-03		J1071		06/01/2015	10/17/2016	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE (U.S.P.)	100	GM	BO	NA	GM	1	MG	1000	06/01/2015	10/17/2016						
49452-7720-01		J2810		06/01/2015	10/17/2016	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE ANHYDROUS (U.S.P.)	100	GM	BO	NA	GM	40	MG	25	06/01/2015	10/17/2016						
49452-7720-02		J2810		06/01/2015	10/17/2016	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE ANHYDROUS (U.S.P.)	500	GM	BO	NA	GM	40	MG	25	06/01/2015	10/17/2016						
49452-7910-01		J3302		06/01/2015	10/17/2016	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (U.S.P.,MICRONIZED)	1	GM	BO	NA	GM	5	MG	200	06/01/2015	10/17/2016						
49452-7910-02		J3302		06/01/2015	10/17/2016	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (U.S.P.,MICRONIZED)	5	GM	BO	NA	GM	5	MG	200	06/01/2015	10/17/2016						
49452-7910-03		J3302		06/01/2015	10/17/2016	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (U.S.P.,MICRONIZED)	10	GM	BO	NA	GM	5	MG	200	06/01/2015	10/17/2016						
49452-7924-01		J3250		06/01/2015	10/17/2016	INJECTION, TRIMETHOBENZAMIDE HCL, UP TO 200 MG	TRIMETHOBENZAMIDE HCL (U.S.P.)	5	GM	BO	NA	GM	200	MG	5	06/01/2015	10/17/2016						
49452-7924-02		J3250		06/01/2015	10/17/2016	INJECTION, TRIMETHOBENZAMIDE HCL, UP TO 200 MG	TRIMETHOBENZAMIDE HCL (U.S.P.)	25	GM	BO	NA	GM	200	MG	5	06/01/2015	10/17/2016						
49452-8070-01		J3350		06/01/2015	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P.,J.P.)	500	GM	BO	NA	GM	40	GM	0.025	06/01/2015	99/99/9999						
49452-8070-02		J3350		06/01/2015	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P.,J.P.)	2500	GM	BO	NA	GM	40	GM	0.025	06/01/2015	99/99/9999						
49452-8070-03		J3350		06/01/2015	99/99/9999	INJECTION, UREA, UP TO 40 GM	UREA (U.S.P.,J.P.)	12000	GM	BO	NA	GM	40	GM	0.025	06/01/2015	99/99/9999						
00603-6330-20		J8499		11/18/2014	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	VALGANCICLOVIR HYDROCHLORIDE (USP,FILM-COATED) 450 MG	60	EA	BO	PO	EA	1	MG	1	11/18/2014	99/99/9999						
68992-3010-01		J7503		01/01/2016	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ENVARUSUS XR), ORAL, 0.25 MG	ENVARUSUS XR 1 MG	100	EA	BO	PO	EA	0.25	MG	4	01/01/2016	99/99/9999						
68992-3010-03		J7503		01/01/2016	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ENVARUSUS XR), ORAL, 0.25 MG	ENVARUSUS XR 1 MG	30	EA	BO	PO	EA	0.25	MG	4	01/01/2016	99/99/9999						
68992-3040-01		J7503		01/01/2016	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ENVARUSUS XR), ORAL, 0.25 MG	ENVARUSUS XR 4 MG	100	EA	BO	PO	EA	0.25	MG	16	01/01/2016	99/99/9999						
68992-3040-03		J7503		01/01/2016	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ENVARUSUS XR), ORAL, 0.25 MG	ENVARUSUS XR 4 MG	30	EA	BO	PO	EA	0.25	MG	16	01/01/2016	99/99/9999						
68992-3075-01		J7503		01/01/2016	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ENVARUSUS XR), ORAL, 0.25 MG	ENVARUSUS XR 0.75 MG	100	EA	BO	PO	EA	0.25	MG	3	01/01/2016	99/99/9999						
68992-3075-03		J7503		01/01/2016	99/99/9999	TACROLIMUS, EXTENDED RELEASE, (ENVARUSUS XR), ORAL, 0.25 MG	ENVARUSUS XR 0.75 MG	30	EA	BO	PO	EA	0.25	MG	3	01/01/2016	99/99/9999						
49452-0011-01		J3490		06/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	5	GM	BO	NA	GM	1	GM	1	06/01/2015	99/99/9999						
49452-0011-02		J3490		06/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	25	GM	BO	NA	GM	1	GM	1	06/01/2015	99/99/9999						
49452-0011-03		J3490		06/01/2015	99/99/9999	UNCLASSIFIED DRUGS	TESTOSTERONE PROPIONATE (U.S.P.,MICRONIZED)	100	GM	BO	NA	GM	1	GM	1	06/01/2015	99/99/9999						
49452-3543-02		J3490		06/01/2015	10/17/2016	UNCLASSIFIED DRUGS	HYALURONIC ACID	1	GM	BO	NA	GM	1	GM	1	06/01/2015	10/17/2016						
49452-5980-01		J7510		06/01/2015	10/17/2016	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (U.S.P.,MICRONIZED)	5	GM	BO	NA	GM	5	MG	200	06/01/2015	10/17/2016						
49452-5980-02		J7510		06/01/2015	10/17/2016	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (U.S.P.,MICRONIZED)	25	GM	BO	NA	GM	5	MG	200	06/01/2015	10/17/2016						
49452-5980-03		J7510		06/01/2015	10/17/2016	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE (U.S.P.,MICRONIZED)	100	GM	BO	NA	GM	5	MG	200	06/01/2015	10/17/2016						
49452-6000-01		J7506		06/01/2015	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,ANH,MICRONIZED)	5	GM	BO	NA	GM	5	MG	200	06/01/2015	12/31/2015						
49452-6000-02		J7506		06/01/2015	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,ANH,MICRONIZED)	25	GM	BO	NA	GM	5	MG	200	06/01/2015	12/31/2015						
49452-6000-03		J7506		06/01/2015	12/31/2015	PREDNISONE, ORAL, PER 5MG	PREDNISONE (U.S.P.,ANH,MICRONIZED)	100	GM	BO	NA	GM	5	MG	200	06/01/2015	12/31/2015						
49452-6061-02		J2675		06/01/2015	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,YAM,MICRONIZED)	25	GM	JR	NA	GM	50	MG	20	06/01/2015	99/99/9999						
49452-6061-03		J2675		06/01/2015	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,YAM,MICRONIZED)	100	GM	JR	NA	GM	50	MG	20	06/01/2015	99/99/9999						
49452-6061-04		J2675		06/01/2015	99/99/9999	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (U.S.P.,YAM,MICRONIZED)	500	GM	JR	NA	GM	50	MG	20	06/01/2015	99/99/9999						
63323-0400-05		J1953		11/13/2015	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (SINGLE USE,LATEX-FREE) 100 MG/1 ML	5	ML	VL	IV	ML	10	MG									

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Suits of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00641-1617-10		J0278		12/02/2015	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE (10X2ML) 250 MG/1 ML	2	ML	VL	IJ	ML	100	MG	2.5	12/02/2015	99/99/9999						
60505-0834-01		J0692		11/02/2015	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME 1 GM	1	EA	VL	IJ	EA	500	MG	2	11/02/2015	99/99/9999						
60505-0681-01		J0692		11/02/2015	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME 2 GM	1	EA	VL	IJ	EA	500	MG	4	11/02/2015	99/99/9999						
00143-9935-01		J0698		11/19/2015	08/23/2018	INJECTION, CEFOTAXIME SODIUM, PER GM	CEFOTAXIME (USP,PHARMACY BULK) 10 GM	1	EA	VL	IV	EA	1	GM	10	11/19/2015	08/23/2018						
67457-0595-08		J1652		11/13/2015	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PF) 10 MG/0.8 ML	0.8	ML	SR	SC	ML	0.5	MG	25	11/13/2015	99/99/9999						
00641-6132-25		J2310		11/09/2015	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL 0.4 MG/1 ML	1	ML	VL	IJ	ML	1	MG	0.4	11/09/2015	99/99/9999						
00781-3344-95		J2543		11/10/2015	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE USE PF) 2 GM-0.25 GM	10	EA	VL	IV	EA	1.125	GM	2	11/10/2015	99/99/9999						
00781-3367-95		J2543		11/10/2015	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE USE PF) 4 GM-0.5 GM	10	EA	VL	IV	EA	1.125	GM	4	11/10/2015	99/99/9999						
00143-9564-10		J2760		11/04/2015	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (LYOPHILIZED) 5 MG	10	EA	VL	IJ	EA	5	MG	1	11/04/2015	99/99/9999						
00074-3109-32		J7502		11/10/2015	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	GENGRAF (BLISTER PACK) 100 MG	30	EA	BX	PO	EA	100	MG	1	11/10/2015	99/99/9999						
00074-3108-32		J7515		12/08/2015	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	GENGRAF (BLISTER PACK) 25 MG	30	EA	BX	PO	EA	25	MG	1	12/08/2015	99/99/9999						
16714-0467-01		None		01/01/2016	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM COATED) 150 MG	60	EA	BO	PO	EA	150	MG	1	01/01/2016	99/99/9999						
16714-0468-01		None		01/01/2016	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (USP,FILM COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	01/01/2016	99/99/9999						
00597-0145-60		J8499		10/16/2014	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	OFEV 100 MG	60	EA	BO	PO	EA	1	EA	1	10/16/2014	99/99/9999						
00487-0201-03		J7620		01/01/2008	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (30X3ML)	3	ML	PC	IH	ML	3	MG	0.33333	01/01/2008	99/99/9999						
00597-0145-60		J8499		10/16/2014	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	OFEV 150 MG	60	EA	BO	PO	EA	1	EA	1	10/16/2014	99/99/9999						
49452-6000-01		J7512		01/01/2016	10/17/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.,ANH,MICRONIZED)	5	GM	BO	NA	GM	1	MG	1000	01/01/2016	10/17/2016						
49452-6000-02		J7512		01/01/2016	10/17/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.,ANH,MICRONIZED)	25	GM	BO	NA	GM	1	MG	1000	01/01/2016	10/17/2016						
49452-6000-03		J7512		01/01/2016	10/17/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE (U.S.P.,ANH,MICRONIZED)	100	GM	BO	NA	GM	1	MG	1000	01/01/2016	10/17/2016						
43598-0409-25		J7614		09/16/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (5X5,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83332	09/16/2014	99/99/9999						
43598-0409-25	KO	J7614	KO	09/16/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (5X5,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83332	09/16/2014	99/99/9999						
68992-3010-01		J7508		09/01/2015	12/31/2015	TACROLIMUS, EXTENDED RELEASE, ORAL, 0.1 MG	ENVARSUS XR 1 MG	0.1	EA	BO	PO	EA	0.1	MG	10	09/01/2015	12/31/2015						
68992-3010-03		J7508		09/01/2015	12/31/2015	TACROLIMUS, EXTENDED RELEASE, ORAL, 0.1 MG	ENVARSUS XR 1 MG	30	EA	BO	PO	EA	0.1	MG	10	09/01/2015	12/31/2015						
68992-3040-01		J7508		09/01/2015	12/31/2015	TACROLIMUS, EXTENDED RELEASE, ORAL, 0.1 MG	ENVARSUS XR 4 MG	100	EA	BO	PO	EA	0.1	MG	40	09/01/2015	12/31/2015						
68992-3040-03		J7508		09/01/2015	12/31/2015	TACROLIMUS, EXTENDED RELEASE, ORAL, 0.1 MG	ENVARSUS XR 4 MG	30	EA	BO	PO	EA	0.1	MG	40	09/01/2015	12/31/2015						
68992-3075-01		J7508		09/01/2015	12/31/2015	TACROLIMUS, EXTENDED RELEASE, ORAL, 0.1 MG	ENVARSUS XR 0.75 MG	100	EA	BO	PO	EA	0.1	MG	7.5	09/01/2015	12/31/2015						
68992-3075-03		J7508		09/01/2015	12/31/2015	TACROLIMUS, EXTENDED RELEASE, ORAL, 0.1 MG	ENVARSUS XR 0.75 MG	30	EA	BO	PO	EA	0.1	MG	7.5	09/01/2015	12/31/2015						
00173-0821-02		J9302		01/05/2016	02/10/2016	INJECTION, OFATUMUMAB, 10 MG	ARZERRA (PF,LATEX-FREE) 20 MG/1 ML	5	ML	VL	IV	ML	10	MG	2	01/05/2016	02/10/2016						
70020-1911-01		J9207		01/01/2016	99/99/9999	INJECTION, IXABEPILONE, 1 MG	IXEMPRA (W/DILUENT) 45 MG	1	EA	VL	IV	EA	1	MG	45	01/01/2016	99/99/9999						
70020-1910-01		J9207		01/01/2016	99/99/9999	INJECTION, IXABEPILONE, 1 MG	IXEMPRA (W/DILUENT) 15 MG	1	EA	VL	IV	EA	1	MG	15	01/01/2016	99/99/9999						
55150-0220-99		J1327		12/14/2015	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (PF,LATEX-FREE) 2 MG/1 ML	100	ML	VL	IV	ML	5	MG	0.4	12/14/2015	99/99/9999						
55150-0218-99		J1327		12/14/2015	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (PF,LATEX-FREE) 0.75 MG/1 ML	100	ML	VL	IV	ML	5	MG	0.15	12/14/2015	99/99/9999						
55150-0219-10		J1327		12/14/2015	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (PF,LATEX-FREE) 2 MG/1 ML	10	ML	VL	IV	ML	5	MG	0.4	12/14/2015	99/99/9999						
00703-1179-01		J1327		12/11/2015	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE 0.75 MG/1 ML	100	ML	VL	IV	ML	5	MG	0.15	12/11/2015	99/99/9999						
63481-0367-06		J3030		11/09/2015	04/13/2018	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	SUMAVEL DOSEPRO 6 MG/0.5 ML	0.5	ML	SR	SC	ML	6	MG	2	11/09/2015	04/13/2018						
17478-0173-24		J7614		12/15/2015	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	12/15/2015	99/99/9999						
63323-0750-20		J9263		12/17/2015	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (SINGLE-USE VIAL; USP,PF) 5 MG/1 ML	20	ML	VL	IV	ML	0.5	MG	10	12/17/2015	99/99/9999						
43598-0410-25		J7614		09/16/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (5X5,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	09/16/2014	99/99/9999						
76204-0600-30		J7620		09/03/2015	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (30 VIALS X 1 POUCH) 3MG/3ML-0.5MG/3ML	3	ML	PC	IH	ML	3	MG	0.33333	09/03/2015	99/99/9999						
17478-0173-24	KO	J7614	KO	12/15/2015	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX (PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	12/15/2015	99/99/9999						
43598-0410-25	KO	J7614	KO	09/16/2014	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (5X5,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	09/16/2014	99/99/9999						
00078-0690-61		J9302		02/11/2016	99/99/9999	INJECTION, OFATUMUMAB, 10 MG	ARZERRA (SINGLE-USE W/2 FILTERS) 20 MG/1 ML	50	ML	VL	IV	ML	10	MG	2	02/11/2016	99/99/9999						
00078-0669-13		J9302		02/11/2016	99/99/9999	INJECTION, OFATUMUMAB, 10 MG	ARZERRA (SINGLE-USE W/2 FILTERS) 20 MG/1 ML	5	ML	VL	IV	ML	10	MG	2	02/11/2016	99/99/9999						
00078-0669-61		J9302		02/11/2016	99/99/9999	INJECTION, OFATUMUMAB, 10 MG	ARZERRA (PF,LATEX-FREE) 20 MG/1 ML	5	ML	VL	IV	ML	10	MG	2	02/11/2016	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00781-9261-95		J0290		12/10/2015	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	PREMIERPRO RX AMPICILLIN 1 GM	10 EA	VL U EA	EA			500 MG		2	12/10/2015	99/99/9999						
00781-9250-95		J0290		12/10/2015	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	PREMIERPRO RX AMPICILLIN 500 MG	10 EA	VL U EA	EA			500 MG		1	12/10/2015	99/99/9999						
00781-9242-95		J0290		12/10/2015	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	PREMIERPRO RX AMPICILLIN 250 MG	10 EA	VL U EA	EA			500 MG		0.5	12/10/2015	99/99/9999						
00781-9273-95		J0290		12/10/2015	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	PREMIERPRO RX AMPICILLIN 2 GM	10 EA	VL U EA	EA			500 MG		4	12/10/2015	99/99/9999						
00409-4684-12	J1450			12/29/2015	09/01/2017	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (LATEX-FREE) 400 MG/200 ML	200 ML	FC IV ML	ML			200 MG		0.01	12/29/2015	09/01/2017						
00409-4688-12	J1450			12/29/2015	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE 400 MG/200 ML	200 ML	FC IV ML	ML			200 MG		0.01	12/29/2015	99/99/9999						
67457-0594-06	J1652			02/11/2016	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	ARIXTRA (PREFL.27GX1/2",PF) 7.5 MG/0.6 ML	0.6 ML	SR SC ML	ML			0.5 MG		25	02/11/2016	99/99/9999						
60505-6097-00	J1740			01/15/2016	99/99/9999	INJECTION, IBANDRONATE SODIUM, 1 MG	IBANDRONATE SODIUM 1 MG/1 ML	3 ML	SR IV ML	ML			1 MG		1	01/15/2016	99/99/9999						
00002-8824-27	J1815			02/29/2016	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMULIN R CONCENTRATED U-500 KWIKPEN 500 U/1 ML	3 ML	SR SC ML	ML			5 U		100	02/29/2016	99/99/9999						
39822-0353-06	J2010			02/01/2016	99/99/9999	INJECTION, LINCOMYCIN HCL, UP TO 300 MG	LINCOMYCIN HCL 300 MG/1 ML	10 ML	VL U ML	ML			300 MG		1	02/01/2016	99/99/9999						
39822-0350-02	J2010			02/01/2016	99/99/9999	INJECTION, LINCOMYCIN HCL, UP TO 300 MG	LINCOMYCIN HCL 300 MG/1 ML	2 ML	VL U ML	ML			300 MG		1	02/01/2016	99/99/9999						
63323-0284-21	J3370			01/22/2016	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF,LATEX-FREE) 1 GM	10 EA	VL IV EA	EA			500 MG		2	01/22/2016	99/99/9999						
00378-9671-30	J7620			01/28/2016	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (30X3ML S VIALS/POUCH)	3 ML	PC IH ML	ML			3 MG		0.33333	01/28/2016	99/99/9999						
49452-2588-04	J1212			09/01/2015	99/99/9999	INJECTION, DMSO, DIMETHYL SULFOXIDE, 50%, 50 ML	DIMETHYL SULFOXIDE (U.S.P.)	100 ML	BO NA ML	ML			50 ML		0.02	09/01/2015	99/99/9999						
00944-2884-01	J0257			10/11/2010	99/99/9999	INJECTION, ALPHA 1 PROTEINASE INHIBITOR (HUMAN), (GLASSIA), 10 MG	GLASSIA (APRX 1000MG/50ML SOLN) 1 MG	1 EA	VL IV EA	EA			10 MG		0.1	10/11/2010	99/99/9999						
76204-0600-60	J7620			09/03/2015	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (30 VIALS X 2 POUCHES) 3MG/3ML-0.5MG/3ML	3 ML	PC IH ML	ML			3 MG		0.33333	09/03/2015	99/99/9999						
00781-7157-29	J7644			09/09/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML,PF) 0.02%	2.5 ML	PC IH ML	ML			1 ML		0.2	09/09/2011	99/99/9999						
00781-7157-29	KO J7644 KO			09/09/2011	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (60X2.5ML,PF) 0.02%	2.5 ML	PC IH ML	ML			1 ML		0.2	09/09/2011	99/99/9999						
55150-0238-05	J1100			02/19/2016	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (USP, MDV,LATEX-FREE) 4 MG/1 ML	5 ML	VL U ML	ML			1 MG		4	02/19/2016	99/99/9999						
55150-0239-30	J1100			02/19/2016	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (USP, MDV,LATEX-FREE) 4 MG/1 ML	30 ML	VL U ML	ML			1 MG		4	02/19/2016	99/99/9999						
55150-0237-01	J1100			02/19/2016	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (USP, SDV,LATEX-FREE) 4 MG/1 ML	1 ML	VL U ML	ML			1 MG		4	02/19/2016	99/99/9999						
00409-4688-18	J1450			12/18/2015	99/99/9999	INJECTION FLUCONAZOLE, 200 MG	FLUCONAZOLE (LATEX-FREE) 200 MG/100 ML	100 ML	FC IV ML	ML			200 MG		0.01	12/18/2015	99/99/9999						
60429-0378-01	J7507			02/10/2016	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100 EA	BO PO EA	EA			1 MG		1	02/10/2016	99/99/9999						
60429-0377-01	J7507			02/10/2016	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 0.5 MG	100 EA	BO PO EA	EA			1 MG		0.5	02/10/2016	99/99/9999						
60429-0379-01	J7507			02/10/2016	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	100 EA	BO PO EA	EA			1 MG		5	02/10/2016	99/99/9999						
00378-9671-60	J7620			03/03/2016	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (STERILE (60X3ML)) 3 MG/3 ML-0.5 MG/3 ML	3 ML	PC IH ML	ML			3 MG		0.33333	03/03/2016	99/99/9999						
00591-3817-66	J7620			02/25/2016	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	3 ML	PC IH ML	ML			3 MG		0.33333	02/25/2016	99/99/9999						
00591-3817-39	J7620			02/25/2016	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	3 ML	PC IH ML	ML			3 MG		0.33333	02/25/2016	99/99/9999						
00093-6817-73	J7626			03/09/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED) 1 MG/2 ML	2 ML	PC IH ML	ML			0.5 MG		1	03/09/2016	99/99/9999						
00143-1473-10	J7512			03/01/2016	06/15/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	1000 EA	BO PO EA	EA			1 MG		10	03/01/2016	06/15/2016						
00143-1477-01	J7512			03/01/2016	06/15/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	100 EA	BO PO EA	EA			1 MG		20	03/01/2016	06/15/2016						
00143-1477-05	J7512			03/01/2016	06/15/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	500 EA	BO PO EA	EA			1 MG		20	03/01/2016	06/15/2016						
00143-1477-10	J7512			03/01/2016	06/15/2016	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	1000 EA	BO PO EA	EA			1 MG		20	03/01/2016	06/15/2016						
49452-0001-04	J0133			09/01/2015	99/99/9999	INJECTION, ACYCLOVIR, 5 MG	ACYCLOVIR (U.S.P.)	100 GM	BO NA GM	GM			5 MG		200	09/01/2015	99/99/9999						
49452-0027-04	J0745			09/01/2015	10/17/2016	INJECTION, CODEINE PHOSPHATE, PER 30 MG	CODEINE PHOSPHATE (U.S.P.)	100 GM	BO NA GM	GM			30 MG		33.33333	09/01/2015	10/17/2016						
49452-0073-03	J0270			09/01/2015	10/17/2016	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	0.1 GM	BO NA GM	GM			1.25 MCG		800000	09/01/2015	10/17/2016						
49452-0073-04	J0270			09/01/2015	10/17/2016	INJECTION, ALPROSTADIL, 1.25 MCG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ALPROSTADIL (U.S.P.)	0.025 GM	BO NA GM	GM			1.25 MCG		800000	09/01/2015	10/17/2016						
49452-0409-01	J3490			09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	25 GM	BO NA GM	GM			1 EA		1	09/01/2015	10/17/2016						
49452-0409-02	J3490			09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	100 GM	BO NA GM	GM			1 EA		1	09/01/2015	10/17/2016						
49452-0409-03	J3490			09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	500 GM	BO NA GM	GM			1 EA		1	09/01/2015	10/17/2016						
49452-0409-04	J3490			09/01/2015	99/99/9999	UNCLASSIFIED DRUGS	AMINOCAPROIC ACID (U.S.P.)	2500 GM	BO NA GM	GM			1 EA		1	09/01/2015	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
49452-0783-01		J7501		09/01/2015	10/17/2016	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE (U.S.P.)	1	GM	BO	NA	GM	100 MG		10	09/01/2015	10/17/2016						
49452-1072-03		J3490		09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	BETAMETHASONE ACETATE MICRONIZED (U.S.P.)	5	GM	BO	NA	GM	1 EA		1	09/01/2015	10/17/2016						
49452-1309-04		J0945		09/01/2015	10/17/2016	INJECTION, BROMPHENIRAMINE MALEATE, PER 10 MG	BROMPHENIRAMINE MALEATE (U.S.P.)	5	GM	BO	NA	GM	10 MG		100	09/01/2015	10/17/2016						
49452-1309-05		J0945		09/01/2015	10/17/2016	INJECTION, BROMPHENIRAMINE MALEATE, PER 10 MG	BROMPHENIRAMINE MALEATE (U.S.P.)	100	GM	BO	NA	GM	10 MG		100	09/01/2015	10/17/2016						
49452-2147-04		J0735		09/01/2015	10/17/2016	INJECTION, CLONIDINE HYDROCHLORIDE, 1 MG	CLONIDINE HCL (U.S.P.)	25	GM	BO	NA	GM	1 MG		1000	09/01/2015	10/17/2016						
49452-2400-04		J3420		09/01/2015	10/17/2016	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (U.S.P.)	25	GM	BO	NA	GM	1000 MCG		1000	09/01/2015	10/17/2016						
49452-2541-02		J1730		09/01/2015	10/17/2016	INJECTION, DIAZOXIDE, UP TO 300 MG	DIAZOXIDE (U.S.P./N.F.)	1	GM	BO	NA	GM	300 MG		3.33333	09/01/2015	10/17/2016						
49452-2541-03		J1730		09/01/2015	10/17/2016	INJECTION, DIAZOXIDE, UP TO 300 MG	DIAZOXIDE (U.S.P./N.F.)	5	GM	BO	NA	GM	300 MG		3.33333	09/01/2015	10/17/2016						
49452-2588-02		J1212		09/01/2015	10/17/2016	INJECTION, DMSO, DIMETHYL SULFOXIDE, 50%, 50 ML	DIMETHYL SULFOXIDE (U.S.P.)	4000	ML	BO	NA	ML	50 %		0.02	09/01/2015	10/17/2016						
49452-6053-01		Q0164		02/01/2016	10/17/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P., N.F.)	5	GM	BO	NA	GM	5 MG		200	02/01/2016	10/17/2016						
49452-6053-02		Q0164		02/01/2016	10/17/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P., N.F.)	25	GM	BO	NA	GM	5 MG		200	02/01/2016	10/17/2016						
49452-6053-03		Q0164		02/01/2016	10/17/2016	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P., N.F.)	100	GM	BO	NA	GM	5 MG		200	02/01/2016	10/17/2016						
49452-6053-05		Q0164		02/01/2016	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE (U.S.P.)	500	GM	BO	NA	GM	5 MG		200	02/01/2016	99/99/9999						
49452-8253-04		J0592		09/01/2015	99/99/9999	INJECTION, BUPRENORPHINE HYDROCHLORIDE, 0.1 MG	BUPRENORPHINE HYDROCHLORIDE (U.S.P.)	5	GM	BO	NA	GM	0.1 MG		10000	09/01/2015	99/99/9999						
00078-0438-15		J8999		04/12/2005	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	GLEEVEC (FILM-COATED) 400 MG	30	EA	BO	PO	EA	1 EA		1	04/12/2005	99/99/9999						
00093-6817-73	KO	J7626	KO	03/09/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (MICRONIZED) 1 MG/2 ML PROMETHAZINE HCL (DOSETTE,VIAL) 25 MG/1 ML	2	ML	PC	IH	ML	0.5 MG		1	03/09/2016	99/99/9999						
00641-0928-25		J2550		12/27/2002	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (DOSETTE,VIAL) 25 MG/1 ML	1	ML	VL	IJ	ML	50 MG		0.5	12/27/2002	99/99/9999						
00641-6173-10		J0500		03/23/2016	99/99/9999	INJECTION, DICYCLIMINE HCL, UP TO 20 MG	DICYCLIMINE 10 MG/1 ML	2	ML	VL	IM	ML	20 MG		0.5	03/23/2016	99/99/9999						
63323-0713-13		J2020		03/25/2016	99/99/9999	INJECTION, LINEZOLID, 200MG	LINEZOLID (LATEX-FREE) 2 MG/1 ML	300	ML	FC	IV	ML	200 MG		0.01	03/25/2016	99/99/9999						
16729-0311-93		J2501		03/15/2016	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (MDV) 0.005 MG/1 ML	2	ML	VL	IV	ML	1 MCG		5	03/15/2016	99/99/9999						
16729-0310-08		J2501		03/15/2016	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (SDV) 0.002 MG/1 ML	1	ML	VL	IV	ML	1 MCG		2	03/15/2016	99/99/9999						
16729-0311-08		J2501		03/15/2016	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (SDV) 0.005 MG/1 ML	1	ML	VL	IV	ML	1 MCG		5	03/15/2016	99/99/9999						
67857-0809-38		J3030		03/17/2016	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ZEMBRACE SYMTOUCH (AUTOINJECTOR) 3 MG/0.5 ML	0.5	ML	SR	SC	ML	6 MG		1	03/17/2016	99/99/9999						
58468-0030-02		J3240		05/01/2016	99/99/9999	INJECTION, THYROTROPIN ALPHA, 0.9 MG, PROVIDED IN 1.1 MG VIAL	THYROGEN (LYOPHILIZED) 1.1 MG	2	EA	VL	IH	EA	1.1 MG		1	05/01/2016	99/99/9999						
76204-0700-24		J7614		04/22/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 0.31 MG/3 ML	3	ML	VL	IH	ML	0.5 MG		0.20666	04/22/2016	99/99/9999						
76204-0900-24		J7614		04/22/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 1.25 MG/3 ML	3	ML	VL	IH	ML	0.5 MG		0.83333	04/22/2016	99/99/9999						
76204-0800-24		J7614		04/22/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 0.63 MG/3 ML	3	ML	VL	IH	ML	0.5 MG		0.42	04/22/2016	99/99/9999						
49452-0029-03		J1170		09/01/2015	10/17/2016	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (U.S.P.)	10	GM	BO	NA	GM	4 MG		250	09/01/2015	10/17/2016						
49452-0031-01		J2175		09/01/2015	10/17/2016	INJECTION, MEPERIDINE HYDROCHLORIDE, PER 100 MG	MEPERIDINE HCL (U.S.P.)	25	GM	BO	NA	GM	100 MG		10	09/01/2015	10/17/2016						
49452-1775-03		J1955		09/01/2015	10/17/2016	INJECTION, LEVOCARNITINE, PER 1 GM	L-CARNITINE FREE BASE	500	GM	BO	NA	GM	1 GM		1	09/01/2015	10/17/2016						
49452-2702-01		J3520		09/01/2015	10/17/2016	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM DIHYDRATE (U.S.P.)	500	GM	BO	NA	GM	150 MG		6.66666	09/01/2015	10/17/2016						
49452-2702-02		J3520		09/01/2015	10/17/2016	EDETATE DISODIUM, PER 150 MG	EDETATE DISODIUM DIHYDRATE (U.S.P.)	2500	GM	BO	NA	GM	150 MG		6.66666	09/01/2015	10/17/2016						
49452-2791-03		J1380		09/01/2015	10/17/2016	INJECTION, ESTRADIOL VALERATE, UP TO 10 MG	ESTRADIOL VALERATE (U.S.P.)	25	GM	BO	NA	GM	10 MG		100	09/01/2015	10/17/2016						
49452-3038-03		J3490		09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	FAMOTIDINE (U.S.P.)	25	GM	BO	NA	GM	1 GM		1	09/01/2015	10/17/2016						
49452-3222-03		J1940		09/01/2015	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (U.S.P./N.F.)	500	GM	BO	NA	GM	20 MG		50	09/01/2015	99/99/9999						
49452-3544-01		J0360		09/01/2015	10/17/2016	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HCL (U.S.P.)	5	GM	BO	NA	GM	20 MG		50	09/01/2015	10/17/2016						
49452-3544-02		J0360		09/01/2015	10/17/2016	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HCL (U.S.P.)	25	GM	BO	NA	GM	20 MG		50	09/01/2015	10/17/2016						
49452-3544-03		J0360		09/01/2015	99/99/9999	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HCL (U.S.P.)	100	GM	BO	NA	GM	20 MG		50	09/01/2015	99/99/9999						
49452-3652-03		J3410		09/01/2015	10/17/2016	INJECTION, HYDROXYZINE HCL, UP TO 25 MG	HYDROXYZINE HCL (U.S.P.)	100	GM	BO	NA	GM	25 MG		40	09/01/2015	10/17/2016						
49452-4836-02		J2310		09/01/2015	10/17/2016	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL DIHYDRATE (U.S.P.)	0.25	GM	BO	NA	GM	1 MG		1000	09/01/2015	10/17/2016						
49452-4836-04		J2310		09/01/2015	10/17/2016	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL DIHYDRATE (U.S.P.)	5	GM	BO	NA	GM	1 MG		1000	09/01/2015	10/17/2016						
49452-4936-01		J2360		09/01/2015	10/17/2016	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORPHENADRINE CITRATE (U.S.P.)	25	GM	BO	NA	GM	60 MG		16.66666	09/01/2015	10/17/2016						
49452-4936-02		J2360		09/01/2015	10/17/2016	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORPHENADRINE CITRATE (U.S.P.)	100	GM	BO	NA	GM	60 MG		16.66666	09/01/2015	10/17/2016						
49452-5217-04		J2760		09/01/2015	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (U.S.P.)	1	GM	BO	NA	GM	5 MG		200	09/01/2015	99/99/9999						
49452-5344-01		J1165		09/01/2015	10/17/2016	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	25	GM	BO	NA	GM	50 MG		20	09/01/2015	10/17/2016						
49452-5344-02		J1165		09/01/2015	10/17/2016	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	100	GM	BO	NA	GM	50 MG		20	09/01/2015	10/17/2016						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
49452-5344-03		J1165		09/01/2015	10/17/2016	INJECTION, PHENYTOIN SODIUM, PER 50 MG	PHENYTOIN SODIUM (U.S.P.)	500	GM	BO	NA	GM	50	MG	20	09/01/2015	10/17/2016							
49452-5390-03		J3430		09/01/2015	10/17/2016	INJECTION, PHYTONADIONE (VITAMIN K), PER 1 MG	PHYTONADIONE (U.S.P.)	25	GM	BO	NA	GM	1	MG	1000	09/01/2015	10/17/2016							
49452-5971-01		J2730		09/01/2015	99/99/9999	INJECTION, PRALIDOXIME CHLORIDE, UP TO 1 GM	PRALIDOXIME CHLORIDE (U.S.P.)	1	GM	BO	NA	GM	1	GM	1	09/01/2015	99/99/9999							
49452-5971-02		J2730		09/01/2015	99/99/9999	INJECTION, PRALIDOXIME CHLORIDE, UP TO 1 GM	PRALIDOXIME CHLORIDE (U.S.P.)	5	GM	BO	NA	GM	1	GM	1	09/01/2015	99/99/9999							
49452-5971-03		J2730		09/01/2015	99/99/9999	INJECTION, PRALIDOXIME CHLORIDE, UP TO 1 GM	PRALIDOXIME CHLORIDE (U.S.P.)	25	GM	BO	NA	GM	1	GM	1	09/01/2015	99/99/9999							
49452-6080-04		J2675		09/01/2015	10/17/2016	INJECTION, PROGESTERONE, PER 50 MG	PROGESTERONE (WETTABLE/U.S.P./PR111)	1000	GM	BO	NA	GM	50	MG	20	09/01/2015	10/17/2016							
49452-6109-01		J2720		09/01/2015	99/99/9999	INJECTION, PROTAMINE SULFATE, PER 10 MG	PROTAMINE SULFATE (U.S.P.)	5	GM	BO	NA	GM	10	MG	100	09/01/2015	99/99/9999							
49452-6109-02		J2720		09/01/2015	99/99/9999	INJECTION, PROTAMINE SULFATE, PER 10 MG	PROTAMINE SULFATE (U.S.P.)	25	GM	BO	NA	GM	10	MG	100	09/01/2015	99/99/9999							
49452-6109-03		J2720		09/01/2015	99/99/9999	INJECTION, PROTAMINE SULFATE, PER 10 MG	PROTAMINE SULFATE (U.S.P.)	100	GM	BO	NA	GM	10	MG	100	09/01/2015	99/99/9999							
49452-6222-04		J3490		09/01/2015	10/17/2016	UNCLASSIFIED DRUGS	RIFAMPIN (U.S.P.)	100	GM	BO	NA	GM	1	GM	1	09/01/2015	10/17/2016							
49452-7720-03		J2810		09/01/2015	10/17/2016	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE ANHYDROUS (U.S.P.)	2500	GM	BO	NA	GM	40	MG	25	09/01/2015	10/17/2016							
49452-7910-04		J3302		09/01/2015	10/17/2016	INJECTION, TRIAMCINOLONE DIACETATE, PER 5MG	TRIAMCINOLONE DIACETATE (MICRONIZED, U.S.P.)	100	GM	BO	NA	GM	5	MG	200	09/01/2015	10/17/2016							
49452-9201-05		J1960		09/01/2015	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	1	GM	BO	NA	GM	2	MG	500	09/01/2015	99/99/9999							
49452-9201-06		J1960		09/01/2015	99/99/9999	INJECTION, LEVORPHANOL TARTRATE, UP TO 2 MG	LEVORPHANOL TARTRATE (U.S.P.)	0.5	GM	BO	NA	GM	2	MG	500	09/01/2015	99/99/9999							
55513-0098-01		J0881		03/16/2015	99/99/9999	INJECTION, DARBEPOETIN ALFA, 1 MCG (NON-ESRD USE)	ARANESP (INNER PACK,PF) 0.01 MG/0.4 ML	0.4	ML	BO	IJ	ML	1	MCG	25	03/16/2015	99/99/9999							
76204-0800-24	KO	J7614	KO	04/22/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 0.63 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.42	04/22/2016	99/99/9999							
76204-0700-24	KO	J7614	KO	04/22/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 0.31 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.20666	04/22/2016	99/99/9999							
76204-0900-24	KO	J7614	KO	04/22/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 1.25 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.83333	04/22/2016	99/99/9999							
55150-0177-05		J1953		04/21/2016	99/99/9999	INJECTION, LEVETIRACETAM, 100 MG	LEVETIRACETAM (LATEX-FREE) 100 MG/1 ML	5	ML	VL	IV	ML	10	MG	10	04/21/2016	99/99/9999							
67457-0291-01		J0360		04/28/2016	99/99/9999	INJECTION, HYDRALAZINE HCL, UP TO 20 MG	HYDRALAZINE HCL (PF) 20 MG/1 ML	1	ML	VL	IJ	ML	20	MG	1	04/28/2016	99/99/9999							
00548-9021-00		J1885		03/01/2016	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/1 ML	1	ML	VL	IJ	ML	15	MG	2	03/01/2016	99/99/9999							
60505-6130-00		J2405		04/28/2016	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON 2 MG/1 ML	2	ML	VL	IJ	ML	1	MG	2	04/28/2016	99/99/9999							
60505-6130-05		J2405		04/28/2016	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON 2 MG/1 ML	2	ML	VL	IJ	ML	1	MG	2	04/28/2016	99/99/9999							
00338-3583-01		J3370		04/18/2016	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL-SODIUM CHLORIDE 0.9%-1 GM	200	ML	VL	IV	ML	500	MG	0.01	04/18/2016	99/99/9999							
00338-3582-01		J3370		05/10/2016	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL-SODIUM CHLORIDE (GALAXY CONTAINER) 0.9%-750 MG/150 ML	150	ML	VL	IV	ML	500	MG	0.01	05/10/2016	99/99/9999							
00338-3581-01		J3370		05/10/2016	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL-SODIUM CHLORIDE (GALAXY CONTAINER) 0.9%-500 MG/100 ML	100	ML	VL	IV	ML	500	MG	0.01	05/10/2016	99/99/9999							
61553-0436-48		J3475		01/01/2016	12/31/2016	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE-SODIUM CHLORIDE (VIAFLEX BAG,PF) 2 GM-0.9% MG/1 ML	100	ML	FC	IV	ML	500	MG	0.04	01/01/2016	12/31/2016							
63323-0642-50		J3475		05/18/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.,PF) 500 MG/1 ML	50	ML	VL	IJ	ML	500	MG	1	05/18/2016	99/99/9999							
63323-0642-20		J3475		05/18/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (S.D.V.,PF) 500 MG/1 ML	20	ML	VL	IJ	ML	500	MG	1	05/18/2016	99/99/9999							
00143-9566-01		J7501		04/21/2016	99/99/9999	AZATHIOPRINE, PARENTERAL, 100 MG	AZATHIOPRINE SODIUM (LYOPHILIZED) 100 MG	1	EA	VL	IV	EA	100	MG	1	04/21/2016	99/99/9999							
17478-0172-24		J7614		04/21/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX PEDIATRIC (PF) 0.31 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.20666	04/21/2016	99/99/9999							
45963-0623-57		J9201		04/12/2016	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 38 MG/1 ML	5.26	ML	VL	IV	ML	200	MG	0.19	04/12/2016	99/99/9999							
45963-0624-58		J9201		04/12/2016	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 38 MG/1 ML	26.3	ML	VL	IV	ML	200	MG	0.19	04/12/2016	99/99/9999							
45963-0636-60		J9201		04/12/2016	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 38 MG/1 ML	52.6	ML	VL	IV	ML	200	MG	0.19	04/12/2016	99/99/9999							
16729-0035-15		J8999		02/08/2011	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOT OTHERWISE SPECIFIED	ANASTROZOLE (FILM-COATED) 1 MG	90	EA	BO	PO	EA	1	MG	1	02/08/2011	99/99/9999							
69656-0101-02		Q9981		07/01/2016	12/31/2016	ROLAPITANT, ORAL, 1 MG	VARUBI (FILM COATED) 90 MG	2	EA	DP	PO	EA	1	MG	90	07/01/2016	12/31/2016							
66887-0004-20		J3490		10/31/2014	99/99/9999	UNCLASSIFIED DRUGS	TESTOPEL PELLETS	100	EA	BX	SC	EA	1	EA	1	10/31/2014	99/99/9999							
17478-0172-24	KO	J7614	KO	04/21/2016	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	XOPENEX PEDIATRIC (PF) 0.31 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.20666	04/21/2016	99/99/9999							
25021-0173-04		J0278		06/15/2016	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE 250 MG/1 ML	4	ML	VL	IJ	ML	100	MG	2.5	06/15/2016	99/99/9999							
25021-0173-02		J0278		06/15/2016	99/99/9999	INJECTION, AMIKACIN SULFATE, 100 MG	AMIKACIN SULFATE 250 MG/1 ML	2	ML	VL	IJ	ML	100	MG	2.5	06/15/2016	99/99/9999							
67457-0523-45		J2543		06/02/2016	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE USE,PF) 4 GM-0.5 GM	10	EA	VL	IV	EA	1.125	GM	4	06/02/2016	99/99/9999							
69452-0153-20		J7507		06/10/2016	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 0.5 MG	100	EA	BO	PO	EA	1	MG	0.5	06/10/2016	99/99/9999							
69452-0154-20		J7507		06/10/2016	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 1 MG	100	EA	BO	PO	EA	1	MG	1	06/10/2016	99/99/9999							
69452-0155-20		J7507		06/10/2016	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	100	EA	BO	PO	EA	1	MG	5	06/10/2016	99/99/9999							
68001-0282-25		J9201		06/07/2016	08/27/2018	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (SINGLE-USE,USP) 200 MG	1	EA	VL	IV	EA	200	MG	1	06/07/2016	08/27/2018							
68001-0282-26		J9201		06/07/2016	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (SINGLE-USE,USP) 1 GM	1	EA	VL	IV	EA	200	MG	5	06/07/2016	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
68001-0282-27	J9201			06/07/2016	08/27/2018	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (SINGLE-USE,USP) 2 GM	1 EA	VL	IV	EA		200 MG		10	06/07/2016	08/27/2018						
00008-4990-02	J3243			05/31/2016	08/14/2017	INJECTION, TIGECYCLINE, 1 MG	TYGACIL (SDV,PF) 50 MG	10 EA	VL	IV	EA		1 MG		50	05/31/2016	08/14/2017						
76388-0635-50	J8999			06/22/2012	10/31/2017	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	LEUKERAN (FILM-COATED) 2 MG	50 EA	BO	PO	EA		1 MG		1	06/22/2012	10/31/2017						
55513-0730-01	J0897			11/20/2010	99/99/9999	INJECTION, DENOSUMAB, 1 MG	XGEVA (PF) 120 MG/1.7 ML	1.7 ML	VL	SC	ML		1 MG		70.58823	11/20/2010	99/99/9999						
55513-0710-01	J0897			06/05/2010	99/99/9999	INJECTION, DENOSUMAB, 1 MG	PROLIA (PF) 60 MG/1 ML	1 ML	SR	SC	ML		1 MG		60	06/05/2010	99/99/9999						
00259-1620-01	J0588			01/25/2016	99/99/9999	INJECTION, INCUBOTULINUMTOXIN A, 1 UNIT	XEOMIN (SINGLE-USE,PF) 200 U	1 EA	VL	IM	EA		1 U		200	01/25/2016	99/99/9999						
00703-1165-01	J1327			07/06/2016	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE 2 MG/1 ML	10 ML	VL	IV	ML		5 MG		0.4	07/06/2016	99/99/9999						
44567-0436-24	J1956			07/01/2016	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (NEXCEL PREMIX BAG,PF) 5%-500 MG/100 ML	100 ML	FC	IV	ML		250 MG		0.02	07/01/2016	99/99/9999						
44567-0435-24	J1956			07/01/2016	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (NEXCEL PREMIX BAG,PF) 5%-250 MG/50 ML	50 ML	FC	IV	ML		250 MG		0.02	07/01/2016	99/99/9999						
44567-0437-24	J1956			07/01/2016	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (NEXCEL PREMIX BAG,PF) 5%-750 MG/150 ML	150 ML	FC	IV	ML		250 MG		0.02	07/01/2016	99/99/9999						
00078-0642-61	J2502			01/05/2016	99/99/9999	INJECTION, PASIREOTIDE LONG ACTING, 1 MG	SIGNIFOR LAR (GML VIAL) 40 MG	1 EA	VL	IM	EA		1 MG		40	01/05/2016	99/99/9999						
00078-0641-61	J2502			01/05/2016	99/99/9999	INJECTION, PASIREOTIDE LONG ACTING, 1 MG	SIGNIFOR LAR (GML VIAL) 20 MG	1 EA	VL	IM	EA		1 MG		20	01/05/2016	99/99/9999						
00078-0643-61	J2502			01/05/2016	99/99/9999	INJECTION, PASIREOTIDE LONG ACTING, 1 MG	SIGNIFOR LAR (GML VIAL) 60 MG	1 EA	VL	IM	EA		1 MG		60	01/05/2016	99/99/9999						
67457-0521-22	J2543			06/23/2016	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 2 GM-0.25 GM	10 EA	VL	IV	EA		1.125 GM		2	06/23/2016	99/99/9999						
17478-0081-30	J2795			06/08/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (PF,LATEX-FREE) 5 MG/1 ML	30 ML	VL	IJ	ML		1 MG		5	06/08/2016	99/99/9999						
55150-0223-10	J2800			07/07/2016	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (LATEX-FREE) 100 MG/1 ML	10 ML	VL	IJ	ML		10 ML		0.1	07/07/2016	99/99/9999						
63323-0106-01	J3475			06/03/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (FREEFLEX BAG,LATEX-FREE) 40 MG/1 ML	100 ML	FC	IV	ML		500 MG		0.08	06/03/2016	99/99/9999						
63323-0108-01	J3475			06/03/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (FREEFLEX BAG,LATEX-FREE) 5%-1 GM/100 ML	100 ML	FC	IV	ML		500 MG		0.02	06/03/2016	99/99/9999						
63323-0106-05	J3475			06/03/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (FREEFLEX BAG,LATEX-FREE) 40 MG/1 ML	50 ML	FC	IV	ML		500 MG		0.08	06/03/2016	99/99/9999						
63323-0107-05	J3475			06/03/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (FREEFLEX BAG,LATEX-FREE) 80 MG/1 ML	50 ML	FC	IV	ML		500 MG		0.16	06/03/2016	99/99/9999						
63323-0106-10	J3475			06/03/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (FREEFLEX BAG,LATEX-FREE) 40 MG/1 ML	1000 ML	FC	IV	ML		500 MG		0.08	06/03/2016	99/99/9999						
63323-0106-15	J3475			06/03/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (FREEFLEX BAG,LATEX-FREE) 40 MG/1 ML	500 ML	FC	IV	ML		500 MG		0.08	06/03/2016	99/99/9999						
00591-5442-43	J7512			04/05/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	48 EA	BX	PO	EA		1 MG		10	04/05/2016	99/99/9999						
00591-5052-21	J7512			04/05/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	21 EA	BX	PO	EA		1 MG		5	04/05/2016	99/99/9999						
00591-5052-43	J7512			04/05/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 5 MG	48 EA	BX	PO	EA		1 MG		5	04/05/2016	99/99/9999						
00591-5442-21	J7512			04/05/2016	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	21 EA	BX	PO	EA		1 MG		10	04/05/2016	99/99/9999						
76075-0102-01	J9047			07/14/2016	99/99/9999	INJECTION, CARFILZOMIB, 1 MG	KYPROLIS (LYOPHILIZED) 30 MG	1 EA	VL	IV	EA		1 MG		30	07/14/2016	99/99/9999						
16714-0500-01	J9171			03/14/2016	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL 20 MG/1 ML	4 ML	VL	IV	ML		1 MG		20	03/14/2016	99/99/9999						
16714-0465-01	J9171			03/14/2016	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL 20 MG/1 ML	1 ML	VL	IV	ML		1 MG		20	03/14/2016	99/99/9999						
68001-0284-25	J9206			06/17/2016	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML SINGLE DOSE,PF) 20 MG/1 ML	5 ML	VL	IV	ML		20 MG		1	06/17/2016	99/99/9999						
68001-0284-34	J9206			06/17/2016	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (PF,LATEX-FREE) 20 MG/1 ML	2 ML	VL	IV	ML		20 MG		1	06/17/2016	99/99/9999						
57237-0076-30	Q0162			04/01/2016	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HCL (FILM-COATED) 8 MG	30 EA	BO	PO	EA		1 MG		8	04/01/2016	99/99/9999						
57237-0078-30	Q0162			02/19/2016	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,STRAWBERRY GUARANA) 8 MG	30 EA	BO	PO	EA		1 MG		8	02/19/2016	99/99/9999						
57237-0077-30	Q0162			02/19/2016	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (USP,STRAWBERRY GUARANA) 4 MG	30 EA	BO	PO	EA		1 MG		4	02/19/2016	99/99/9999						
57237-0075-30	Q0162			04/01/2016	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HCL (FILM-COATED) 4 MG	30 EA	BO	PO	EA		1 MG		4	04/01/2016	99/99/9999						
70332-0103-01	Q0163			04/01/2016	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	RAPIDPAQ DICOPANOL (1X150ML) 5 MG/1 ML	150 ML	BO	PO	ML		50 MG		0.1	04/01/2016	99/99/9999						
10702-0003-50	Q0169			06/08/2016	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL (USP) 25 MG	500 EA	BO	PO	EA		12.5 MG		2	06/08/2016	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00003-2188-51		J0129		06/13/2016	99/99/9999	INJECTION, ABATACEPT, 10 MG	ORENCIA CLICKJECT (PF) 125 MG/1 ML	1	ML	SR	SC	ML	10	MG	12.5	06/13/2016	99/99/9999						
00641-0367-25		J1100		04/27/1983	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1MG	DEXAMETHASONE SODIUM PHOSPHATE (VIAL, DOSETTE) 10 MG/1 ML	1	ML	VL	IJ	ML	1	MG	10	04/27/1983	99/99/9999						
55513-0078-01		J9999		10/28/2015	99/99/9999	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	IMLYGIC (PF) 1000000 PFU/1 ML	1	ML	VL	IJ	ML	1	U	1	10/28/2015	99/99/9999						
55513-0079-01		J9999		10/28/2015	99/99/9999	NOT OTHERWISE CLASSIFIED, ANTINEOPLASTIC DRUGS	IMLYGIC (PF) 100000000 PFU/1 ML	1	ML	VL	IJ	ML	1	U	1	10/28/2015	99/99/9999						
55513-0221-01		J2796		08/25/2008	99/99/9999	INJECTION, ROMIPLOSTIM, 10 MICROGRAMS	NPLATE (PF,STERILE, LYOPHILIZED) 250 MCG	1	EA	VL	SC	EA	10	MCG	25	08/25/2008	99/99/9999						
55513-0222-01		J2796		08/25/2008	99/99/9999	INJECTION, ROMIPLOSTIM, 10 MICROGRAMS	NPLATE (PF,STERILE, LYOPHILIZED) 500 MCG	1	EA	VL	SC	EA	10	MCG	50	08/25/2008	99/99/9999						
69097-0173-53		J7620		07/01/2015	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (30X3ML 5 VIALS/POUCH) 3MG/3ML-0.5MG/3ML	3	ML	PC	IH	ML	3	MG	0.33333	07/01/2015	99/99/9999						
76075-0101-01		J9047		07/20/2012	99/99/9999	INJECTION, CARFILZOMIB, 1 MG	KYPROLIS 60 MG	1	EA	VL	IV	EA	1	MG	60	07/20/2012	99/99/9999						
00487-9601-01		J7626		06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .25MG/2ML	30	ML	PC	IH	ML	0.5	MG	0.25	06/13/2016	99/99/9999						
00487-9601-30		J7626		06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .25MG/2ML	30	ML	PC	IH	ML	0.5	MG	0.25	06/13/2016	99/99/9999						
00487-9701-01		J7626		06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .5MG/2ML	30	ML	PC	IH	ML	0.5	MG	0.5	06/13/2016	99/99/9999						
00487-9701-30		J7626		06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .5MG/2ML	30	ML	AM	IH	ML	0.5	MG	0.5	06/13/2016	99/99/9999						
00487-9601-01	KO	J7626	KO	06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .25MG/2ML	30	ML	PC	IH	ML	0.5	MG	0.25	06/13/2016	99/99/9999						
00487-9601-30	KO	J7626	KO	06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .25MG/2ML	30	ML	PC	IH	ML	0.5	MG	0.25	06/13/2016	99/99/9999						
00487-9701-01	KO	J7626	KO	06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .5MG/2ML	30	ML	PC	IH	ML	0.5	MG	0.5	06/13/2016	99/99/9999						
00487-9701-30	KO	J7626	KO	06/13/2016	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30x2mL) .5MG/2ML	30	ML	AM	IH	ML	0.5	MG	0.5	06/13/2016	99/99/9999						
55513-0209-01		J1442		08/08/2000	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN (26GX5/8",PF,SINGLEJECT) 480 MCG/0.8 ML	0.8	ML	SR	IJ	ML	1	MCG	600	08/08/2000	99/99/9999						
55513-0209-10		J1442		08/08/2000	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN (26GX5/8".0.8MLX10,PF) 480 MCG/0.8 ML	0.8	ML	SR	IJ	ML	1	MCG	600	08/08/2000	99/99/9999						
55513-0530-01		J1442		03/17/1997	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN (S.D.V.,PF) 300 MCG/1 ML	1	ML	VL	IJ	ML	1	MCG	300	03/17/1997	99/99/9999						
55513-0530-10		J1442		03/17/1997	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN (SDV,1MLX10,PF) 300 MCG/1 ML	1	ML	VL	IJ	ML	1	MCG	300	03/17/1997	99/99/9999						
55513-0546-01		J1442		03/17/1997	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN (S.D.V.,PF) 480 MCG/1.6 ML	1.6	ML	VL	IJ	ML	1	MCG	300	03/17/1997	99/99/9999						
55513-0546-10		J1442		03/17/1997	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN (SDV,1.6MLX10,PF) 480 MCG/1.6 ML	1.6	ML	VL	IJ	ML	1	MCG	300	03/17/1997	99/99/9999						
55513-0924-01		J1442		08/08/2000	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN ((26GX5/8"),SINGLE-USE) 300 MCG/0.5 ML	0.5	ML	SR	IJ	ML	1	MCG	600	08/08/2000	99/99/9999						
55513-0924-10		J1442		08/08/2000	99/99/9999	INJECTION, FILGRASTIM (G-CSF), EXCLUDES BIOSIMILARS, 1 MICROGRAM	NEUPOGEN (26GX5/8".0.5MLX10,PF) 300 MCG/0.5 ML	0.5	ML	SR	IJ	ML	1	MCG	600	08/08/2000	99/99/9999						
00143-9558-01		J0641		08/01/2016	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (PF,LYOPHILIZED) 50 MG	1	EA	VL	IV	EA	0.5	MG	100	08/01/2016	99/99/9999						
39822-0617-01		J0770		07/01/2016	99/99/9999	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE (LYOPHILIZED CAKE) 150 MG	1	EA	VL	IJ	EA	150	MG	1	07/01/2016	99/99/9999						
00409-6557-01		J1071		07/19/2016	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE (MDV) 100 MG/1 ML	10	ML	VL	IM	ML	1	MG	100	07/19/2016	99/99/9999						
00409-6562-01		J1071		07/19/2016	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE 200 MG/1 ML	1	ML	VL	IM	ML	1	MG	200	07/19/2016	99/99/9999						
00409-6562-20		J1071		07/19/2016	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1MG	TESTOSTERONE CYPIONATE (MDV) 200 MG/1 ML	10	ML	VL	IM	ML	1	MG	200	07/19/2016	99/99/9999						
61553-0242-52		J1170		04/01/2016	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL-SODIUM CHLORIDE (LIFECARE BAG,LATEX-FREE) 1 MG/1 ML 0.9%	100	ML	FC	IV	ML	4	MG	0.25	04/01/2016	99/99/9999						
61553-0915-04		J1644		04/01/2016	03/31/2017	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM-SODIUM CHLORIDE (VIAFLEX BAG,LATEX-FREE) 1000 U/1000 ML-0.9%	1000	ML	FC	IV	ML	1000	U	0.001	04/01/2016	03/31/2017						
00143-9673-25		J1953		07/29/2016	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM 100 MG/1 ML	5	ML	VL	IV	ML	10	MG	10	07/29/2016	99/99/9999						
00781-3433-95		J2020		08/02/2016	99/99/9999	INJECTION, LINEZOLID, 200MG	LINEZOLID (10X300ML BAGS) 2 MG/1 ML	300	ML	FC	IV	ML	200	MG	0.01	08/02/2016	99/99/9999						
39822-5525-03		J2550		08/01/2016	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (25X1ML,USP) 25 MG/1 ML	1	ML	AM	IJ	ML	50	MG	0.5	08/01/2016	99/99/9999						
39822-5550-06		J2550		08/01/2016	99/99/9999	INJECTION, PROMETHAZINE HCL, UP TO 50 MG	PROMETHAZINE HCL (25X1ML,USP) 50 MG/1 ML	1	ML	AM	IJ	ML	50	MG	1	08/01/2016	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00093-2014-12		J3030		07/20/2016	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	SUMATRIPTAN SUCCINATE 6 MG/0.5 ML	0.5	ML	SR	SC	ML	6	MG	2	07/20/2016	99/99/9999						
00093-2013-12		J3030		07/20/2016	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	SUMATRIPTAN SUCCINATE 4 MG/0.5 ML CYANOCOBALAMIN (M.D.V.,25X1ML) 1000 MCG/1 ML	0.5	ML	SR	SC	ML	6	MG	1.33333	07/20/2016	99/99/9999						
70069-0005-10		J3420		07/28/2016	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CAPECITABINE (USP,FILM-COATED) 150 MG	1	ML	VL	IJ	ML	1000	MCG	1	07/28/2016	99/99/9999						
00054-0271-21		None		07/18/2016	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM COATED) 500 MG	60	EA	BO	PO	EA	150	MG	1	07/18/2016	99/99/9999						
00054-0272-23		None		07/18/2016	99/99/9999	CAPECITABINE, 500 MG, ORAL	CYTARABINE (SDV,PF,LATEX-FREE) 20 MG/1 ML	120	EA	BO	PO	EA	500	MG	1	07/18/2016	99/99/9999						
67457-0455-52		J9100		07/22/2016	99/99/9999	INJECTION, CYTARABINE, 100 MG	DOCETAXEL 20 MG/1 ML	5	ML	VL	IJ	ML	100	MG	0.2	07/22/2016	99/99/9999						
00409-0367-01		J9171		07/08/2016	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL 20 MG/1 ML	4	ML	VL	IV	ML	1	MG	20	07/08/2016	99/99/9999						
00409-0366-01		J9171		07/08/2016	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL 20 MG/1 ML	1	ML	VL	IV	ML	1	MG	20	07/08/2016	99/99/9999						
00944-3810-01		J9266		08/16/2016	99/99/9999	INJECTION, PEGASPARGASE, PER SINGLE DOSE VIAL	ONCASPAR (S.D.V. PF) 750 IU/1 ML	5	ML	VL	IJ	ML	1	VL	0.2	08/16/2016	99/99/9999						
61755-0005-02		J0178		11/21/2011	99/99/9999	INJECTION, AFLIBERCEPT, 1 MG	EYLEA (PF) 40 MG/1 ML	0.05	ML	VL	IO	ML	1	MG	40	11/21/2011	99/99/9999						
59676-0320-04		J0885		01/01/2016	99/99/9999	INJECTION, EPOTIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	PROCRIT (MULTIDOSE) 20000 U/ML	1	ML	VL	IJ	ML	1000	U	20	01/01/2016	99/99/9999						
00074-4339-07		J0135		03/19/2009	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (SINGLE-USE PEN; 4X1ML) 40 MG/0.8 ML	4	EA	BX	SC	EA	20	MG	2	03/19/2009	99/99/9999						
00074-6347-02		J0135		10/15/2014	99/99/9999	INJECTION, ADALIMUMAB, 20 MG	HUMIRA (PRE-FILLED SYRINGE,PF) 10 MG/0.2 ML	2	EA	BX	SC	EA	20	MG	0.5	10/15/2014	99/99/9999						
13533-0703-10		J0256		08/31/2016	99/99/9999	INJECTION, ALPHA 1 PROTEINASE INHIBITOR (HUMAN), NOT OTHERWISE SPECIFIED, 10 MG	PROLASTIN-C (1000MG,L.YOPHILIZED) 1 MG	1	EA	VL	IV	EA	10	MG	0.1	08/31/2016	99/99/9999						
67457-0350-10		J0290		09/12/2016	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,CRYSTALLINE) 500 MG	10	EA	VL	IJ	EA	500	MG	1	09/12/2016	99/99/9999						
67457-0404-10		J0290		09/12/2016	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,CRYSTALLINE) 10 GM	1	EA	VL	IV	EA	500	MG	20	09/12/2016	99/99/9999						
67457-0351-10		J0290		09/12/2016	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,CRYSTALLINE) 1 GM	10	EA	VL	IJ	EA	500	MG	2	09/12/2016	99/99/9999						
00143-9552-01		J0640		08/24/2016	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (PF,L.YOPHILIZED) 350 MG	1	EA	VL	IJ	EA	50	MG	7	08/24/2016	99/99/9999						
00703-0125-01		J0878		09/14/2016	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (PF,L.YOPHILIZED) 500 MG	1	EA	VL	IV	EA	1	MG	500	09/14/2016	99/99/9999						
55150-0243-46		J1956		09/01/2016	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (24X50ML, SINGLE-USE,PF) 5%-250 MG/50 ML	50	ML	FC	IV	ML	250	MG	0.02	09/01/2016	99/99/9999						
55150-0244-47		J1956		09/01/2016	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (24X100ML, SINGLE-USE,PF) 5%-500 MG/100 ML	100	ML	FC	IV	ML	250	MG	0.02	09/01/2016	99/99/9999						
55150-0245-52		J1956		09/01/2016	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (24X150ML, SINGLE-USE,PF) 5%-750 MG/150 ML	150	ML	FC	IV	ML	250	MG	0.02	09/01/2016	99/99/9999						
00781-3000-96		J2185		09/12/2016	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM 500 MG	25	EA	VL	IV	EA	100	MG	5	09/12/2016	99/99/9999						
00781-3000-95		J2185		09/12/2016	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM 500 MG	10	EA	VL	IV	EA	100	MG	5	09/12/2016	99/99/9999						
00781-3098-96		J2185		09/12/2016	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM 1 GM	25	EA	VL	IV	EA	100	MG	10	09/12/2016	99/99/9999						
00781-3098-95		J2185		09/12/2016	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM 1 GM	10	EA	VL	IV	EA	100	MG	10	09/12/2016	99/99/9999						
67457-0299-10		J2310		09/14/2016	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL 0.4 MG/1 ML	10	ML	VL	IJ	ML	1	MG	0.4	09/14/2016	99/99/9999						
00143-9890-10		J2405		09/14/2016	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (USP,MULTIDOSE) 2 MG/1 ML	20	ML	VL	IJ	ML	1	MG	2	09/14/2016	99/99/9999						
62847-0001-01		J3095		10/01/2016	99/99/9999	INJECTION, TELEVANICIN, 10 MG	VIBATIV (SDV,PF,L.YOPHILIZED) 750 MG	10	EA	VL	IV	EA	10	MG	75	10/01/2016	99/99/9999						
25208-0001-04		J3246		09/01/2016	99/99/9999	INJECTION, TIROFIBAN HCL, 0.25MG	AGGRASTAT (PF) 0.25 MG/1 ML	15	ML	PC	IV	ML	0.25	MG	1	09/01/2016	99/99/9999						
25208-0002-03		J3246		09/01/2016	99/99/9999	INJECTION, TIROFIBAN HCL, 0.25MG	AGGRASTAT (1X100ML) 0.05 MG/1 ML	100	ML	PC	IV	ML	0.25	mg	0.2	09/01/2016	99/99/9999						
67457-0281-01		J3415		09/01/2016	99/99/9999	INJECTION, PYRIDOXINE HCL, 100 MG	PYRIDOXINE HCL 100 MG/1 ML	1	ML	VL	IJ	ML	100	MG	1	09/01/2016	99/99/9999						
68001-0283-27		J9060		09/12/2016	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (MDV,LATEX-FREE) 1 MG/1 ML	50	ML	VL	IV	ML	10	MG	0.1	09/12/2016	99/99/9999						
68001-0283-32		J9060		09/12/2016	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (MDV,LATEX-FREE) 1 MG/1 ML	100	ML	VL	IV	ML	10	MG	0.1	09/12/2016	99/99/9999						
25021-0215-98		J9190		09/29/2016	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (BULK PACKAGE,PF) 50 MG/1 ML	50	ML	VL	IV	ML	500	MG	0.1	09/29/2016	99/99/9999						
42023-0149-01		J9245		08/24/2016	99/99/9999	INJECTION, MELPHALAN HYDROCHLORIDE, 50 MG	MELPHALAN HYDROCHLORIDE (W/ 10ML DILUENT) 50 MG	1	EA	VL	IV	EA	50	MG	1	08/24/2016	99/99/9999						
59572-0984-01		J9315		09/16/2016	99/99/9999	INJECTION, ROMIDEPSON, 1 MG	ISTODAX (W/DILUENT) 10 MG	1	EA	VL	IV	EA	1	MG	10	09/16/2016	99/99/9999						
38779-1816-05		J2810		08/01/2016	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE ANHYDROUS (USP)	100	GM	BO	NA	GM	40	MG	25	08/01/2016	99/99/9999						
38779-1816-08		J2810		08/01/2016	99/99/9999	INJECTION, THEOPHYLLINE, PER 40 MG	THEOPHYLLINE ANHYDROUS (USP)	500	GM	BO	NA	GM	40	MG	25	08/01/2016	99/99/9999						
66302-0206-03		J7686		01/01/2011	99/99/9999	TREPROSTINIL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 1.74 MG	TYVASO (4X2.9ML) 0.6 MG/1 ML	2.9	ML	PC	IH	ML	1.74	MG	0.34482	01/01/2011	99/99/9999						
66302-0206-03	KO	J7686	KO	01/01/2011	99/99/9999	TREPROSTINIL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, 1.74 MG	TYVASO (4X2.9ML) 0.6 MG/1 ML	2.9	ML	PC	IH	ML	1.74	MG	0.34482	01/01/2011	99/99/9999						
00409-0805-11		J0690		12/15/2015	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN (INNER NDC) 1 GM	1	EA	VL	IJ	EA	500	MG	2	12/15/2015	99/99/9999						
51862-0460-47		J7502		08/03/2016	99/99/9999	CYCLOSPORINE, ORAL, 100 MG	CYCLOSPORINE (USP,SOFT GELATIN) 100 MG	30	EA	BX	PO	EA	100	MG	1	08/03/2016	99/99/9999						
60505-6110-00		J3489		10/04/2013	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SDV) 4 MG/5 ML	5	ML	VL	IV	ML	1	MG	0.8	10/04/2013	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
67457-0352-10		J0290		10/06/2016	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,CRYSTALLINE) 2 GM	10 EA	VL	U	EA		500 MG		4	10/06/2016	99/99/9999						
67457-0353-10		J0290		10/06/2016	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,CRYSTALLINE) 250 MG	10 EA	VL	U	EA		500 MG		0.5	10/06/2016	99/99/9999						
67457-0675-02		J0630		09/16/2016	99/99/9999	INJECTION, CALCITONIN SALMON, UP TO 400 UNITS	MIACALCIN 200 IU/1 ML	2 ML	VL	U	ML		400 IU		0.5	09/16/2016	99/99/9999						
55150-0242-51		J2020		09/26/2016	99/99/9999	INJECTION, LINEZOLID, 200MG	LINEZOLID 2 MG/1 ML	300 ML	FC	IV	ML		200 MG		0.01	09/26/2016	99/99/9999						
70121-1453-07		J2185		10/03/2016	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM (USP) 1 GM	10 EA	VL	IV	EA		100 MG		10	10/03/2016	99/99/9999						
70121-1454-07		J2185		10/03/2016	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM (USP) 500 MG	10 EA	VL	IV	EA		100 MG		5	10/03/2016	99/99/9999						
16729-0297-83		J2405		10/08/2016	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (5X2ML,SINGLE DOSE) 2 MG/1 ML	2 ML	VL	IJ	ML		1 MG		2	10/08/2016	99/99/9999						
16729-0298-05		J2405		10/08/2016	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV) 2 MG/1 ML	20 ML	VL	IJ	ML		1 MG		2	10/08/2016	99/99/9999						
43598-0565-10		J2501		09/16/2016	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (MDV) 0.005 MG/1 ML	2 ML	VL	IV	ML		1 MCG		5	09/16/2016	99/99/9999						
43598-0564-25		J2501		09/16/2016	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (SDV) 0.005 MG/1 ML	1 ML	VL	IV	ML		1 MCG		5	09/16/2016	99/99/9999						
43598-0563-25		J2501		09/16/2016	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (SDV) 0.002 MG/1 ML	1 ML	VL	IV	ML		1 MCG		2	09/16/2016	99/99/9999						
57894-0054-27		J3357		09/27/2016	12/31/2016	USTEKINUMAB, FOR SUBCUTANEOUS INJECTION, 1 MG	STELARA (SDV,PF) 5 MG/1 ML	26 ML	VL	IV	ML		1 MG		5	09/27/2016	12/31/2016						
63323-0203-20		J3370		10/03/2016	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (FLIP TOP VIAL) 750 MG	10 EA	VL	IV	EA		500 MG		1.5	10/03/2016	99/99/9999						
44567-0410-24		J3475		10/24/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	MAGNESIUM SULFATE-DEXTROSE (LATEX-FREE) 5%-1 GM/100 ML	100 ML	FC	IV	ML		500 MG		0.02	10/24/2016	99/99/9999						
70644-0899-99		J7682		10/01/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN INHALATION SOLUTION PAK (PF) 300 MG/5 ML	5 ML	PC	IH	ML		300 MG		0.2	10/01/2016	99/99/9999						
00591-2897-49		J9025		09/16/2016	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (SDV,PF,LATEX-FREE) 100 MG	1 EA	VL	IJ	EA		1 MG		100	09/16/2016	99/99/9999						
25021-0215-99		J9190		09/29/2016	99/99/9999	INJECTION, FLUOROURACIL, 500 MG	FLUOROURACIL (BULK PACKAGE,PF) 50 MG/1 ML	100 ML	VL	IV	ML		500 MG		0.1	09/29/2016	99/99/9999						
00069-0809-01		Q5102		10/17/2016	03/31/2018	INJECTION, INFLIXIMAB, BIOSIMILAR, 10 MG	INFLECTRA (SDV,PF) 100 MG	1 EA	VL	IV	EA		10 MG		10	10/17/2016	03/31/2018						
55150-0259-30		J0132		10/06/2016	99/99/9999	INJECTION, ACETYLCYSTEINE, 100 MG	ACETYLCYSTEINE (SDV; 4X30ML,PF) 200 MG/1 ML	30 ML	VL	IV	ML		100 MG		2	10/06/2016	99/99/9999						
00378-5260-14	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	14 EA	BO	PO	EA		5 MG		1	06/29/2016	99/99/9999						
00378-5260-98	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	5 EA	BO	PO	EA		5 MG		1	06/29/2016	99/99/9999						
00378-5261-14	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14 EA	BO	PO	EA		20 MG		1	06/29/2016	99/99/9999						
00378-5261-98	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5 EA	BO	PO	EA		20 MG		1	06/29/2016	99/99/9999						
00378-5262-14	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14 EA	BO	PO	EA		100 MG		1	06/29/2016	99/99/9999						
00378-5262-98	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5 EA	BO	PO	EA		100 MG		1	06/29/2016	99/99/9999						
00378-5263-14	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	14 EA	BO	PO	EA		20 MG		7	06/29/2016	99/99/9999						
00378-5263-98	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	5 EA	BO	PO	EA		20 MG		7	06/29/2016	99/99/9999						
00378-5264-14	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	14 EA	BO	PO	EA		20 MG		9	06/29/2016	99/99/9999						
00378-5264-98	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	5 EA	BO	PO	EA		20 MG		9	06/29/2016	99/99/9999						
00378-5265-98	None			06/29/2016	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE 250 MG	5 EA	BO	PO	EA		250 MG		1	06/29/2016	99/99/9999						
00944-2850-01		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (1GM,PF,LATEX-FREE) 20% CUVITRU (1GM, INNER PACK NDC,PF) 20%	5 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
00944-2850-02		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (2GM,PF,LATEX-FREE) 20% CUVITRU (2GM, INNER PACK NDC,PF) 20%	10 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
00944-2850-03		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (2GM,PF,LATEX-FREE) 20% CUVITRU (2GM, INNER PACK NDC,PF) 20%	10 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
00944-2850-04		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (4GM,PF,LATEX-FREE) 20% CUVITRU (4GM, INNER PACK NDC,PF) 20%	20 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
00944-2850-05		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (4GM,PF,LATEX-FREE) 20% CUVITRU (4GM, INNER PACK NDC,PF) 20%	20 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
00944-2850-06		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (8GM,PF,LATEX-FREE) 20% CUVITRU (8GM, INNER PACK NDC,PF) 20%	40 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
00944-2850-07		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (8GM,PF,LATEX-FREE) 20% CUVITRU (8GM, INNER PACK NDC,PF) 20%	40 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
00944-2850-08		J7799		09/26/2016	12/31/2017	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	CUVITRU (8GM,PF,LATEX-FREE) 20% CUVITRU (8GM, INNER PACK NDC,PF) 20%	40 ML	VL	SC	ML		1 GM		2	09/26/2016	12/31/2017						
70644-0899-99	KO	J7682	KO	10/01/2016	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG TOBRAMYCIN, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, UNIT DOSE FORM, ADMINISTERED THROUGH DME, PER 300 MILLIGRAMS	TOBRAMYCIN INHALATION SOLUTION PAK (PF) 300 MG/5 ML	5 ML	PC	IH	ML		300 MG		0.2	10/01/2016	99/99/9999						
00781-7515-87	KO	J7626	KO	08/20/2015	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.25 MG/2 ML	2 ML	PC	IH	ML		0.5 MG		0.25	08/20/2015	99/99/9999						
61553-0243-72		J0171		07/01/2016	06/30/2017	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE HCL-SODIUM CHLORIDE (BD SYRINGE,PF) 50 MCG/1 ML-0.9%	10 ML	SR	IV	ML		0.1 MG		0.5	07/01/2016	06/30/2017						
67457-0349-10		J0295		10/31/2016	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM 2 GM-1 GM	10 EA	VL	IJ	EA		1.5 GM		2	10/31/2016	99/99/9999						
00641-6135-25		J0780		10/31/2016	99/99/9999	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE 5 MG/1 ML	2 ML	VL	IJ	ML		10 MG		0.5	10/31/2016	99/99/9999						
00143-9659-01		J1071		11/08/2016	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE 200 MG/1 ML	1 ML	VL	IM	ML		1 MG		200	11/08/2016	99/99/9999						
76329-3399-05		J2690		11/07/2016	99/99/9999	INJECTION, PROCAINAMIDE HCL, UP TO 1 GM	PROCAINAMIDE HCL (LUER-JET, LUER-LOCK) 100 MG/1 ML	10 ML	VL	IJ	ML		1 GM		0.1	11/07/2016	99/99/9999						
55150-0200-10		J2795		10/31/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (SDV,PF,LATEX-FREE) 10 MG/1 ML	10 ML	VL	IJ	ML		1 MG		10	10/31/2016	99/99/9999						
55150-0201-20		J2795		10/31/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (SDV,PF,LATEX-FREE) 10 MG/1 ML	20 ML	VL	IJ	ML		1 MG		10	10/31/2016	99/99/9999						
55150-0195-20		J2795		10/31/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (SDV,PF,LATEX-FREE) 2 MG/1 ML	20 ML	VL	IJ	ML		1 MG		2	10/31/2016	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
55150-0199-20		J2795		10/31/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (SDV,PF,LATEX-FREE) 7.5 MG/1 ML	20	ML	VL	IJ	ML	1	MG	7.5	10/31/2016	99/99/9999							
55150-0198-30		J2795		10/31/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (SDV,PF,LATEX-FREE) 5 MG/1 ML	30	ML	VL	IJ	ML	1	MG	5	10/31/2016	99/99/9999							
55150-0196-99		J2795		10/31/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (SDV,PF,LATEX-FREE) 2 MG/1 ML	100	ML	BO	IJ	ML	1	MG	2	10/31/2016	99/99/9999							
55150-0197-20	J2795			10/31/2016	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (SDV,PF,LATEX-FREE) 5 MG/1 ML	20	ML	VL	IJ	ML	1	MG	5	10/31/2016	99/99/9999							
00904-6574-61	J7509			11/07/2016	01/08/2018	METHYLPREDNISOLONE ORAL, PER 4 MG	METHYLPREDNISOLONE (10X10) 4 MG	100	EA	BX	PO	EA	4	MG	1	11/07/2016	01/08/2018							
43975-0256-05	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	5	EA	BO	PO	EA	20	MG	9	08/02/2016	99/99/9999							
43975-0256-14	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	14	EA	BO	PO	EA	20	MG	9	08/02/2016	99/99/9999							
43975-0255-05	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	5	EA	BO	PO	EA	20	MG	7	08/02/2016	99/99/9999							
43975-0254-14	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14	EA	BO	PO	EA	100	MG	1	08/02/2016	99/99/9999							
43975-0254-05	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5	EA	BO	PO	EA	100	MG	1	08/02/2016	99/99/9999							
43975-0253-14	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14	EA	BO	PO	EA	20	MG	1	08/02/2016	99/99/9999							
43975-0253-05	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5	EA	BO	PO	EA	20	MG	1	08/02/2016	99/99/9999							
43975-0252-14	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	14	EA	BO	PO	EA	5	MG	1	08/02/2016	99/99/9999							
43975-0252-05	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	5	EA	BO	PO	EA	5	MG	1	08/02/2016	99/99/9999							
43975-0257-05	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE 250 MG	5	EA	BO	PO	EA	250	MG	1	08/02/2016	99/99/9999							
43975-0255-14	None			08/02/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	14	EA	BO	PO	EA	20	MG	7	08/02/2016	99/99/9999							
00143-9547-01	J9000			11/04/2016	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 2 MG/1 ML	25	ML	VL	IV	ML	10	MG	0.2	11/04/2016	99/99/9999							
00143-9548-10	J9000			11/04/2016	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 2 MG/1 ML	10	ML	VL	IV	ML	10	MG	0.2	11/04/2016	99/99/9999							
00143-9549-10	J9000			11/04/2016	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (S.D.V.,PF) 2 MG/1 ML	5	ML	VL	IV	ML	10	MG	0.2	11/04/2016	99/99/9999							
00143-9546-01	J9000			11/04/2016	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	ADRIAMYCIN (M.D.V.,PF) 2 MG/1 ML	100	ML	VL	IV	ML	10	MG	0.2	11/04/2016	99/99/9999							
44567-0511-01	J9060			10/17/2016	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (MDV,PF) 1 MG/1 ML	200	ML	VL	IV	ML	10	MG	0.1	10/17/2016	99/99/9999							
45963-0620-60	J9201			10/21/2016	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE HCL (PF,LATEX-FREE) 2 GM	1	EA	VL	IV	EA	200	MG	10	10/21/2016	99/99/9999							
00078-0683-06	J9261			10/11/2016	99/99/9999	INJECTION, NELARABINE, 50 MG	ARRANON (6X50ML,LATEX-FREE) 5 MG/1 ML	50	ML	VL	IV	ML	50	MG	0.1	10/11/2016	99/99/9999							
00078-0683-61	J9261			10/11/2016	99/99/9999	INJECTION, NELARABINE, 50 MG	ARRANON (LATEX-FREE) 5 MG/1 ML	50	ML	VL	IV	ML	50	MG	0.1	10/11/2016	99/99/9999							
00781-7515-87	J7626			08/20/2015	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.25 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	0.25	08/20/2015	99/99/9999							
63323-0300-30	J2543			09/24/2012	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE USE,PF) 3 GM-0.375 GM	10	EA	VL	IV	EA	1.125	GM	3	09/24/2012	99/99/9999							
00264-7751-00	J7121			01/01/2016	99/99/9999	5% DEXTROSE IN LACTATED RINGERS INFUSION, UP TO 1000 CC	DEXTROSE 5%/LACTATED RINGERS (EXCEL)	1000	ML	FC	IV	ML	1000	ML	0.001	01/01/2016	99/99/9999							
00264-7751-10	J7121			01/01/2016	99/99/9999	5% DEXTROSE IN LACTATED RINGERS INFUSION, UP TO 1000 CC	DEXTROSE 5%/LACTATED RINGERS (EXCEL)	500	ML	FC	IV	ML	1000	ML	0.001	01/01/2016	99/99/9999							
00338-0125-03	J7121			01/01/2016	99/99/9999	5% DEXTROSE IN LACTATED RINGERS INFUSION, UP TO 1000 CC	LACTATED RINGER'S AND 5% DEXTROSE (VIAFLEX)	500	ML	FC	IV	ML	1000	ML	0.001	01/01/2016	99/99/9999							
00338-0125-04	J7121			01/01/2016	99/99/9999	5% DEXTROSE IN LACTATED RINGERS INFUSION, UP TO 1000 CC	LACTATED RINGER'S AND 5% DEXTROSE (VIAFLEX, 14X1000ML)	1000	ML	FC	IV	ML	1000	ML	0.001	01/01/2016	99/99/9999							
00409-7929-03	J7121			01/01/2016	99/99/9999	5% DEXTROSE IN LACTATED RINGERS INFUSION, UP TO 1000 CC	DEXTROSE 5% IN RINGERS (LATEX-FREE)	500	ML	FC	IV	ML	1000	ML	0.001	01/01/2016	99/99/9999							
00409-7929-09	J7121			01/01/2016	99/99/9999	5% DEXTROSE IN LACTATED RINGERS INFUSION, UP TO 1000 CC	DEXTROSE 5% IN RINGERS (LIFECARE,LATEX-FREE)	1000	ML	FC	IV	ML	1000	ML	0.001	01/01/2016	99/99/9999							
69656-0101-02	J8670			01/01/2017	99/99/9999	ROLAPITANT, 1 MG	VARUBI (FILM COATED) 90 MG	2	EA	DP	PO	EA	1	MG	90	01/01/2017	99/99/9999							
68001-0286-38	J0640			11/23/2016	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (SDV,PF,LATEX-FREE) 350 MG	1	EA	VL	IJ	EA	50	MG	7	11/23/2016	99/99/9999							
68001-0285-40	J0640			11/23/2016	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (SDV,PF,LATEX-FREE) 50 MG	10	EA	VL	IJ	EA	50	MG	1	11/23/2016	99/99/9999							
68001-0285-36	J0640			11/23/2016	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (PF,LATEX-FREE) 100 MG	1	EA	VL	IJ	EA	50	MG	2	11/23/2016	99/99/9999							
68001-0285-37	J0640			11/23/2016	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (SDV,PF,LATEX-FREE) 200 MG	1	EA	VL	IJ	EA	50	MG	4	11/23/2016	99/99/9999							
63323-0871-15	J0878			08/30/2016	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (PF,LYPHILIZED) 500 MG	1	EA	VL	IV	EA	1	MG	500	08/30/2016	99/99/9999							
60505-6160-04	J1267			12/12/2016	99/99/9999	INJECTION, DORIPENEM, 10 MG	DORIPENEM 250 MG	10	EA	VL	IV	EA	10	MG	25	12/12/2016	99/99/9999							
60505-6161-00	J1267			12/12/2016	99/99/9999	INJECTION, DORIPENEM, 10 MG	DORIPENEM 500 MG	1	EA	VL	IV	EA	10	MG	50	12/12/2016	99/99/9999							
60505-6161-04	J1267			12/12/2016	99/99/9999	INJECTION, DORIPENEM, 10 MG	DORIPENEM 500 MG	10	EA	VL	IV	EA	10	MG	50	12/12/2016	99/99/9999							
60505-6160-00	J1267			12/12/2016	99/99/9999	INJECTION, DORIPENEM, 10 MG	DORIPENEM 250 MG	1	EA	VL	IV	EA	10	MG	25	12/12/2016	99/99/9999							
70121-1002-01	J1327			12/14/2016	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (SDV) 2 MG/1 ML	10	ML	VL	IV	ML	5	MG	0.4	12/14/2016	99/99/9999							
70121-1003-01	J1327			12/14/2016	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (SDV) 0.75 MG/1 ML	100	ML	VL	IV	ML	5	MG	0.15	12/14/2016	99/99/9999							
51991-0144-17	J2210			11/10/2016	99/99/9999	INJECTION, METHYLERGONOVINE MALEATE, UP TO 0.2 MG	METHYLERGONOVINE MALEATE (USP) 0.2 MG/1 ML	1	ML	AM	IJ	ML	0.2	MG	1	11/10/2016	99/99/9999							
00078-0818-81	J2353			12/08/2016	99/99/9999	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1 1/2"X19G) 20 MG	1	EA	BX	IM	EA	1	MG	20	12/08/2016	99/99/9999							
00078-0825-81	J2353			12/06/2016	99/99/9999	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1 1/2"X19G) 30 MG	1	EA	BX	IM	EA	1	MG	30	12/06/2016	99/99/9999							
00143-9529-01	J2680			12/12/2016	99/99/9999	INJECTION, FLUPHENAZINE DECAANOATE, UP TO 25 MG	FLUPHENAZINE DECAANOATE 25 MG/1 ML	5	ML	VL	IJ	ML	25	MG	1	12/12/2016	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
36000-0242-01	J3260			09/17/2016	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (MDV,USP,LATEX-FREE) 40 MG/1 ML	30 ML	VL	IJ	ML		80 MG		0.5	09/17/2016	99/99/9999						
36000-0244-25	J3260			09/17/2016	99/99/9999	INJECTION, TOBRAMYCIN SULFATE, UP TO 80 MG	TOBRAMYCIN SULFATE (MDV,USP,LATEX-FREE) 40 MG/1 ML	2 ML	VL	IJ	ML		80 MG		0.5	09/17/2016	99/99/9999						
51862-0083-14	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	14 EA	BO	PO	EA		5 MG		1	11/18/2016	99/99/9999						
51862-0087-51	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5 EA	BO	PO	EA		20 MG		9	11/18/2016	99/99/9999						
51862-0088-51	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE 250 MG	5 EA	BO	PO	EA		250 MG		1	11/18/2016	99/99/9999						
51862-0084-14	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14 EA	BO	PO	EA		20 MG		1	11/18/2016	99/99/9999						
51862-0086-51	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5 EA	BO	PO	EA		20 MG		7	11/18/2016	99/99/9999						
51862-0084-51	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5 EA	BO	PO	EA		20 MG		1	11/18/2016	99/99/9999						
51862-0085-14	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14 EA	BO	PO	EA		100 MG		1	11/18/2016	99/99/9999						
51862-0087-14	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14 EA	BO	PO	EA		100 MG		9	11/18/2016	99/99/9999						
51862-0083-51	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 5MG, ORAL	TEMOZOLOMIDE 5 MG	5 EA	BO	PO	EA		5 MG		1	11/18/2016	99/99/9999						
51862-0085-51	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5 EA	BO	PO	EA		100 MG		1	11/18/2016	99/99/9999						
51862-0086-14	None			11/18/2016	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14 EA	BO	PO	EA		20 MG		7	11/18/2016	99/99/9999						
16729-0288-11	J9060			12/07/2016	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (LATEX-FREE) 1 MG/1 ML	50 ML	VL	IV	ML		10 MG		0.1	12/07/2016	99/99/9999						
16729-0288-38	J9060			12/07/2016	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (LATEX-FREE) 1 MG/1 ML	100 ML	VL	IV	ML		10 MG		0.1	12/07/2016	99/99/9999						
00955-1022-08	J9171			11/17/2016	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (1X8ML,SINGLE USE) 20 MG/1 ML	8 ML	VL	IV	ML		1 MG		20	11/17/2016	99/99/9999						
57894-0200-01	J0130			01/01/2017	99/99/9999	INJECTION, ALCIXIMAB, 10 MG	REOPRO (VIAL,PF) 2 MG/1 ML	5 ML	VL	IV	ML		10 MG		0.2	01/01/2017	99/99/9999						
00009-0274-01	J1020			02/02/1987	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 20 MG	DEPO-MEDROL (M.D.V.) 20 MG/1 ML	5 ML	VL	IJ	ML		20 MG		1	02/02/1987	99/99/9999						
45963-0613-59	J9267			01/13/2015	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV,PF) 6 MG/1 ML	50 ML	VL	IV	ML		1 MG		6	01/13/2015	99/99/9999						
54569-1818-04	None			01/08/2015	10/17/2016	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	12 EA	BO	PO	EA		2.5 MG		1	01/08/2015	10/17/2016						
60687-0149-11	None			03/11/2016	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (INNER NDC,FILM-COATED) 500 MG	1 EA	BP	PO	EA		500 MG		1	03/11/2016	99/99/9999						
60687-0149-94	None			03/11/2016	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (2X10,FILM-COATED) 500 MG	20 EA	BX	PO	EA		500 MG		1	03/11/2016	99/99/9999						
49502-0101-02	J0171			12/15/2016	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE (0.15 MG/DELIVERY) 0.15 MG/0.3 ML	2 EA	SR	MR	EA		0.1 MG		1.5	12/15/2016	99/99/9999						
49502-0102-02	J0171			12/15/2016	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE AUTO-INJECTORS (0.3 MG/DELIVERY) 0.3 MG/0.3 ML	2 EA	SR	MR	EA		0.1 MG		3	12/15/2016	99/99/9999						
63323-0707-20	J0290			01/05/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM, 250 MG	10 EA	VL	IJ	EA		500 MG		0.5	01/05/2017	99/99/9999						
63323-0705-08	J0290			01/05/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM, 2 GM	10 EA	VL	IJ	EA		500 MG		4	01/05/2017	99/99/9999						
00143-9559-01	J0883			12/27/2016	99/99/9999	INJECTION, ARGATROBAN, 1 MG (FOR NON-ESRD USE)	ARGATROBAN (SDV,PF) 1 MG/1 ML	50 ML	VL	IV	ML		1 MG		1	12/27/2016	99/99/9999						
50242-0080-03	J2778			01/30/2017	99/99/9999	INJECTION, RANIBIZUMAB, 0.1 MG	INJECTION, RHO D IMMUNE GLOBULIN, INTRAVENOUS, HUMAN, SOLVENT	0.05 ML	SR	IO	ML		0.1 MG		100	01/30/2017	99/99/9999						
70504-3500-02	J2792			01/01/2017	99/99/9999	DETERGENT, 100 IU	WINRHO SDF (1X2.2ML,SDV) 2500 IU	2.2 ML	VL	IV	ML		100 IU		11.36363	01/01/2017	99/99/9999						
70504-3300-02	J2792			01/01/2017	99/99/9999	DETERGENT, 100 IU	WINRHO SDF (1X1.3ML,SDV) 1500 IU	1.3 ML	VL	IV	ML		100 IU		11.53846	01/01/2017	99/99/9999						
70504-3100-02	J2792			01/01/2017	99/99/9999	DETERGENT, 100 IU	WINRHO SDF (1X4.4ML,SDV) 5000 IU	4.4 ML	VL	IV	ML		100 IU		11.36363	01/01/2017	99/99/9999						
70504-3000-02	J2792			01/01/2017	99/99/9999	DETERGENT, 100 IU	WINRHO SDF (SDV) 15000 IU	13 ML	VL	IV	ML		100 IU		11.53846	01/01/2017	99/99/9999						
64679-0096-01	J9025			12/23/2016	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (SDV) 100 MG	1 EA	VL	IJ	EA		1 MG		100	12/23/2016	99/99/9999						
45963-0765-52	J9171			12/22/2016	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SINGLE-USE VIAL,PF) 20 MG/1 ML	4 ML	VL	IV	ML		1 MG		20	12/22/2016	99/99/9999						
25021-0242-02	J9185			12/19/2016	99/99/9999	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE (1X2ML,SDV,USP,PF) 25 MG/1 ML	2 ML	VL	IV	ML		50 MG		0.5	12/19/2016	99/99/9999						
60505-6132-07	J9263			01/05/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (1X20ML,SINGLE USE,PF) 5 MG/1 ML	20 ML	VL	IV	ML		0.5 MG		10	01/05/2017	99/99/9999						
60505-6132-06	J9263			01/05/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (1X10ML,SINGLE USE,PF) 5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		10	01/05/2017	99/99/9999						
00078-0674-61	J9351			01/05/2017	99/99/9999	INJECTION, TOPOTECAN, 0.1 MG	HYCANTIN (S.D.V.) 4 MG	1 EA	VL	IV	EA		0.1 MG		40	01/05/2017	99/99/9999						
60842-0023-01	J0171			01/19/2017	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	AUVI-Q 0.3 MG/0.3 ML	2 EA	BX	IJ	EA		0.1 MG		3	01/19/2017	99/99/9999						
60842-0022-01	J0171			01/19/2017	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	AUVI-Q 0.15 MG/0.15 ML	2 EA	BX	IJ	EA		0.1 MG		1.5	01/19/2017	99/99/9999						
00115-1695-49	J0171			02/10/2017	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE 0.15 MG/0.15 ML	2 EA	BX	IJ	EA		0.1 MG		1.5	02/10/2017	99/99/9999						
00115-1694-49	J0171			02/15/2017	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE (USP) 0.3 MG/0.3 ML	2 EA	BX	IJ	EA		0.1 MG		3	02/15/2017	99/99/9999						
70860-0100-10	J0456			02/01/2017	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN (SDV,LYOPHILIZED) 500 MG	10 EA	VL	IV	EA		500 MG		1	02/01/2017	99/99/9999						
45963-0762-57	J0641			02/14/2017	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (SDV,PF,LATEX-FREE) 50 MG	1 EA	VL	IV	EA		0.5 MG		100	02/14/2017	99/99/9999						
70121-1099-01	J0641			02/16/2017	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (SDV,PF,LYOPHILIZED) 50 MG	1 EA	VL	IV	EA		0.5 MG		100	02/16/2017	99/99/9999						
00591-4130-54	J0641			02/06/2017	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (SDV,PF,LATEX-FREE) 175 MG	1 EA	VL	IV	EA		0.5 MG		350	02/06/2017	99/99/9999						
23155-0294-41	J0780			01/09/2017	99/99/9999	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE 5 MG/1 ML	2 ML	VL	IJ	ML		10 MG		0.5	01/09/2017	99/99/9999						
00409-0106-01	J0878			01/04/2017	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (SDV,PF,LYOPHILIZED) 500 MG	1 EA	VL	IV	EA		1 MG		500	01/04/2017	99/99/9999						
00641-6145-25	J1100			01/20/2017	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1 MG	DEXAMETHASONE SODIUM PHOSPHATE 4 MG/1 ML	2 ML	VL	IJ	ML		1 MG		4	01/20/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00641-6146-25	J1100			01/20/2017	99/99/9999	INJECTION, DEXAMETHASONE SODIUM PHOSPHATE, 1 MG	DEXAMETHASONE SODIUM PHOSPHATE 4 MG/1 ML	5 ML	VL	IJ	ML		1 MG		4	01/20/2017	99/99/9999						
00004-6940-04	J1570			03/01/2017	99/99/9999	INJECTION, GANCICLOVIR SODIUM, 500 MG	CYTOVENE IV 500 MG	5 EA	VL	IV	EA		500 MG		1	03/01/2017	99/99/9999						
55150-0246-47	J1953			01/06/2017	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM-SODIUM CHLORIDE (LATEX-FREE) 500 MG/100 ML-0.82%	100 ML	BG	IV	ML		10 MG		0.5	01/06/2017	99/99/9999						
55150-0248-47	J1953			01/06/2017	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM-SODIUM CHLORIDE (LATEX-FREE) 1500 MG/100 ML-0.54%	100 ML	BG	IV	ML		10 MG		1.5	01/06/2017	99/99/9999						
55150-0247-47	J1953			01/06/2017	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM-SODIUM CHLORIDE (LATEX-FREE) 1000 MG/100 ML-0.75%	100 ML	BG	IV	ML		10 MG		1	01/06/2017	99/99/9999						
70860-0600-02	J2250			02/01/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (SDV) 1 MG/1 ML	2 ML	VL	IJ	ML		1 MG		1	02/01/2017	99/99/9999						
70860-0601-05	J2250			02/01/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (MDV) 5 MG/1 ML	5 ML	VL	IJ	ML		1 MG		5	02/01/2017	99/99/9999						
70860-0601-10	J2250			02/01/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (MDV) 5 MG/1 ML	10 ML	VL	IJ	ML		1 MG		5	02/01/2017	99/99/9999						
23155-0601-42	J2250			01/30/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (MDV) 5 MG/1 ML	10 ML	VL	IJ	ML		1 MG		5	01/30/2017	99/99/9999						
23155-0600-41	J2250			01/30/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (SDV) 1 MG/1 ML	2 ML	VL	IJ	ML		1 MG		1	01/30/2017	99/99/9999						
23155-0601-41	J2250			01/30/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM (MDV) 5 MG/1 ML	5 ML	VL	IJ	ML		1 MG		5	01/30/2017	99/99/9999						
70860-0776-02	J2405			02/01/2017	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON HCL (SDV,PF) 2 MG/1 ML	2 ML	VL	IJ	ML		1 MG		2	02/01/2017	99/99/9999						
70860-0777-20	J2405			02/01/2017	99/99/9999	INJECTION, ONDANSETRON HYDROCHLORIDE, PER 1 MG	ONDANSETRON (MDV) 2 MG/1 ML	20 ML	VL	IJ	ML		1 MG		2	02/01/2017	99/99/9999						
39822-0123-02	J2543			02/13/2017	99/99/9999	GRAMS) INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (PF,LATEX-FREE) 2 GM-0.25 GM	10 EA	VL	IV	EA		1.125 GM		2	02/13/2017	99/99/9999						
39822-0125-04	J2543			02/13/2017	99/99/9999	GRAMS) INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (PF,LATEX-FREE) 3 GM-0.375 GM	10 EA	VL	IV	EA		1.125 GM		3	02/13/2017	99/99/9999						
39822-0127-06	J2543			02/13/2017	99/99/9999	GRAMS) INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (PF,LATEX-FREE) 4 GM-0.5 GM	10 EA	VL	IV	EA		1.125 GM		4	02/13/2017	99/99/9999						
39822-0139-07	J2543			02/13/2017	99/99/9999	GRAMS) INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (PHARMACY BULK PACKAGE) 36 GM-4.5 GM	1 EA	VL	IV	EA		1.125 GM		36	02/13/2017	99/99/9999						
00409-2999-14	J2543			01/23/2017	99/99/9999	GRAMS) INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (PF,LATEX-FREE) 12 GM-1.5 GM	1 EA	BO	IV	EA		1.125 GM		12	01/23/2017	99/99/9999						
52565-0102-01	J2780			01/11/2017	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC (M.D.V.) 25 MG/1 ML	6 ML	VL	IJ	ML		25 MG		1	01/11/2017	99/99/9999						
52565-0101-10	J2780			01/11/2017	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC 25 MG/1 ML	2 ML	VL	IJ	ML		25 MG		1	01/11/2017	99/99/9999						
52565-0096-01	J2780			01/11/2017	99/99/9999	INJECTION, RANITIDINE HYDROCHLORIDE, 25 MG	ZANTAC 25 MG/1 ML	40 ML	VL	IJ	ML		25 MG		1	01/11/2017	99/99/9999						
70860-0105-20	J3370			02/01/2017	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF,LATEX-FREE) 1 GM	10 EA	VL	IV	EA		500 MG		2	02/01/2017	99/99/9999						
70860-0104-10	J3370			02/01/2017	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF) 500 MG	10 EA	VL	IV	EA		500 MG		1	02/01/2017	99/99/9999						
49702-0213-26	J3485			01/05/2017	99/99/9999	INJECTION, ZIDOVUDINE, 10 MG	RETROVIR (SINGLE USE PF) 10 MG/1 ML	20 ML	VL	IV	ML		10 MG		1	01/05/2017	99/99/9999						
00121-0777-08	J7510			02/10/2017	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (AF,DYE-FREE) 20 MG/5 ML	237 ML	BO	PO	ML		5 MG		0.8	02/10/2017	99/99/9999						
00121-0773-08	J7510			02/10/2017	99/99/9999	PREDNISOLONE ORAL, PER 5 MG	PREDNISOLONE SODIUM PHOSPHATE (AF,DYE-FREE,GRAPE) 10 MG/5 ML	237 ML	BO	PO	ML		5 MG		0.4	02/10/2017	99/99/9999						
49884-0373-01	J8540			01/25/2017	01/05/2018	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE 6 MG	100 EA	BO	PO	EA		0.25 MG		24	01/25/2017	01/05/2018						
45963-0687-49	J9245			01/19/2017	99/99/9999	INJECTION, MELPHALAN HYDROCHLORIDE, 50 MG	MELPHALAN HYDROCHLORIDE (INNER VIAL NDC,PF) 50 MG	1 EA	VL	IV	EA		50 MG		1	01/19/2017	99/99/9999						
45963-0686-02	J9245			01/19/2017	99/99/9999	INJECTION, MELPHALAN HYDROCHLORIDE, 50 MG	MELPHALAN HYDROCHLORIDE (W/ 10ML DILUENT,PF) 50 MG	1 EA	VL	IV	EA		50 MG		1	01/19/2017	99/99/9999						
43825-0102-01	J0131			01/03/2011	99/99/9999	INJECTION, ACETAMINOPHEN, 10 MG	OFIRMEV 10 MG/1 ML	100 ML	VL	IV	ML		10 MG		1	01/03/2011	99/99/9999						
51862-0458-47	J7515			07/18/2016	99/99/9999	CYCLOSPORINE, ORAL, 25 MG	CYCLOSPORINE (USP,MODIFIED) 25 MG	30 EA	ST	PO	EA		25 MG		1	07/18/2016	99/99/9999						
00023-5902-04	J3315			03/13/2017	99/99/9999	INJECTION, TRIPTORELIN PAMOATE, 3.75 MG	TRELSTAR (W/MIXJECT SYSTEM) 3.75 MG	1 EA	VL	IM	EA		3.75 MG		1	03/13/2017	99/99/9999						
00023-5904-12	J3315			03/13/2017	99/99/9999	INJECTION, TRIPTORELIN PAMOATE, 3.75 MG	TRELSTAR (W/MIXJECT SYSTEM) 11.25 MG	1 EA	VL	IM	EA		3.75 MG		3	03/13/2017	99/99/9999						
00078-0734-61	J0638			03/08/2017	99/99/9999	INJECTION, CANAKINUMAB, 1 MG	ILARIS (PF) 150 MG/1 ML	1 ML	VL	SC	ML		1 MG		150	03/08/2017	99/99/9999						
00409-1140-01	J0883			02/22/2017	99/99/9999	INJECTION, ARGATROBAN, 1 MG (FOR NON-ESRD USE)	ARGATROBAN (SDV,PF) 100 MG/1 ML	2.5 ML	VL	IV	ML		1 MG		100	02/22/2017	99/99/9999						
00781-7146-63	J7620			02/21/2017	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (60X3ML) 3 MG/3 ML-0.5 MG/3 ML	3 ML	VL	IH	ML		3 MG		0.33333	02/21/2017	99/99/9999						
00781-7146-87	J7620			03/15/2017	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE (30X3ML) 3 MG/3 ML-0.5 MG/3 ML	3 ML	VL	IH	ML		3 MG		0.33333	03/15/2017	99/99/9999						
00904-6623-61	J7507			03/20/2017	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 0.5 MG	100 EA	ST	PO	EA		1 MG		0.5	03/20/2017	99/99/9999						
00904-6624-61	J7507			03/20/2017	99/99/9999	TACROLIMUS, IMMEDIATE RELEASE, ORAL, 1 MG	TACROLIMUS (HARD GELATIN) 5 MG	100 EA	ST	PO	EA		1 MG		5	03/20/2017	99/99/9999						
16729-0224-05	J0894			03/03/2017	99/99/9999	INJECTION, DECITABINE, 1 MG	DECITABINE (LYOPHILIZED) 50 MG	1 EA	VL	IV	EA		1 MG		50	03/03/2017	99/99/9999						
17478-0931-01	J0636			02/28/2017	99/99/9999	INJECTION, CALCITRIOL, 0.1 MCG	CALCITRIOL (10 X 1ML) 1 MCG/1 ML	1 ML	AM	IV	ML		0.1 MCG		10	02/28/2017	99/99/9999						
45963-0621-51	J9185			03/02/2017	99/99/9999	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE (PF,LATEX-FREE) 25 MG/1 ML	2 ML	VL	IV	ML		50 MG		0.5	03/02/2017	99/99/9999						
63323-0106-26	J3475			03/14/2017	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	PREMIERPRO RX MAGNESIUM SULFATE (FREEFLEX BAG,LATEX-FREE) 40 MG/1 ML	50 ML	BG	IV	ML		500 MG		0.08	03/14/2017	99/99/9999						
63323-0108-26	J3475			03/14/2017	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	PREMIERPRO RX MAGNESIUM SULFATE-DEXTROSE (FREEFLEX BAG,LATEX-FREE) 5%-1 GM/100 ML	100 ML	BG	IV	ML		500 MG		0.02	03/14/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
69639-0101-01		J8655		04/01/2017	99/99/9999	Netupitant 300 mg and palonosetron 0.5 mg, oral	AKYNZEO (HARD GELATIN) 300 MG-0.5 MG	1	EA	ST	PO	EA	300.5	MG	1	04/01/2017	99/99/9999						
70121-1000-05		J2920		02/28/2017	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 40 MG	METHYLPREDNISOLONE SODIUM SUCCINATE (SDV,LYOPHILIZED) 40 MG	25	EA	VL	IJ	EA	40	MG	1	02/28/2017	99/99/9999						
70121-1001-05		J2930		02/28/2017	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	METHYLPREDNISOLONE SODIUM SUCCINATE (SDV,LYOPHILIZED) 125 MG	25	EA	VL	IJ	EA	125	MG	1	02/28/2017	99/99/9999						
00409-7241-10		J0171		09/01/2016	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE (INNER NDC) 1 MG/1 ML	1	ML	AM	IJ	ML	0.1	MG	10	09/01/2016	99/99/9999						
16729-0048-53		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	5	EA	BO	PO	EA	5	MG	1	02/28/2017	99/99/9999						
16729-0048-54		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	14	EA	BO	PO	EA	5	MG	1	02/28/2017	99/99/9999						
16729-0049-53		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5	EA	BO	PO	EA	20	MG	1	02/28/2017	99/99/9999						
16729-0049-54		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14	EA	BO	PO	EA	20	MG	1	02/28/2017	99/99/9999						
16729-0050-53		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5	EA	BO	PO	EA	100	MG	1	02/28/2017	99/99/9999						
16729-0050-54		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14	EA	BO	PO	EA	100	MG	1	02/28/2017	99/99/9999						
16729-0051-53		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE 250 MG	5	EA	BO	PO	EA	250	MG	1	02/28/2017	99/99/9999						
16729-0129-53		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	5	EA	BO	PO	EA	20	MG	7	02/28/2017	99/99/9999						
16729-0129-54		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	14	EA	BO	PO	EA	20	MG	7	02/28/2017	99/99/9999						
16729-0130-53		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	5	EA	BO	PO	EA	20	MG	9	02/28/2017	99/99/9999						
16729-0130-54		None		02/28/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	14	EA	BO	PO	EA	20	MG	9	02/28/2017	99/99/9999						
64980-0276-06		None		03/15/2017	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM COATED) 150 MG	60	EA	BO	PO	EA	150	MG	1	03/15/2017	99/99/9999						
64980-0277-12		None		03/15/2017	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (USP,FILM COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	03/15/2017	99/99/9999						
65162-0843-06		None		03/10/2017	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (USP,FILM COATED) 150 MG	60	EA	BO	PO	EA	150	MG	1	03/10/2017	99/99/9999						
65162-0844-16		None		03/10/2017	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (USP,FILM COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	03/10/2017	99/99/9999						
68382-0775-01		None		02/27/2017	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE (USP) 2.5 MG	100	EA	BO	PO	EA	2.5	MG	1	02/27/2017	99/99/9999						
00078-0673-01		None		03/21/2017	99/99/9999	TOPOTECAN, ORAL, 0.25 MG	HYCAMTIN 1 MG	10	EA	BO	PO	EA	0.25	MG	4	03/21/2017	99/99/9999						
00143-9875-25		J0282		03/30/2017	99/99/9999	INJECTION, AMIODARONE HYDROCHLORIDE, 30 MG	AMIODARONE HCL (10X3ML) 50 MG/1 ML	3	ML	VL	IV	ML	30	MG	1.66666	03/30/2017	99/99/9999						
00517-0920-01		J0594		04/01/2017	99/99/9999	INJECTION, BUSULFAN, 1 MG	BUSULFAN 6 MG/1 ML	10	ML	VL	IV	ML	1	MG	6	04/01/2017	99/99/9999						
00517-0920-08		J0594		04/01/2017	99/99/9999	INJECTION, BUSULFAN, 1 MG	BUSULFAN (8X10ML SINGLE-USE) 6 MG/1 ML	10	ML	VL	IV	ML	1	MG	6	04/01/2017	99/99/9999						
16714-0221-10		Q0166		03/17/2017	99/99/9999	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (INNER NDC,FILM-COATED) 1 MG	1	EA	ST	PO	EA	1	MG	1	03/17/2017	99/99/9999						
16714-0221-12		Q0166		03/17/2017	99/99/9999	GRANISETRON HYDROCHLORIDE, 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 24 HOUR DOSAGE REGIMEN	GRANISETRON HYDROCHLORIDE (FILM-COATED) 1 MG	10	EA	ST	PO	EA	1	MG	1	03/17/2017	99/99/9999						
25021-0221-60		J9245		04/21/2017	99/99/9999	INJECTION, MELPHALAN HYDROCHLORIDE, 50 MG	MELPHALAN HYDROCHLORIDE (W/10ML DILUENT,PF) 50 MG	1	EA	VL	IV	EA	50	MG	1	04/21/2017	99/99/9999						
25021-0807-05		J2920		04/17/2017	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 40 MG	METHYLPREDNISOLONE SODIUM SUCCINATE (LYOPHILIZED) 40 MG	10	EA	VL	IJ	EA	40	MG	1	04/17/2017	99/99/9999						
25021-0808-10		J2930		04/17/2017	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	METHYLPREDNISOLONE SODIUM SUCCINATE (LYOPHILIZED) 125 MG	10	EA	VL	IJ	EA	125	MG	1	04/17/2017	99/99/9999						
25021-0810-30		J2930		04/17/2017	99/99/9999	INJECTION, METHYLPREDNISOLONE SODIUM SUCCINATE, UP TO 125 MG	METHYLPREDNISOLONE SODIUM SUCCINATE (LATEX-FREE,LYOPHILIZED) 1 GM	1	EA	VL	IJ	EA	125	MG	8	04/17/2017	99/99/9999						
42023-0173-25		J1570		04/05/2017	99/99/9999	INJECTION, GANCICLOVIR SODIUM, 500 MG	GANCICLOVIR (SDV,PF,LYOPHILIZED) 500 MG	25	EA	VL	IV	EA	500	MG	1	04/05/2017	99/99/9999						
42023-0191-10		J2185		04/05/2017	12/21/2017	INJECTION, MEROPENEM, 100 MG	MEROPENEM (SDV,USP) 500 MG	10	EA	VL	IV	EA	100	MG	5	04/05/2017	12/21/2017						
42023-0192-10		J2185		04/05/2017	12/21/2017	INJECTION, MEROPENEM, 100 MG	MEROPENEM (SDV,USP) 1 GM	10	EA	VL	IV	EA	100	MG	10	04/05/2017	12/21/2017						
50419-0537-01		J2280		04/01/2017	99/99/9999	INJECTION, MOXIFLOXACIN, 100 MG	AVELOX I.V. (SINGLE-DOSE FLEXIBAG,PF) 400 MG/250 ML	250	ML	BG	IV	ML	100	MG	0.016	04/01/2017	99/99/9999						
55150-0207-20		J2185		03/27/2017	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM (USP) 500 MG	10	EA	VL	IV	EA	100	MG	5	03/27/2017	99/99/9999						
55150-0208-30		J2185		03/27/2017	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM (USP) 1 GM	10	EA	VL	IV	EA	100	MG	10	03/27/2017	99/99/9999						
60505-6146-00		J0692		04/03/2017	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP,SDV) 1 GM	1	EA	VL	IJ	EA	500	MG	2	04/03/2017	99/99/9999						
60505-6146-04		J0692		04/03/2017	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP,SDV) 1 GM	10	EA	VL	IJ	EA	500	MG	2	04/03/2017	99/99/9999						
60505-6147-00		J0692		04/03/2017	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP,SDV) 2 GM	1	EA	VL	IJ	EA	500	MG	4	04/03/2017	99/99/9999						
60505-6147-04		J0692		04/03/2017	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP,SDV) 2 GM	10	EA	VL	IJ	EA	500	MG	4	04/03/2017	99/99/9999						
63323-0771-39		J9025		04/13/2017	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (SDV) 100 MG	1	EA	VL	IJ	EA	1	MG	100	04/13/2017	99/99/9999						
63323-0966-00		J3489		03/31/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SDV) 5 MG/100 ML	100	ML	VL	IV	ML	1	MG	0.05	03/31/2017	99/99/9999						
64208-8235-05		J1557		04/01/2017	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), 500 MG	GAMMAPLEX 10% (PF,LATEX-FREE) 100 MG/1 ML	50	ML	VL	IV	ML	500	MG	0.2	04/01/2017	99/99/9999						
64208-8235-06		J1557		04/01/2017	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), 500 MG	GAMMAPLEX 10% (PF,LATEX-FREE) 100 MG/1 ML	100	ML	VL	IV	ML	500	MG	0.2	04/01/2017	99/99/9999						
64208-8235-07		J1557		04/01/2017	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), 500 MG	GAMMAPLEX 10% (PF,LATEX-FREE) 100 MG/1 ML	200	ML	VL	IV	ML	500	MG	0.2	04/01/2017	99/99/9999						
50111-0788-10		Q0144		04/05/2017	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (FILM-COATED) 500 MG	30	EA	BO	PO	EA	1	GM	0.5	04/05/2017	99/99/9999						
50268-0761-11		None		03/24/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (INNER PACK) 20 MG	1	EA	ST	PO	EA	20	MG	1	03/24/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
50268-0761-12		None		03/24/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (4 X 5) 20 MG	20	EA	ST	PO	EA	20	MG	1	03/24/2017	99/99/9999							
50268-0762-11		None		03/24/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE (INNERPACK) 100 MG	1	EA	ST	PO	EA	100	MG	1	03/24/2017	99/99/9999							
50268-0762-12		None		03/24/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	20	EA	ST	PO	EA	100	MG	1	03/24/2017	99/99/9999							
50268-0763-11		None		03/24/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (INNERPACK) 140 MG	1	EA	ST	PO	EA	20	MG	7	03/24/2017	99/99/9999							
50268-0763-12		None		03/24/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	20	EA	ST	PO	EA	20	MG	7	03/24/2017	99/99/9999							
00143-9501-25		J1630		04/17/2017	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL LACTATE 5 MG/1 ML	1	ML	VL	IM	ML	5	MG	1	04/17/2017	99/99/9999							
00143-9502-01		J1630		04/17/2017	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL LACTATE 5 MG/1 ML	10	ML	VL	IM	ML	5	MG	1	04/17/2017	99/99/9999							
00409-0528-15		J1956		05/15/2017	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (24X50ML, SINGLE-USE,PF) 5%-250 MG/50 ML	50	ML	BG	IV	ML	250	MG	0.02	05/15/2017	99/99/9999							
00409-0528-25		J1956		05/15/2017	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (24X100ML, SINGLE-USE,PF) 5%-500 MG/100 ML	100	ML	BG	IV	ML	250	MG	0.02	05/15/2017	99/99/9999							
00409-0528-35		J1956		05/15/2017	99/99/9999	INJECTION, LEVOFLOXACIN, 250 MG	LEVOFLOXACIN IN 5% DEXTROSE (24X150ML, SINGLE-USE,PF) 5%-750 MG/150 ML	150	ML	BG	IV	ML	250	MG	0.02	05/15/2017	99/99/9999							
39822-2180-01		J9171		05/05/2017	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SDV) 20 MG/1 ML	4	ML	VL	IV	ML	1	MG	20	05/05/2017	99/99/9999							
39822-2200-01		J9171		05/05/2017	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SDV) 20 MG/1 ML	10	ML	VL	IV	ML	1	MG	20	05/05/2017	99/99/9999							
44087-0016-01		J2941		04/21/2017	99/99/9999	INJECTION, SOMATROPIN, 1 MG	SAIZEN SAIZENPREP CARTRIDGE (W/DILUENT) 8.8 MG	1	EA	CT	IJ	EA	1	MG	8.8	04/21/2017	99/99/9999							
50242-0080-02		J2778		05/15/2017	04/30/2018	INJECTION, RANIBIZUMAB, 0.1 MG	LUCENTIS (INTRAVITREAL INJECTION) 0.5 MG/0.05 ML	0.05	ML	VL	IO	ML	0.1	MG	100	05/15/2017	04/30/2018							
50242-0082-02		J2778		05/15/2017	99/99/9999	INJECTION, RANIBIZUMAB, 0.1 MG	LUCENTIS (INTRAVITREAL INJECTION) 0.3 MG/0.05 ML	0.05	ML	VL	IO	ML	0.1	MG	60	05/15/2017	99/99/9999							
51754-2500-03		J1570		06/01/2017	99/99/9999	INJECTION, GANCICLOVIR SODIUM, 500 MG	GANCICLOVIR-SODIUM CHLORIDE (PF) 500 MG/250 ML-0.8%	250	ML	BG	IV	ML	500	MG	0.004	06/01/2017	99/99/9999							
53964-0001-01		J9340		04/21/2017	99/99/9999	INJECTION, THIOTEPA, 15 MG	TEPADINA 15 MG	1	EA	VL	IJ	EA	15	MG	1	04/21/2017	99/99/9999							
53964-0002-02		J9340		04/21/2017	99/99/9999	INJECTION, THIOTEPA, 15 MG	TEPADINA 100 MG	1	EA	VL	IJ	EA	15	MG	6.66666	04/21/2017	99/99/9999							
63323-0572-70		J9027		04/25/2017	99/99/9999	INJECTION, CLOFARABINE, 1 MG	CLOFARABINE (PF,LATEX-FREE) 1 MG/1 ML	20	ML	VL	IV	ML	1	MG	1	04/25/2017	99/99/9999							
62991-2700-01		J3121		10/17/2016	99/99/9999	INJECTION, TESTOSTERONE ENANTHATE, 1 MG	TESTOSTERONE ENANTHATE (USP, 1X1000GM)	1000	GM	BO	NA	GM	1	MG	1000	10/17/2016	99/99/9999							
00781-7517-87		J7626		07/27/2015	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (SINGLE DOSE) 1 MG/2 ML RASUVO (1X4 AUTO INJECTORS,PF) 10 MG/0.2 ML	2	ML	AM	IH	ML	0.5	MG	1	07/27/2015	99/99/9999							
59137-0510-04		J9250		09/22/2014	99/99/9999	METHOTREXATE SODIUM, 5 MG	CAPECITABINE (USP, FILM COATED) 500 MG	0.2	ML	CT	SC	ML	5	MG	10	09/22/2014	99/99/9999							
42291-0167-12		None		04/14/2017	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (USP, FILM COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	04/14/2017	99/99/9999							
00781-7517-87	KO	J7626	KO	07/27/2015	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (SINGLE DOSE) 1 MG/2 ML	2	ML	AM	IH	ML	0.5	MG	1	07/27/2015	99/99/9999							
59137-0510-04		J9250		09/22/2014	99/99/9999	METHOTREXATE SODIUM, 5 MG	CAPECITABINE (USP, FILM COATED) 500 MG	0.2	ML	CT	SC	ML	5	MG	10	09/22/2014	99/99/9999							
42291-0167-12		None		04/14/2017	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (USP, FILM COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	04/14/2017	99/99/9999							
00781-7517-87	KO	J7626	KO	07/27/2015	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (SINGLE DOSE) 1 MG/2 ML	2	ML	AM	IH	ML	0.5	MG	1	07/27/2015	99/99/9999							
00003-2814-11		J0129		04/06/2017	99/99/9999	INJECTION, ABATACEPT, 10 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ORENCIA (PF LYOPHILIZED) 50 MG/0.4 ML	0.4	ML	SR	SC	ML	10	MG	12.5	04/06/2017	99/99/9999							
00003-2818-11		J0129		04/06/2017	99/99/9999	INJECTION, ABATACEPT, 10 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ORENCIA (SD PREFILLED SYRINGE,PF) 87.5 MG/0.7 ML	0.7	ML	SR	SC	ML	10	MG	12.5	04/06/2017	99/99/9999							
00023-5906-23		J3315		06/08/2017	99/99/9999	INJECTION, TRIPTORELIN PAMOATE, 3.75 MG	TRELSTAR (W/MIXJECT SYSTEM) 22.5 MG SANDOSTATIN LAR DEPOT (1 1/2"X19G) 10 MG	1	EA	VL	IM	EA	3.75	MG	6	06/08/2017	99/99/9999							
00078-0811-81		J2353		05/10/2017	99/99/9999	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (1 1/2"X19G) 10 MG	1	EA	BX	IM	EA	1	MG	10	05/10/2017	99/99/9999							
00085-4320-01		J0702		05/16/2017	99/99/9999	INJECTION, BETAMETHASONE ACETATE 3 MG AND BETAMETHASONE SODIUM PHOSPHATE 3 MG	CELESTONE SOLUSPAN (MDV) 3 MG/1 ML 3 MG/1 ML	5	ML	VL	IJ	ML	6	MG	1	05/16/2017	99/99/9999							
00121-0489-00		Q0163		06/07/2017	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	5	ML	CP	PO	ML	50	MG	0.05	06/07/2017	99/99/9999							
00121-0489-00		Q0163		06/07/2017	99/99/9999	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DIPHENHYDRAMINE HCL 12.5 MG/5 ML	10	ML	CP	PO	ML	50	MG	0.05	06/07/2017	99/99/9999							
00955-1746-01		J9027		05/30/2017	99/99/9999	INJECTION, CLOFARABINE, 1 MG	CLOFARABINE (PF) 1 MG/1 ML	20	ML	VL	IV	ML	1	MG	1	05/30/2017	99/99/9999							
25021-0241-10		J0594		06/19/2017	99/99/9999	INJECTION, BUSULFAN, 1 MG	BUSULFAN 6 MG/1 ML	10	ML	VL	IV	ML	1	MG	6	06/19/2017	99/99/9999							
25021-0676-20		J2515		05/10/2017	99/99/9999	INJECTION, PENTOBARBITAL SODIUM, PER 50 MG	PENTOBARBITAL SODIUM (MDV,PF,LATEX-FREE) 50 MG/1 ML	20	ML	VL	IJ	ML	50	MG	1	05/10/2017	99/99/9999							
42023-0188-10		J2710		05/22/2017	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 0.5 MG/1 ML	10	ML	VL	IV	ML	0.5	MG	1	05/22/2017	99/99/9999							
42023-0189-10		J2710		05/22/2017	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 1 MG/1 ML	10	ML	VL	IV	ML	0.5	MG	2	05/22/2017	99/99/9999							
50242-0132-01		J9355		05/30/2017	99/99/9999	INJECTION, TRASTUZUMAB, 10 MG	HERCEPTIN (SDV,PF,LYOPHILIZED) 150 MG	1	EA	VL	IV	EA	10	MG	15	05/30/2017	99/99/9999							
55111-0652-07		J0583		05/31/2017	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (SINGLE-USE LYOPHILIZED) 250 MG	1	EA	VL	IV	EA	1	MG	250	05/31/2017	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55111-0652-37		J0583		05/31/2017	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (SINGLE-USE,LYOPHILIZED) 250 MG	10	EA	VL	IV	EA	1	MG	250	05/31/2017	99/99/9999						
64679-0012-01		J2543		06/12/2017	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 4 GM-0.5 GM	10	EA	VL	IV	EA	1.125	GM	4	06/12/2017	99/99/9999						
64679-0034-01		J2543		06/12/2017	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 2 GM-0.25 GM	10	EA	VL	IV	EA	1.125	GM	2	06/12/2017	99/99/9999						
64679-0056-01		J2543		06/12/2017	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 3 GM-0.375 GM	10	EA	VL	IV	EA	1.125	GM	3	06/12/2017	99/99/9999						
64679-0679-01		J2543		06/12/2017	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 36 GM-4.5 GM	1	EA	VL	IV	EA	1.125	GM	36	06/12/2017	99/99/9999						
65757-0404-03		J1942		06/05/2017	99/99/9999	INJECTION, ARIPIRAZOLE LAUROXIL, 1 MG	ARISTADA 1064 MG/3.9 ML	3.9	ML	SR	IM	ML	1	MG	272.8205	06/05/2017	99/99/9999						
67457-0619-10		J3489		05/19/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID 5 MG/100 ML	100	ML	VL	IV	ML	1	MG	0.05	05/19/2017	99/99/9999						
70655-0071-25		J2800		04/01/2017	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (LATEX-FREE) 100 MG/1 ML	10	ML	VL	IJ	ML	10	ML	0.1	04/01/2017	99/99/9999						
71297-0127-27		J8540		03/17/2017	03/21/2018	DEXAMETHASONE, ORAL, 0.25 MG	LOCORT (7-DAY) 1.5 MG	27	EA	ST	PO	EA	0.25	MG	6	03/17/2017	03/21/2018						
71297-0211-41		J8540		03/17/2017	03/21/2018	DEXAMETHASONE, ORAL, 0.25 MG	LOCORT (11-DAY) 1.5 MG	41	EA	ST	PO	EA	0.25	MG	6	03/17/2017	03/21/2018						
76204-0700-01		J7614		05/19/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.20666	05/19/2017	99/99/9999						
76204-0800-01		J7614		05/19/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.63 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.42	05/19/2017	99/99/9999						
76204-0900-01		J7614		05/19/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 1.25 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.83333	05/19/2017	99/99/9999						
67919-0030-01		J0695		12/22/2014	99/99/9999	INJECTION, CEFTIOZANE 50 MG AND TAZOBACTAM 25 MG	ZERBAXA (PF) 1 GM-0.5 GM	10	EA	VL	IV	EA	75	MG	20	12/22/2014	99/99/9999						
76204-0700-01	KO	J7614	KO	05/19/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.20666	05/19/2017	99/99/9999						
76204-0800-01	KO	J7614	KO	05/19/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.63 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.42	05/19/2017	99/99/9999						
76204-0900-01	KO	J7614	KO	05/19/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 1.25 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.83333	05/19/2017	99/99/9999						
60219-1076-01		J7500		04/13/2017	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE (USP) 50 MG	100	EA	BO	PO	EA	50	MG	1	04/13/2017	99/99/9999						
00006-3061-00		J1453		06/19/2017	99/99/9999	INJECTION, FOSAPREPITANT, 1 MG	EMEND (LYOPHILIZED) 150 MG	1	EA	VL	IV	EA	1	MG	150	06/19/2017	99/99/9999						
00078-0790-61		J2353		07/11/2017	99/99/9999	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (INNER PACK) 10 MG	1	EA	VL	IM	EA	1	MG	10	07/11/2017	99/99/9999						
00078-0797-61		J2353		07/11/2017	99/99/9999	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (INNER PACK) 20 MG	1	EA	VL	IM	EA	1	MG	20	07/11/2017	99/99/9999						
00078-0804-61		J2353		07/11/2017	99/99/9999	INJECTION, OCTREOTIDE, DEPOT FORM FOR INTRAMUSCULAR INJECTION, 1 MG	SANDOSTATIN LAR DEPOT (INNER PACK) 30 MG	1	EA	VL	IM	EA	1	MG	30	07/11/2017	99/99/9999						
13925-0523-01		J9025		07/07/2017	02/13/2018	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (PF,LYOPHILIZED) 100 MG	1	EA	VL	IJ	EA	1	MG	100	07/07/2017	02/13/2018						
47781-0200-50		None		06/27/2017	99/99/9999	MELPHALAN, 2 MG, ORAL	MELPHALAN (FILM COATED) 2 MG	50	EA	BO	PO	EA	2	MG	1	06/27/2017	99/99/9999						
55292-0702-54		J1640		07/01/2017	99/99/9999	INJECTION, HEMIN, 1 MG	PANHEMATIN (PF,LYOPHILIZED) 350 MG	1	EA	VL	IV	EA	1	MG	350	07/01/2017	99/99/9999						
55292-0702-55		J1640		07/01/2017	99/99/9999	INJECTION, HEMIN, 1 MG	PANHEMATIN (PF,LYOPHILIZED) 350 MG	1	EA	VL	IV	EA	1	MG	350	07/01/2017	99/99/9999						
60505-6148-00		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (CRYSTALLINE) 1 GM	1	EA	VL	IJ	EA	250	MG	4	06/23/2017	99/99/9999						
60505-6148-04		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (10X20ML,CRYSTALLINE) 1 GM	10	EA	VL	IJ	EA	250	MG	4	06/23/2017	99/99/9999						
60505-6149-00		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (CRYSTALLINE) 2 GM	1	EA	VL	IJ	EA	250	MG	8	06/23/2017	99/99/9999						
60505-6149-04		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (10X20ML,CRYSTALLINE) 2 GM	10	EA	VL	IJ	EA	250	MG	8	06/23/2017	99/99/9999						
60505-6151-01		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (SDV,CRYSTALLINE) 250 MG	10	EA	VL	IJ	EA	250	MG	1	06/23/2017	99/99/9999						
60505-6151-04		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (SDV,CRYSTALLINE) 250 MG	1	EA	VL	IJ	EA	250	MG	1	06/23/2017	99/99/9999						
60505-6152-01		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (10X10ML,CRYSTALLINE) 500 MG	10	EA	VL	IJ	EA	250	MG	2	06/23/2017	99/99/9999						
60505-6152-04		J0696		06/23/2017	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE (CRYSTALLINE) 500 MG	1	EA	VL	IJ	EA	250	MG	2	06/23/2017	99/99/9999						
63323-0704-08		J0290		06/23/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM (VIAL) 1 GM	10	EA	VL	IJ	EA	500	MG	2	06/23/2017	99/99/9999						
67457-0399-25		J3420		07/06/2017	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN 1000 MCG/1 ML	10	ML	VL	IJ	ML	1000	MCG	1	07/06/2017	99/99/9999						
67457-0400-05		J3420		07/06/2017	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN 1000 MCG/1 ML	30	ML	VL	IJ	ML	1000	MCG	1	07/06/2017	99/99/9999						
68001-0323-31		J2185		07/14/2017	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM (SDV,USP) 500 MG	10	EA	VL	IV	EA	100	MG	5	07/14/2017	99/99/9999						
68001-0324-57		J2185		07/14/2017	99/99/9999	INJECTION, MEROPENEM, 100 MG	MEROPENEM (SDV,USP) 1 GM	10	EA	VL	IV	EA	100	MG	10	07/14/2017	99/99/9999						
70257-0562-55		J0476		07/10/2017	99/99/9999	INJECTION, BACLOFEN, 50 MCG FOR INTRATHECAL TRIAL	LIORESAL INTRATHECAL (SCREENING #8563,PF) 0.05 MG/1 ML	1	ML	AM	IN	ML	50	MCG	1	07/10/2017	99/99/9999						
70860-0200-05		J9267		06/29/2017	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV,PF,LATEX-FREE) 6 MG/1 ML	5	ML	VL	IV	ML	1	MG	6	06/29/2017	99/99/9999						
70860-0200-17		J9267		06/29/2017	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV,PF,LATEX-FREE) 6 MG/1 ML	16.7	ML	VL	IV	ML	1	MG	6	06/29/2017	99/99/9999						
70860-0200-50		J9267		06/29/2017	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV,PF,LATEX-FREE) 6 MG/1 ML	50	ML	VL	IV	ML	1	MG	6	06/29/2017	99/99/9999						
70860-0201-10		J9263		06/29/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (MDV,PF,LATEX-FREE) 5 MG/1 ML	10	ML	VL	IV	ML	0.5	MG	10	06/29/2017	99/99/9999						
70860-0201-20		J9263		06/29/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (MDV,PF,LATEX-FREE) 5 MG/1 ML	20	ML	VL	IV	ML	0.5	MG	10	06/29/2017	99/99/9999						
70860-0700-01		J1885		07/01/2017	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (PF,LATEX-FREE) 15 MG/1 ML	1	ML	VL	IJ	ML	15	MG	1	07/01/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
70860-0701-01		J1885		07/01/2017	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (PF,LATEX-FREE) 30 MG/1 ML	1 ML	VL	IJ	ML		15 MG		2	07/01/2017	99/99/9999						
70860-0701-02		J1885		07/01/2017	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (PF,LATEX-FREE) 30 MG/1 ML	2 ML	VL	IM	ML		15 MG		2	07/01/2017	99/99/9999						
70860-0801-01		J3105		06/12/2017	99/99/9999	INJECTION, TERBUTALINE SULFATE, UP TO 1 MG	TERBUTALINE SULFATE (PF,LATEX-FREE) 1 MG/1 ML	1 ML	VL	SC	ML		1 MG		1	06/12/2017	99/99/9999						
63323-0130-11		J3490		10/29/2003	99/99/9999	UNCLASSIFIED DRUGS	DOXY 100 (VIAL,PF) 100 MG	10 EA	VL	IV	EA		1 MG		1	10/29/2003	99/99/9999						
00006-4305-02		Q5102		07/25/2017	03/31/2018	INJECTION, INFLIXIMAB, BIOSIMILAR, 10 MG	RENFLIXIS (PF,LYOPHILIZED) 100 MG	1 EA	VL	IV	EA		10 MG		10	07/25/2017	03/31/2018						
00037-9001-05		J1980		08/07/2017	99/99/9999	INJECTION, HYOSCYAMINE SULFATE, UP TO 0.25 MG	LEVSIN (5X1ML) 0.5 MG/1 ML	1 ML	AM	IJ	ML		0.25 MG		2	08/07/2017	99/99/9999						
00078-0672-01		None		07/31/2017	99/99/9999	TOPOTECAN, ORAL, 0.25 MG	HYCAMTIN 0.25 MG	10 EA	BO	PO	EA		0.25 MG		1	07/31/2017	99/99/9999						
00143-9217-01		J9211		07/18/2017	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (PF) 1 MG/1 ML	5 ML	VL	IV	ML		5 MG		0.2	07/18/2017	99/99/9999						
00143-9218-01		J9211		07/18/2017	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (PF) 1 MG/1 ML	10 ML	VL	IV	ML		5 MG		0.2	07/18/2017	99/99/9999						
00143-9219-01		J9211		07/18/2017	99/99/9999	INJECTION, IDARUBICIN HYDROCHLORIDE, 5 MG	IDARUBICIN HYDROCHLORIDE (PF) 1 MG/1 ML	20 ML	VL	IV	ML		5 MG		0.2	07/18/2017	99/99/9999						
00143-9261-10		J0690		07/27/2017	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN NOVAPLUS (PF,LATEX-FREE) 10 GM	10 EA	VL	IV	EA		500 MG		20	07/27/2017	99/99/9999						
00143-9262-25		J0690		07/27/2017	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN NOVAPLUS (PF,LATEX-FREE) 1 GM	25 EA	VL	IJ	EA		500 MG		2	07/27/2017	99/99/9999						
00781-3411-95		J0330		07/17/2017	99/99/9999	INJECTION, SUCCINYLCHOLINE CHLORIDE, UP TO 20 MG	ANECTAN (MDV) 20 MG/1 ML	10 ML	VL	IV	ML		20 MG		1	07/17/2017	99/99/9999						
17478-0041-01		J2310		08/07/2017	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (SDV,PF) 0.4 MG/1 ML	1 ML	VL	IJ	ML		1 MG		0.4	08/07/2017	99/99/9999						
17478-0042-10		J2310		08/14/2017	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (MDV) 0.4 MG/1 ML	10 ML	VL	IJ	ML		1 MG		0.4	08/14/2017	99/99/9999						
51991-0922-98		J9263		07/19/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF,LATEX-FREE) 5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		10	07/19/2017	99/99/9999						
51991-0923-98		J9263		07/19/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF,LATEX-FREE) 5 MG/1 ML	20 ML	VL	IV	ML		0.5 MG		10	07/19/2017	99/99/9999						
51991-0936-98		J9267		07/19/2017	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV) 6 MG/1 ML	5 ML	VL	IV	ML		1 MG		6	07/19/2017	99/99/9999						
51991-0937-98		J9267		07/19/2017	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV) 6 MG/1 ML	16.7 ML	VL	IV	ML		1 MG		6	07/19/2017	99/99/9999						
51991-0938-98		J9267		07/19/2017	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV) 6 MG/1 ML	50 ML	VL	IV	ML		1 MG		6	07/19/2017	99/99/9999						
57664-0683-31		J2020		08/10/2017	99/99/9999	INJECTION, LINEZOLID, 200 MG	LINEZOLID (INNER PACK,LATEX-FREE) 2 MG/1 ML	300 ML	BG	IV	ML		200 MG		0.01	08/10/2017	99/99/9999						
57664-0683-57		J2020		08/10/2017	99/99/9999	INJECTION, LINEZOLID, 200 MG	LINEZOLID (10X300ML BAGS) 2 MG/1 ML	300 ML	BG	IV	ML		200 MG		0.01	08/10/2017	99/99/9999						
59676-0966-01		Q2050		07/24/2017	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, LIPOSOMAL, NOT OTHERWISE SPECIFIED, 10 MG	DOXORUBICIN HCL LIPOSOME 2 MG/1 ML	10 ML	VL	IV	ML		10 MG		0.2	07/24/2017	99/99/9999						
60505-6101-04		J0583		07/17/2017	99/99/9999	INJECTION, BIVALIRUDIN, 1 MG	BIVALIRUDIN (SDV,LYOPHILIZED) 250 MG	10 EA	VL	IV	EA		1 MG		250	07/17/2017	99/99/9999						
60505-6142-00		J0690		08/07/2017	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN (INNER PACK,PF) 1 GM	1 EA	VL	IJ	EA		500 MG		2	08/07/2017	99/99/9999						
60505-6142-05		J0690		08/07/2017	99/99/9999	INJECTION, CEFAZOLIN SODIUM, 500 MG	CEFAZOLIN (USP,PF,LATEX-FREE) 1 GM	25 EA	VL	IJ	EA		500 MG		2	08/07/2017	99/99/9999						
63323-0356-10		J0637		07/28/2017	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LATEX-FREE) 50 MG	10 EA	VL	IV	EA		5 MG		10	07/28/2017	99/99/9999						
63323-0358-10		J0637		07/28/2017	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LATEX-FREE) 70 MG	10 EA	VL	IV	EA		5 MG		14	07/28/2017	99/99/9999						
66220-0110-01		J1190		07/25/2017	99/99/9999	INJECTION, DEXRAXOXANE HYDROCHLORIDE, PER 250 MG	TOTECT (LYOPHILIZED) 500 MG	1 EA	VL	IV	EA		250 MG		2	07/25/2017	99/99/9999						
67457-0790-05		J1953		07/24/2017	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (SDV) 100 MG/1 ML	5 ML	VL	IV	ML		10 MG		10	07/24/2017	99/99/9999						
68001-0313-56		J9025		08/16/2017	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (PF,LATEX-FREE) 100 MG	1 EA	VL	IJ	EA		1 MG		100	08/16/2017	99/99/9999						
70069-0071-10		J2310		08/09/2017	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (SINGLE-DOSE) 0.4 MG/1 ML	1 ML	VL	IJ	ML		1 MG		0.4	08/09/2017	99/99/9999						
70069-0072-10		J2310		08/09/2017	99/99/9999	INJECTION, NALOXONE HYDROCHLORIDE, PER 1 MG	NALOXONE HCL (MDV) 0.4 MG/1 ML	10 ML	VL	IJ	ML		1 MG		0.4	08/09/2017	99/99/9999						
70069-0172-10		J3420		07/31/2017	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (MDV,LATEX-FREE) 1000 MCG/1 ML	10 ML	VL	IJ	ML		1000 MCG		1	07/31/2017	99/99/9999						
70121-1408-05		J1270		07/10/2017	99/99/9999	INJECTION, DOXERCALCIFEROL, 1 MCG	DOXERCALCIFEROL (MDV) 2 MCG/1 ML	2 ML	VL	IV	ML		1 MCG		2	07/10/2017	99/99/9999						
70257-0563-01		J0475		07/24/2017	99/99/9999	INJECTION, BACLOFEN, 10 MG	LIRESAL INTRATHECAL REFILL KIT (PF) 2 MG/1 ML	20 ML	AM	IN	ML		10 MG		0.2	07/24/2017	99/99/9999						
70257-0563-02		J0475		07/24/2017	99/99/9999	INJECTION, BACLOFEN, 10 MG	LIRESAL INTRATHECAL REFILL KIT (PF) 2 MG/1 ML	20 ML	AM	IN	ML		10 MG		0.2	07/24/2017	99/99/9999						
76204-0700-25		J7614		07/17/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.20666	07/17/2017	99/99/9999						
76204-0800-25		J7614		07/17/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.63 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.42	07/17/2017	99/99/9999						
76204-0900-25		J7614		07/17/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 1.25 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.83333	07/17/2017	99/99/9999						
76204-0700-25	KO	J7614	KO	07/17/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.20666	07/17/2017	99/99/9999						
76204-0800-25	KO	J7614	KO	07/17/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.63 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.42	07/17/2017	99/99/9999						
76204-0900-25	KO	J7614	KO	07/17/2017	99/99/9999	INJECTION, LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 1.25 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.83333	07/17/2017	99/99/9999						
00002-7714-59		J1815		08/14/2017	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	HUMALOG JUNIOR KWIKPEN 100 U/1 ML	3 ML	BX	SC	ML		5 U		20	08/14/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00078-0679-19		Q0162		08/30/2017	10/17/2018	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT (3X10) 4 MG	30	EA	ST	PO	EA	1	MG	4	08/30/2017	10/17/2018						
00409-3718-01	J0290			08/07/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,PF,LATEX-FREE) 500 MG	10	EA	VL	IJ	EA	500	MG	1	08/07/2017	99/99/9999						
00409-3719-01	J0290			08/07/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,PF,LATEX-FREE) 250 MG	10	EA	VL	IJ	EA	500	MG	0.5	08/07/2017	99/99/9999						
00409-3720-01	J0290			08/01/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (PF,LATEX-FREE) 2 GM	10	EA	VL	IJ	EA	500	MG	4	08/01/2017	99/99/9999						
00409-3725-01	J0290			08/07/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,PF,LATEX-FREE) 10 GM	10	EA	VL	IJ	EA	500	MG	20	08/07/2017	99/99/9999						
00409-3726-01	J0290			08/01/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (PF,LATEX-FREE) 1 GM	10	EA	VL	IJ	EA	500	MG	2	08/01/2017	99/99/9999						
00409-4215-01	J3489			08/21/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SINGLE USE,LATEX-FREE) 5 MG/100 ML	5	ML	VL	IV	ML	1	MG	0.8	08/21/2017	99/99/9999						
00409-4228-01	J3489			08/21/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SINGLE USE,LATEX-FREE) 4 MG/100 ML	100	ML	BG	IV	ML	1	MG	0.05	08/21/2017	99/99/9999						
00409-4229-01	J3489			08/21/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SINGLE USE,LATEX-FREE) 4 MG/100 ML	100	ML	BG	IV	ML	1	MG	0.04	08/21/2017	99/99/9999						
00517-1980-05	J0500			08/30/2017	99/99/9999	INJECTION, DICYCLOMINE HCL, UP TO 20 MG	DICYCLOMINE 10 MG/1 ML	2	ML	VL	IM	ML	20	MG	0.5	08/30/2017	99/99/9999						
16729-0189-29	J7518			09/07/2017	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (DELAYED RELEASE) 360 MG	120	EA	BO	PO	EA	180	MG	2	09/07/2017	99/99/9999						
16729-0261-29	J7518			09/07/2017	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (DELAYED RELEASE) 180 MG	120	EA	BO	PO	EA	180	MG	1	09/07/2017	99/99/9999						
16729-0295-12	J9045			09/14/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	60	ML	VL	IV	ML	50	MG	0.2	09/14/2017	99/99/9999						
16729-0295-31	J9045			09/14/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	5	ML	VL	IV	ML	50	MG	0.2	09/14/2017	99/99/9999						
16729-0295-33	J9045			09/14/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	15	ML	VL	IV	ML	50	MG	0.2	09/14/2017	99/99/9999						
16729-0295-34	J9045			09/14/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	45	ML	VL	IV	ML	50	MG	0.2	09/14/2017	99/99/9999						
24338-0150-20	J3315			09/25/2017	99/99/9999	INJECTION, TRIPTORELIN PAMOATE, 3.75 MG	TRIPTODUR (LYOPHILIZED) 22.5 MG	1	EA	VL	IM	EA	3.75	MG	6	09/25/2017	99/99/9999						
47781-0588-68	J2250			08/21/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (LATEX-FREE) 1 MG/1 ML	2	ML	VL	IJ	ML	1	MG	1	08/21/2017	99/99/9999						
47781-0589-17	J2250			08/21/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (LATEX-FREE) 5 MG/1 ML	5	ML	VL	IJ	ML	1	MG	5	08/21/2017	99/99/9999						
47781-0589-91	J2250			08/21/2017	99/99/9999	INJECTION, MIDAZOLAM HYDROCHLORIDE, PER 1 MG	MIDAZOLAM HCL (LATEX-FREE) 5 MG/1 ML	10	ML	VL	IJ	ML	1	MG	5	08/21/2017	99/99/9999						
51991-0942-98	J1190			09/15/2017	99/99/9999	INJECTION, DEXRAZOXANE HYDROCHLORIDE, PER 250 MG	DEXRAZOXANE (LYOPHILIZED) 500 MG	1	EA	VL	IV	EA	250	MG	2	09/15/2017	99/99/9999						
55566-1502-01	J0725			09/15/2017	99/99/9999	INJECTION, CHORIONIC GONADOTROPIN, PER 1,000 USP UNITS	NOVAREL (10MLVIALBACTRIOSTTICH20) 5000 U	1	EA	VL	IM	EA	1000	U	5	09/15/2017	99/99/9999						
59676-0966-02	Q2050			08/28/2017	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, LIPOSOMAL, NOT OTHERWISE SPECIFIED, 10 MG	DOXORUBICIN HCL LIPOSOME 2 MG/1 ML	25	ML	VL	IV	ML	10	MG	0.2	08/28/2017	99/99/9999						
63323-0360-19	J0610			08/31/2017	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (PF,LATEX-FREE) 100 MG/1 ML	10	ML	VL	IV	ML	10	ML	0.1	08/31/2017	99/99/9999						
63323-0360-59	J0610			08/31/2017	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (PF,LATEX-FREE) 100 MG/1 ML	50	ML	VL	IV	ML	10	ML	0.1	08/31/2017	99/99/9999						
63323-0360-61	J0610			08/31/2017	99/99/9999	INJECTION, CALCIUM GLUCONATE, PER 10 ML	CALCIUM GLUCONATE (PF,LATEX-FREE) 100 MG/1 ML	100	ML	VL	IV	ML	10	ML	0.1	08/31/2017	99/99/9999						
67457-0856-20	J0153			08/31/2017	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (1X20ML,USP,SDV,PF) 3 MG/1 ML	20	ML	VL	IV	ML	1	MG	3	08/31/2017	99/99/9999						
67457-0857-30	J0153			08/31/2017	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (1X30ML,USP,SDV,PF) 3 MG/1 ML	30	ML	VL	IV	ML	1	MG	3	08/31/2017	99/99/9999						
70069-0101-05	J2800			09/12/2017	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL 100 MG/1 ML	10	ML	VL	IJ	ML	10	ML	0.1	09/12/2017	99/99/9999						
70069-0101-25	J2800			09/12/2017	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL 100 MG/1 ML	10	ML	VL	IJ	ML	10	ML	0.1	09/12/2017	99/99/9999						
70121-1630-01	J9340			09/11/2017	99/99/9999	INJECTION, THIOTEPA, 15 MG	TEPADINA 15 MG	1	EA	VL	IJ	EA	15	MG	1	09/11/2017	99/99/9999						
70121-1631-01	J9340			09/11/2017	99/99/9999	INJECTION, THIOTEPA, 15 MG	TEPADINA 100 MG	1	EA	VL	IJ	EA	15	MG	6.66666	09/11/2017	99/99/9999						
75987-0080-10	J2507			08/25/2017	99/99/9999	INJECTION, PEGLOTICASE, 1 MG	KRYSTEXXA (LATEX-FREE) 8 MG/1 ML	1	ML	VL	IV	ML	1	MG	8	08/25/2017	99/99/9999						
00487-9007-60	A4216			03/13/2017	99/99/9999	STERILE WATER, SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (30 x 4ML,PF) 7%	4	ML	VL	IH	ML	10	ML	0.1	03/13/2017	99/99/9999						
00008-4510-01	J9300			09/01/2017	12/31/2017	INJECTION, GEMTUZUMAB OZOGAMICIN, 5 MG	MYLOTARG (PF,LYOPHILIZED CAKE) 4.5 MG	1	EA	VL	IV	EA	5	MG	0.9	09/01/2017	12/31/2017						
00078-0880-19	Q0162			09/19/2017	10/17/2018	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN ODT 8 MG	30	EA	ST	PO	EA	1	MG	8	09/19/2017	10/17/2018						
00143-9209-10	J2400			09/28/2017	99/99/9999	INJECTION, CHLOROPROCAINE HYDROCHLORIDE, PER 30 ML	CHLOROPROCAINE HCL (400MG/20ML, SDV, USP,PF) 2%	20	ML	VL	IJ	ML	30	ML	0.03333	09/28/2017	99/99/9999						
00143-9210-10	J2400			09/28/2017	99/99/9999	INJECTION, CHLOROPROCAINE HYDROCHLORIDE, PER 30 ML	CHLOROPROCAINE HCL (600MG/20ML, SDV, USP,PF) 3%	20	ML	VL	IJ	ML	30	ML	0.03333	09/28/2017	99/99/9999						
00169-3201-11	J1817			09/29/2017	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	FIASP 100 U/1 ML	10	ML	VL	IJ	ML	50	U	2	09/29/2017	99/99/9999						
00169-3204-15	J1815			09/29/2017	99/99/9999	INJECTION, INSULIN, PER 5 UNITS	FIASP FLEXTOUCH (PREFILLED PEN, SU) 100 U/1 ML	3	ML	CT	SC	ML	5	U	20	09/29/2017	99/99/9999						
00378-6960-93	J1595			10/04/2017	99/99/9999	INJECTION, GLATIRAMER ACETATE, 20 MG	GLATIRAMER ACETATE 20 MG/1 ML	1	ML	SR	SC	ML	20	MG	1	10/04/2017	99/99/9999						
00548-9601-00	J2710			10/10/2017	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 0.5 MG/1 ML	10	ML	VL	IV	ML	0.5	MG	1	10/10/2017	99/99/9999						
00548-9602-00	J2710			10/10/2017	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 1 MG/1 ML	10	ML	VL	IV	ML	0.5	MG	2	10/10/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
15054-0043-01		J9205		10/16/2017	99/99/9999	INJECTION, IRINOTECAN LIPOSOME, 1 MG	ONIVYDE (SDV) 4.3 MG/1 ML	10 ML	VL	IV	ML		1 MG		4.3	10/16/2017	99/99/9999						
16714-0725-01		J9206		11/01/2017	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV,PF,LATEX-FREE) 20 MG/1 ML	2 ML	VL	IV	ML		20 MG		1	11/01/2017	99/99/9999						
16714-0726-01		J9206		11/01/2017	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV,PF,LATEX-FREE) 20 MG/1 ML	5 ML	VL	IV	ML		20 MG		1	11/01/2017	99/99/9999						
16714-0727-01		J9263		11/06/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (1X10ML,SINGLE DOSE,PF) 5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		10	11/06/2017	99/99/9999						
16714-0728-01		J9263		11/06/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (1X20ML,SINGLE DOSE,PF) 5 MG/1 ML	20 ML	VL	IV	ML		0.5 MG		10	11/06/2017	99/99/9999						
16714-0742-01		Q2050		10/04/2017	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, LIPOSOMAL, NOT OTHERWISE SPECIFIED, 10 MG	DOXORUBICIN HCL LIPOSOME 2 MG/1 ML	10 ML	VL	IV	ML		10 MG		0.2	10/04/2017	99/99/9999						
16714-0856-01		Q2050		10/04/2017	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, LIPOSOMAL, NOT OTHERWISE SPECIFIED, 10 MG	DOXORUBICIN HCL LIPOSOME 2 MG/1 ML	25 ML	VL	IV	ML		10 MG		0.2	10/04/2017	99/99/9999						
16729-0242-31		J3489		10/04/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SDV) 4 MG/5 ML	5 ML	VL	IV	ML		1 MG		0.8	10/04/2017	99/99/9999						
47781-0578-07		J1190		09/14/2017	99/99/9999	INJECTION, DEXRAZOXANE HYDROCHLORIDE, PER 250 MG	DEXRAZOXANE (SDV,W/DILUENT) 500 MG	1 EA	VL	IV	EA		250 MG		2	09/14/2017	99/99/9999						
47781-0583-68		J1885		10/10/2017	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV,25X1ML,PF) 15 MG/1 ML	1 ML	VL	IJ	ML		15 MG		1	10/10/2017	99/99/9999						
47781-0584-68		J1885		10/10/2017	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (SDV,25X1ML,PF) 30 MG/1 ML	1 ML	VL	IJ	ML		15 MG		2	10/10/2017	99/99/9999						
47781-0609-25		J9060		10/09/2017	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (PF,LATEX-FREE) 1 MG/1 ML	50 ML	VL	IV	ML		10 MG		0.1	10/09/2017	99/99/9999						
47781-0610-23		J9060		10/09/2017	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (PF,LATEX-FREE) 1 MG/1 ML	100 ML	VL	IV	ML		10 MG		0.1	10/09/2017	99/99/9999						
51991-0218-98		J9263		09/27/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (SINGLE-USE,PF) 50 MG	1 EA	VL	IV	EA		0.5 MG		100	09/27/2017	99/99/9999						
51991-0219-98		J9263		09/27/2017	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (SINGLE-USE,PF) 100 MG	1 EA	VL	IV	EA		0.5 MG		200	09/27/2017	99/99/9999						
51991-0797-98		J9025		09/25/2017	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (PF,LYOPHILIZED) 100 MG	1 EA	VL	IJ	EA		1 MG		100	09/25/2017	99/99/9999						
58406-0456-01		J1438		11/17/2017	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (MINI,PF) 50 MG/1 ML	0.98 ML	BX	SC	ML		25 MG		2	11/17/2017	99/99/9999						
58406-0456-04		J1438		11/17/2017	99/99/9999	INJECTION, ETANERCEPT, 25 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	ENBREL (MINI,PF) 50 MG/1 ML	0.98 ML	BX	SC	ML		25 MG		2	11/17/2017	99/99/9999						
66794-0151-01		J0476		11/01/2017	99/99/9999	INJECTION, BACLOFEN, 50 MCG FOR INTRATHECAL TRIAL	BACLOFEN (1X1ML,SINGLE USE) 0.05 MG/1 ML	1 ML	SR	IN	ML		50 MCG		1	11/01/2017	99/99/9999						
67457-0323-25		J2280		10/03/2017	99/99/9999	INJECTION, MOXIFLOXACIN, 100 MG	MOXIFLOXACIN HCL (FLEXIBAG,LATEX-FREE) 400 MG/250 ML	250 ML	BG	IV	ML		100 MG		0.016	10/03/2017	99/99/9999						
67457-0831-50		J0637		09/29/2017	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (SDV,PF,LYOPHILIZED) 50 MG	1 EA	VL	IV	EA		5 MG		10	09/29/2017	99/99/9999						
68462-0583-85		J8501		10/13/2017	99/99/9999	APREPITANT, ORAL, 5 MG	APREPITANT (1X5,HARD GELATIN) 40 MG	5 EA	ST	PO	EA		5 MG		8	10/13/2017	99/99/9999						
68462-0584-58		J8501		10/13/2017	99/99/9999	APREPITANT, ORAL, 5 MG	APREPITANT (2-DAY PACK,HARD GELATIN) 80 MG	2 EA	ST	PO	EA		5 MG		16	10/13/2017	99/99/9999						
68462-0584-76		J8501		10/13/2017	99/99/9999	APREPITANT, ORAL, 5 MG	APREPITANT (1X6,HARD GELATIN) 80 MG	6 EA	ST	PO	EA		5 MG		16	10/13/2017	99/99/9999						
68462-0585-76		J8501		10/13/2017	99/99/9999	APREPITANT, ORAL, 5 MG	APREPITANT (1X6,HARD GELATIN) 125 MG	6 EA	ST	PO	EA		5 MG		25	10/13/2017	99/99/9999						
69784-0205-60		J7631		10/18/2017	99/99/9999	CROMOLYN SODIUM, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM 10 MG/1 ML	2 ML	VL	IH	ML		10 MG		1	10/18/2017	99/99/9999						
70860-0206-50		J9060		09/15/2017	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (PF,LATEX-FREE) 1 MG/1 ML	50 ML	VL	IV	ML		10 MG		0.1	09/15/2017	99/99/9999						
70860-0206-51		J9060		09/15/2017	99/99/9999	INJECTION, CISPLATIN, POWDER OR SOLUTION, 10 MG	CISPLATIN (PF,LATEX-FREE) 1 MG/1 ML	100 ML	VL	IV	ML		10 MG		0.1	09/15/2017	99/99/9999						
68462-0502-01		J7500		11/20/2008	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	AZATHIOPRINE 50 MG	100 EA	BO	PO	EA		50 MG		1	11/20/2008	99/99/9999						
69784-0205-60	KO	J7631	KO	10/18/2017	99/99/9999	CROMOLYN SODIUM, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER 10 MILLIGRAMS	CROMOLYN SODIUM 10 MG/1 ML	2 ML	VL	IH	ML		10 MG		1	10/18/2017	99/99/9999						
64011-0247-02		J1726		01/01/2018	99/99/9999	INJECTION, HYDROXYPROGESTERONE CAPROATE, (MAKENA), 10 MG	MAKENA 250 MG/1 ML	1 ML	VL	IM	ML		10 MG		25	01/01/2018	99/99/9999						
62559-0540-15		J1729		01/01/2018	99/99/9999	INJECTION, HYDROXYPROGESTERONE CAPROATE, NOT OTHERWISE SPECIFIED, 10 MG	HYDROXYPROGESTERONE CAPROATE 250 MG/1 ML	5 ML	VL	IM	ML		10 MG		25	01/01/2018	99/99/9999						
00008-4510-01		J9203		01/01/2018	99/99/9999	INJECTION, GEMTUZUMAB OZOGAMICIN, 0.1 MG	MYLOTARG (PF,LYOPHILIZED CAKE) 4.5 MG	1 EA	VL	IV	EA		0.1 MG		45	01/01/2018	99/99/9999						
64980-0336-05	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	5 EA	BO	PO	EA		20 MG		7	05/25/2017	99/99/9999						
64980-0336-14	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	14 EA	BO	PO	EA		20 MG		7	05/25/2017	99/99/9999						
64980-0337-05	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	5 EA	BO	PO	EA		20 MG		9	05/25/2017	99/99/9999						
00115-1687-74		J7626		11/10/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.25 MG/2 ML	2 ML	AM	IH	ML		0.5 MG		0.25	11/10/2017	99/99/9999						
00115-1689-74		J7626		11/07/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.5 MG/2 ML	2 ML	AM	IH	ML		0.5 MG		0.5	11/07/2017	99/99/9999						
00338-9586-24		J2001		10/02/2017	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL-DEXTROSE 5%-0.4%	500 ML	BG	IV	ML		10 MG		0.4	10/02/2017	99/99/9999						
00338-9590-30		J2001		10/02/2017	99/99/9999	INJECTION, LIDOCAINE HCL FOR INTRAVENOUS INFUSION, 10 MG	LIDOCAINE HCL-DEXTROSE 5%-0.4%	250 ML	BG	IV	ML		10 MG		0.4	10/02/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
00641-6174-10	J2354			10/20/2017	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS INJECTION, 25 MCG	OCTREOTIDE ACETATE 50 MCG/1 ML	1	ML	VL	IJ	ML	25	MCG	2	10/20/2017	99/99/9999							
00641-6175-10	J2354			10/20/2017	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS INJECTION, 25 MCG	OCTREOTIDE ACETATE 100 MCG/1 ML	1	ML	VL	IJ	ML	25	MCG	4	10/20/2017	99/99/9999							
00641-6176-10	J2354			10/20/2017	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS INJECTION, 25 MCG	OCTREOTIDE ACETATE 500 MCG/1 ML	1	ML	VL	IJ	ML	25	MCG	20	10/20/2017	99/99/9999							
00641-6177-01	J2354			10/20/2017	99/99/9999	INJECTION, OCTREOTIDE, NON-DEPOT FORM FOR SUBCUTANEOUS OR INTRAVENOUS INJECTION, 25 MCG	OCTREOTIDE ACETATE 200 MCG/1 ML	5	ML	VL	IJ	ML	25	MCG	8	10/20/2017	99/99/9999							
00641-6178-01	J2354			10/20/2017	99/99/9999	INJECTION, 25 MCG	OCTREOTIDE ACETATE 1000 MCG/1 ML	5	ML	VL	IJ	ML	25	MCG	40	10/20/2017	99/99/9999							
00641-6182-02	J2360			11/07/2017	99/99/9999	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	ORPHENADRINE CITRATE 30 MG/1 ML	2	ML	VL	IJ	ML	60	MG	0.5	11/07/2017	99/99/9999							
17478-0380-20	J1230			11/13/2017	99/99/9999	INJECTION, METHADONE HCL, UP TO 10 MG	METHADONE HCL 10 MG/1 ML	20	ML	VL	IJ	ML	10	MG	1	11/13/2017	99/99/9999							
43598-0309-20	J9027			11/08/2017	99/99/9999	INJECTION, CLOFARABINE, 1 MG DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	CLOFARABINE (PF) 1 MG/1 ML	20	ML	VL	IV	ML	1	MG	1	11/08/2017	99/99/9999							
50580-0226-50	Q0163			10/30/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	BENADRYL ALLERGY (ULTRATAB) 25 MG ZOLEDRONIC ACID (1X100ML SINGLE USE) 5 MG/100 ML	100	EA	BX	PO	EA	50	MG	0.5	10/30/2017	99/99/9999							
51991-0064-98	J3489			10/30/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SINGLE-USE) 4 MG/5 ML	5	ML	VL	IV	ML	1	MG	0.8	10/30/2017	99/99/9999							
51991-0065-98	J3489			10/30/2017	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	VANCOMYCIN HCL (PHARMACY BULK PACKAGE) 10 MG	1	EA	VL	IV	EA	500	MG	20	10/26/2017	99/99/9999							
63323-0314-68	J3370			10/26/2017	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	ARGATROBAN (SDV,PF) 100 MG/1 ML	2.5	ML	VL	IV	ML	1	MG	100	11/14/2017	99/99/9999							
67457-0212-02	J0883			11/14/2017	99/99/9999	INJECTION, ORPHENADRINE CITRATE, UP TO 60 MG	CLOFARABINE (PF) 1 MG/1 ML	20	ML	VL	IV	ML	1	MG	1	11/06/2017	99/99/9999							
67457-0546-20	J9027			11/06/2017	99/99/9999	INJECTION, CLOFARABINE, 1 MG	CASPOFUNGIN ACETATE (PF,LYOPHILIZED) 70 MG	1	EA	VL	IV	EA	5	MG	14	11/15/2017	99/99/9999							
67457-0832-70	J0637			11/15/2017	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/1 ML	1	ML	VL	IJ	ML	15	MG	2	11/16/2017	99/99/9999							
69543-0386-25	J1885			11/16/2017	99/99/9999	INJECTION, CLOFARABINE, 1 MG	CLOFARABINE (PF) 1 MG/1 ML	20	ML	VL	IV	ML	1	MG	1	11/06/2017	99/99/9999							
70121-1236-01	J9027			11/06/2017	99/99/9999	INJECTION, OCRELIZUMAB, 1 MG	OCREVUS (SDV,PF) 30 MG/1 ML	10	ML	VL	IV	ML	1	MG	30	01/01/2018	99/99/9999							
50242-0150-01	J2350			01/01/2018	99/99/9999	INJECTION, OCRELIZUMAB, 1 MG	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (PF) 10%	30	ML	VL	IH	ML	1	GM	0.1	07/14/2014	99/99/9999						
63323-0691-30	J7608			07/14/2014	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (PF) 20%	4	ML	VL	PO	ML	1	GM	0.2	12/10/2013	99/99/9999							
63323-0694-04	J7608			12/10/2013	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	TEMOZOLOMIDE 5 MG	14	EA	BO	PO	EA	5	MG	1	05/25/2017	99/99/9999							
64980-0333-14	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 100 MG	14	EA	BO	PO	EA	100	MG	1	05/25/2017	99/99/9999							
64980-0335-14	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 180 MG	14	EA	BO	PO	EA	20	MG	9	05/25/2017	99/99/9999							
64980-0337-14	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 250 MG	5	EA	BO	PO	EA	250	MG	1	05/25/2017	99/99/9999							
64980-0338-05	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	CUVITRU (1GM,PF,LATEX-FREE) 20%	5	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-01	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	CUVITRU (1GM,INNER PACK NDC,PF) 20%	5	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-02	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	CUVITRU (2GM,PF,LATEX-FREE) 20%	10	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-03	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	CUVITRU (2GM,INNER PACK NDC,PF) 20%	10	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-04	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	CUVITRU (4GM,PF,LATEX-FREE) 20%	20	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-05	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	CUVITRU (4GM,INNER PACK NDC,PF) 20%	20	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-06	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	CUVITRU (8GM,PF,LATEX-FREE) 20%	40	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-07	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	CUVITRU (8GM,INNER PACK NDC,PF) 20%	40	ML	VL	SC	ML	100	MG	2	01/01/2018	99/99/9999							
00944-2850-08	J1555			01/01/2018	99/99/9999	INJECTION, IMMUNE GLOBULIN (CUVITRU), 100 MG	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.25 MG/2 ML	2	ML	AM	IH	ML	0.5	MG	0.25	11/10/2017	99/99/9999						
00115-1687-74	KO	J7626	KO	11/10/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,SINGLE-DOSE) 0.5 MG/2 ML	2	ML	AM	IH	ML	0.5	MG	0.5	11/07/2017	99/99/9999							
00115-1689-74	KO	J7626	KO	11/07/2017	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (PF) 10%	30	ML	VL	IH	ML	1	GM	0.1	07/14/2014	99/99/9999						
63323-0691-30	KO	J7608	KO	07/14/2014	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	ACETYLCYSTEINE (PF) 20%	4	ML	VL	PO	ML	1	GM	0.2	12/10/2013	99/99/9999							
63323-0694-04	KO	J7608	KO	12/10/2013	99/99/9999	ACETYLCYSTEINE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER GRAM	HYDROXYPROGESTERONE CAPROATE 250 MG/1 ML	5	ML	VL	IM	ML	10	MG	25	07/01/2017	12/31/2017							
62559-0540-15	Q9985			07/01/2017	12/31/2017	INJECTION, HYDROXYPROGESTERONE CAPROATE, NOT OTHERWISE SPECIFIED, 10 MG	MAKENA 250 MG/1 ML	1	ML	VL	IM	ML	10	MG	25	07/01/2017	12/31/2017							
64011-0247-02	Q9986			07/01/2017	12/31/2017	INJECTION, HYDROXYPROGESTERONE CAPROATE, (MAKENA), 10 MG	TEMOZOLOMIDE 5 MG	5	EA	BO	PO	EA	5	MG	1	05/25/2017	99/99/9999							
64980-0333-05	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 20 MG	5	EA	BO	PO	EA	20	MG	1	05/25/2017	99/99/9999							
64980-0334-05	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14	EA	BO	PO	EA	20	MG	1	05/25/2017	99/99/9999							
64980-0334-14	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 100 MG	5	EA	BO	PO	EA	100	MG	1	05/25/2017	99/99/9999							
64980-0335-05	None			05/25/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	DIHYDROERGOTAMINE MESYLATE 1 MG/1 ML	1	ML	AM	IJ	ML	1	MG	1	11/28/2017	99/99/9999							
00143-9273-10	J1110			11/28/2017	99/99/9999	INJECTION, DIHYDROERGOTAMINE MESYLATE, PER 1 MG																		

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00143-9553-01	J0640			06/14/2017	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (PF,LATEX-FREE) 200 MG	1 EA	VL	IJ	EA		50 MG		4	06/14/2017	99/99/9999						
00143-9554-01	J0640			06/14/2017	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (PF,LATEX-FREE) 100 MG	1 EA	VL	IJ	EA		50 MG		2	06/14/2017	99/99/9999						
00143-9555-01	J0640			06/14/2017	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (PF,LATEX-FREE) 50 MG	1 EA	VL	IJ	EA		50 MG		1	06/14/2017	99/99/9999						
00143-9830-01	J9260			11/20/2017	99/99/9999	METHOTREXATE SODIUM, 50 MG	METHOTREXATE (SINGLE USE VIAL,PF) 1 GM	1 EA	VL	IJ	EA		50 MG		20	11/20/2017	99/99/9999						
00409-0368-01	J9171			12/08/2017	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL 20 MG/1 ML	8 ML	VL	IV	ML		1 MG		20	12/08/2017	99/99/9999						
00517-0650-02	J1439			12/01/2017	99/99/9999	INJECTION, FERRIC CARBOXYMALTOSE, 1 MG	INJECTAFER (2 X15ML) 50 MG/1 ML	15 ML	VL	IV	ML		1 MG		50	12/01/2017	99/99/9999						
00781-3481-92	J3243			11/30/2017	99/99/9999	INJECTION, TIGECYCLINE, 1 MG	TIGECYCLINE (10ML VIALS,PF) 50 MG	10 EA	VL	IV	EA		1 MG		50	11/30/2017	99/99/9999						
16714-0749-01	J0894			12/19/2017	99/99/9999	INJECTION, DECATIBINE, 1 MG	DECATIBINE (LYOPHILIZED) 50 MG	1 EA	VL	IV	EA		1 MG		50	12/19/2017	99/99/9999						
55513-0073-30	J0604			04/05/2004	99/99/9999	CINACALCET, ORAL, 1 MG, (FOR ESRD ON DIALYSIS)	SENSIPAR (FILM COATED) 30 MG	30 EA	BO	PO	EA		1 MG		30	04/05/2004	99/99/9999						
17478-0902-10	J1327			11/20/2017	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (SDV) 2 MG/1 ML	10 ML	VL	IV	ML		5 MG		0.4	11/20/2017	99/99/9999						
17478-0902-90	J1327			11/20/2017	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE (SDV) 2 MG/1 ML	100 ML	VL	IV	ML		5 MG		0.4	11/20/2017	99/99/9999						
17478-0903-90	J1327			11/20/2017	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE 0.75 MG/1 ML	100 ML	VL	IV	ML		5 MG		0.15	11/20/2017	99/99/9999						
25021-0831-01	J1631			12/11/2017	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (SDV,LATEX-FREE) 50 MG/1 ML	1 ML	VL	IM	ML		50 MG		1	12/11/2017	99/99/9999						
25021-0833-01	J1631			12/11/2017	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (SDV,LATEX-FREE) 100 MG/1 ML	1 ML	VL	IM	ML		50 MG		2	12/11/2017	99/99/9999						
25021-0834-05	J1631			12/11/2017	99/99/9999	INJECTION, HALOPERIDOL DECAANOATE, PER 50 MG	HALOPERIDOL DECAANOATE (SDV,LATEX-FREE) 100 MG/1 ML	5 ML	VL	IM	ML		50 MG		2	12/11/2017	99/99/9999						
47781-0585-68	J1885			11/22/2017	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (USP,25X2ML,SDV) 30 MG/1 ML	2 ML	VL	IM	ML		15 MG		2	11/22/2017	99/99/9999						
47781-0613-07	J0637			12/11/2017	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LATEX-FREE) 50 MG	1 EA	VL	IV	EA		5 MG		10	12/11/2017	99/99/9999						
47781-0614-07	J0637			12/11/2017	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LATEX-FREE) 70 MG	1 EA	VL	IV	EA		5 MG		14	12/11/2017	99/99/9999						
55513-0740-01	J0606			10/09/2017	99/99/9999	INJECTION, ETELCALCETIDE, 0.1 MG	PARSABIV (PF) 2.5 MG/0.5 ML	0.5 ML	VL	IV	ML		0.1 MG		50	10/09/2017	99/99/9999						
55513-0740-10	J0606			10/09/2017	99/99/9999	INJECTION, ETELCALCETIDE, 0.1 MG	PARSABIV (PF) 2.5 MG/0.5 ML	0.5 ML	VL	IV	ML		0.1 MG		50	10/09/2017	99/99/9999						
55513-0741-01	J0606			10/09/2017	99/99/9999	INJECTION, ETELCALCETIDE, 0.1 MG	PARSABIV (PF) 5 MG/1 ML	1 ML	VL	IV	ML		0.1 MG		50	10/09/2017	99/99/9999						
55513-0741-10	J0606			10/09/2017	99/99/9999	INJECTION, ETELCALCETIDE, 0.1 MG	PARSABIV (PF) 5 MG/1 ML	1 ML	VL	IV	ML		0.1 MG		50	10/09/2017	99/99/9999						
55513-0742-01	J0606			10/09/2017	99/99/9999	INJECTION, ETELCALCETIDE, 0.1 MG	PARSABIV (SDV,PF) 10 MG/2 ML	2 ML	VL	IV	ML		0.1 MG		50	10/09/2017	99/99/9999						
55513-0742-10	J0606			10/09/2017	99/99/9999	INJECTION, ETELCALCETIDE, 0.1 MG	PARSABIV (SDV,PF) 10 MG/2 ML	2 ML	VL	IV	ML		0.1 MG		50	10/09/2017	99/99/9999						
63323-0708-00	J0290			12/01/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN SODIUM 500 MG	10 EA	VL	IJ	EA		500 MG		1	12/01/2017	99/99/9999						
63459-0601-06	J9017			12/05/2017	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	TRISENOX (PF) 2 MG/1 ML	6 ML	VL	IV	ML		1 MG		2	12/05/2017	99/99/9999						
65862-0942-03	J7612			12/07/2017	99/99/9999	INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, CONCENTRATED FORM, 0.5 MG	LEVALBUTEROL (CONCENTRATE,PF) 1.25 MG/0.5 ML	30 EA	VL	IH	EA		0.5 MG		2.5	12/07/2017	99/99/9999						
65862-0943-24	J7614			12/07/2017	99/99/9999	INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.20666	12/07/2017	99/99/9999						
65862-0944-24	J7614			12/07/2017	99/99/9999	INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 0.63 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.42	12/07/2017	99/99/9999						
65862-0945-24	J7614			12/07/2017	99/99/9999	INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (2X12 POUCHES,PF) 1.25 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.83333	12/07/2017	99/99/9999						
66794-0155-01	J0475			01/01/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	GABLOFEN (1X20ML,SINGLE USE) 0.5 MG/1 ML	20 ML	SR	IN	ML		10 MG		0.05	01/01/2018	99/99/9999						
66794-0157-01	J0475			01/01/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	GABLOFEN (1X20ML,SINGLE USE) 2 MG/1 ML	20 ML	SR	IN	ML		10 MG		0.2	01/01/2018	99/99/9999						
66993-0489-83	J9120			12/07/2017	99/99/9999	INJECTION, DACTINOMYCIN, 0.5 MG	DACTINOMYCIN (SDV,PF,LYOPHILIZED) 0.5 MG	1 EA	VL	IV	EA		0.5 MG		1	12/07/2017	99/99/9999						
67457-0348-10	J0295			12/01/2017	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM 1 GM-0.5 GM	10 EA	VL	IJ	EA		1.5 GM		1	12/01/2017	99/99/9999						
67457-0886-05	J1729			09/22/2017	99/99/9999	INJECTION, HYDROXYPROGESTERONE CAPROATE, NOT OTHERWISE SPECIFIED, 10 MG	HYDROXYPROGESTERONE CAPROATE 250 MG/1 ML	5 ML	VL	IM	ML		10 MG		25	09/22/2017	99/99/9999						
67457-0893-08	J0594			11/21/2017	99/99/9999	INJECTION, BUSULFAN, 1 MG	BUSULFAN (8X10ML,SINGLE-USE) 6 MG/1 ML	10 ML	VL	IV	ML		1 MG		6	11/21/2017	99/99/9999						
67877-0537-07	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	5 EA	BO	PO	EA		5 MG		1	04/26/2017	99/99/9999						
67877-0537-14	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	14 EA	BO	PO	EA		5 MG		1	04/26/2017	99/99/9999						
67877-0538-07	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5 EA	BO	PO	EA		20 MG		1	04/26/2017	99/99/9999						
67877-0538-14	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14 EA	BO	PO	EA		20 MG		1	04/26/2017	99/99/9999						
67877-0539-07	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5 EA	BO	PO	EA		100 MG		1	04/26/2017	99/99/9999						
67877-0539-14	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14 EA	BO	PO	EA		100 MG		1	04/26/2017	99/99/9999						
67877-0540-07	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	5 EA	BO	PO	EA		20 MG		7	04/26/2017	99/99/9999						
67877-0540-14	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	14 EA	BO	PO	EA		20 MG		7	04/26/2017	99/99/9999						
67877-0541-07	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	5 EA	BO	PO	EA		20 MG		9	04/26/2017	99/99/9999						
67877-0541-14	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	14 EA	BO	PO	EA		20 MG		9	04/26/2017	99/99/9999						
67877-0542-07	None			04/26/2017	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE 250 MG	5 EA	BO	PO	EA		250 MG		1	04/26/2017	99/99/9999						
69097-0285-37	J0894			11/17/2017	99/99/9999	INJECTION, DECATIBINE, 1 MG	DECATIBINE (LYOPHILIZED) 50 MG	1 EA	VL	IV	EA		1 MG		50	11/17/2017	99/99/9999						
70121-1049-05	J3301			12/12/2017	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE 40 MG/1 ML	1 ML	VL	IJ	ML		10 MG		4	12/12/2017	99/99/9999						
70121-1168-01	J3301			12/12/2017	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE 40 MG/1 ML	5 ML	VL	IJ	ML		10 MG		4	12/12/2017	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
70121-1169-01		J3301		12/12/2017	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE 40 MG/1 ML	10	ML	VL	IJ	ML	10	MG	4	12/12/2017	99/99/9999							
47335-0235-83		None		12/01/2017	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	100	EA	BO	PO	EA	2.5	MG	1	12/01/2017	99/99/9999							
47335-0235-96		None		12/01/2017	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE 2.5 MG	36	EA	BO	PO	EA	2.5	MG	1	12/01/2017	99/99/9999							
55513-0074-30		J0604		04/05/2004	99/99/9999	CINACALCET, ORAL, 1 MG, (FOR ESRD ON DIALYSIS)	SENSIPAR (FILM COATED) 60 MG	30	EA	BO	PO	EA	1	MG	60	04/05/2004	99/99/9999							
55513-0075-30		J0604		04/05/2004	99/99/9999	CINACALCET, ORAL, 1 MG, (FOR ESRD ON DIALYSIS)	SENSIPAR (FILM COATED) 90 MG	30	EA	BO	PO	EA	1	MG	90	04/05/2004	99/99/9999							
63323-0721-10		J9041		11/17/2017	12/31/2018	INJECTION, BORTEZOMIB, 0.1 MG	BORTEZOMIB, (SDV,LATEX-FREE) 3.5 MG	1	EA	VL	IV	EA	0.1	MG	35	11/17/2017	12/31/2018							
67877-0568-60		Q0167		09/22/2017	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (SOFT GELATIN) 2.5 MG	60	EA	BO	PO	EA	2.5	MG	1	09/22/2017	99/99/9999							
67877-0569-60		Q0167		09/22/2017	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (SOFT GELATIN) 5 MG	60	EA	BO	PO	EA	2.5	MG	2	09/22/2017	99/99/9999							
67877-0570-60		Q0167		09/22/2017	99/99/9999	DRONABINOL, 2.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	DRONABINOL (SOFT GELATIN) 10 MG	60	EA	BO	PO	EA	2.5	MG	4	09/22/2017	99/99/9999							
65862-0943-24	KO	J7614	KO	12/07/2017	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (PF) 0.31 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.20666	12/07/2017	99/99/9999							
65862-0944-24	KO	J7614	KO	12/07/2017	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (2X12 POUCHES,PF) 0.63 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.42	12/07/2017	99/99/9999							
65862-0945-24	KO	J7614	KO	12/07/2017	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (2X12 POUCHES,PF) 1.25 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.83333	12/07/2017	99/99/9999							
00003-3734-13		J9299		01/02/2018	99/99/9999	INJECTION, NIVOLUMAB, 1 MG	OPDIVO (PF) 10 MG/1 ML	24	ML	VL	IV	ML	1	MG	10	01/02/2018	99/99/9999							
00069-0983-01		J9315		01/04/2018	99/99/9999	INJECTION, ROMIDEPSIN, 1 MG	ROMIDEPSIN (W/DILUENT) 10 MG	1	EA	VL	IV	EA	1	MG	10	01/04/2018	99/99/9999							
00078-0676-15		Q0162		01/11/2018	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ZOFRAN (FILM COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	01/11/2018	99/99/9999							
00115-9930-78		J7614		01/09/2018	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (PF) 0.31 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.20666	01/09/2018	99/99/9999							
00115-9931-78		J7614		01/09/2018	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (PF) 0.63 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.42	01/09/2018	99/99/9999							
00115-9932-78		J7614		01/09/2018	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (PF) 1.25 MG/3 ML	3	ML	VL	IH	ML	0.5	MG	0.83333	01/09/2018	99/99/9999							
00143-9530-01		J9208		01/11/2018	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (S.D.V, 1X60ML,PF) 3 GM/60 ML	60	ML	VL	IV	ML	1	GM	0.05	01/11/2018	99/99/9999							
00143-9531-01		J9208		12/14/2017	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE (S.D.V, 1X20ML) 1 GM/20 ML	20	ML	VL	IV	ML	1	GM	0.05	12/14/2017	99/99/9999							
00406-8020-03		J0574		01/05/2018	99/99/9999	BUPRENORPHINE/NALOXONE, ORAL, GREATER THAN 6 MG, BUT LESS THAN OR EQUAL TO 10 MG BUPRENORPHINE	BUPRENORPHINE-NALOXONE (LEMON) 8 MG-2 MG	30	EA	BO	SL	EA	8	MG	1	01/05/2018	99/99/9999							
00409-1007-01		J2501		01/01/2018	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL (LATEX-FREE) 0.002 MG/1 ML	1	ML	VL	IV	ML	1	MCG	2	01/01/2018	99/99/9999							
00548-5400-00		J1050		01/15/2018	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	MEDROXYPROGESTERONE ACETATE 150 MG/1 ML	1	ML	VL	IM	ML	1	MG	150	01/15/2018	99/99/9999							
00548-5400-25		J1050		02/05/2018	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	MEDROXYPROGESTERONE ACETATE 150 MG/1 ML	1	ML	VL	IM	ML	1	MG	150	02/05/2018	99/99/9999							
00548-5701-00		J1050		01/15/2018	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	MEDROXYPROGESTERONE ACETATE (PRE-FILLED SYRINGE) 150 MG/1 ML	1	ML	SR	IM	ML	1	MG	150	01/15/2018	99/99/9999							
13533-0705-01		J0256		01/09/2018	99/99/9999	INJECTION, ALPHA 1 PROTEINASE INHIBITOR (HUMAN), NOT OTHERWISE SPECIFIED, 10 MG	PROLASTIN-C (APPROX 1000MG,PF) 1 MG	1	EA	VL	IV	EA	10	MG	0.1	01/09/2018	99/99/9999							
16729-0391-30		J9201		01/15/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 100 MG/1 ML	2	ML	VL	IV	ML	200	MG	0.5	01/15/2018	99/99/9999							
16729-0419-30		J9201		01/15/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 100 MG/1 ML	10	ML	VL	IV	ML	200	MG	0.5	01/15/2018	99/99/9999							
16729-0423-33		J9201		01/15/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 100 MG/1 ML	15	ML	VL	IV	ML	200	MG	0.5	01/15/2018	99/99/9999							
16729-0426-05		J9201		01/15/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 100 MG/1 ML	20	ML	VL	IV	ML	200	MG	0.5	01/15/2018	99/99/9999							
43598-0392-48		J9245		12/21/2017	99/99/9999	INJECTION, MELPHALAN HYDROCHLORIDE, 50 MG	MELPHALAN HYDROCHLORIDE (W/ 10ML DILUENT) 50 MG	1	EA	VL	IV	EA	50	MG	1	12/21/2017	99/99/9999							
43598-0678-11		J9025		12/21/2017	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE 100 MG	1	EA	VL	IJ	EA	1	MG	100	12/21/2017	99/99/9999							
45963-0640-77		J0594		01/04/2018	99/99/9999	INJECTION, BUSULFAN, 1 MG	BUSULFAN (8X10ML,SINGLE-USE,PF) 6 MG/1 ML	10	ML	VL	IV	ML	1	MG	6	01/04/2018	99/99/9999							
51224-0012-20		J2760		01/31/2018	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (LYOPHILIZED) 5 MG	1	EA	VL	IJ	EA	5	MG	1	01/31/2018	99/99/9999							
54766-0590-10		J7500		01/01/2018	99/99/9999	AZATHIOPRINE, ORAL, 50 MG	IMURAN 50 MG	100	EA	BO	PO	EA	50	MG	1	01/01/2018	99/99/9999							
55150-0230-10		J1652		01/12/2018	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PF) 2.5 MG/0.5 ML	0.5	ML	SR	SC	ML	0.5	MG	10	01/12/2018	99/99/9999							
55150-0231-10		J1652		01/12/2018	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PF) 5 MG/0.4 ML	0.4	ML	SR	SC	ML	0.5	MG	25	01/12/2018	99/99/9999							
55150-0232-10		J1652		01/12/2018	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PF) 7.5 MG/0.6 ML	0.6	ML	SR	SC	ML	0.5	MG	25	01/12/2018	99/99/9999							
55150-0233-10		J1652		01/12/2018	99/99/9999	INJECTION, FONDAPARINUX SODIUM, 0.5 MG	FONDAPARINUX SODIUM (PF) 10 MG/0.8 ML	0.8	ML	SR	SC	ML	0.5	MG	25	01/12/2018	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
60505-6128-00		J9206		01/10/2018	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV;USP,PF) 20 MG/1 ML	2 ML	VL	IV	ML		20 MG		1	01/10/2018	99/99/9999						
60505-6128-01		J9206		01/10/2018	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV;USP,PF) 20 MG/1 ML	5 ML	VL	IV	ML		20 MG		1	01/10/2018	99/99/9999						
60505-6166-00		J9027		01/09/2018	99/99/9999	INJECTION, CLOFARABINE, 1 MG	CLOFARABINE (SDV,PF) 1 MG/1 ML	20 ML	VL	IV	ML		1 MG		1	01/09/2018	99/99/9999						
63323-0221-38		J3370		01/10/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF,LATEX-FREE) 500 MG	25 EA	VL	IV	EA		500 MG		1	01/10/2018	99/99/9999						
67457-0513-99		J9120		01/01/2018	02/27/2018	INJECTION, DACTINOMYCIN, 0.5 MG	DACTINOMYCIN (PF,LYOPHILIZED) 0.5 MG	12 EA	VL	IV	EA		0.5 MG		1	01/01/2018	02/27/2018						
67457-0616-10		J9201		01/03/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (1X5.26ML) 38 MG/1 ML	5.26 ML	VL	IV	ML		200 MG		0.19	01/03/2018	99/99/9999						
67457-0617-30		J9201		12/18/2017	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (1X26.3ML) 38 MG/1 ML	26.3 ML	VL	IV	ML		200 MG		0.19	12/18/2017	99/99/9999						
67457-0618-10		J9201		12/18/2017	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 38 MG/1 ML	52.6 ML	VL	IV	ML		200 MG		0.19	12/18/2017	99/99/9999						
68094-0101-10		J2760		12/19/2017	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (LYOPHILIZED) 5 MG	10 EA	VL	IJ	EA		5 MG		1	12/19/2017	99/99/9999						
68094-0101-20		J2760		12/19/2017	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (LYOPHILIZED) 5 MG	1 EA	VL	IJ	EA		5 MG		1	12/19/2017	99/99/9999						
70121-1244-07		J0594		12/28/2017	99/99/9999	INJECTION, BUSULFAN, 1 MG	BUSULFAN 6 MG/1 ML	10 ML	VL	IV	ML		1 MG		6	12/28/2017	99/99/9999						
70257-0560-01		J0475		01/25/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT 0.5 MG/1 ML	20 ML	AM	IN	ML		10 MG		0.05	01/25/2018	99/99/9999						
70257-0560-02		J0475		01/25/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT 0.5 MG/1 ML	20 ML	AM	IN	ML		10 MG		0.05	01/25/2018	99/99/9999						
70257-0561-02		J0475		01/25/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	LIORESAL INTRATHECAL REFILL KIT 2 MG/1 ML	5 ML	AM	IN	ML		10 MG		0.2	01/25/2018	99/99/9999						
70515-0260-10		J1160		01/17/2018	99/99/9999	INJECTION, DIGOXIN, UP TO 0.5 MG	LANOXIN 0.25 MG/1 ML	2 ML	AM	IJ	ML		0.5 MG		0.5	01/17/2018	99/99/9999						
70515-0262-10		J1160		01/17/2018	99/99/9999	INJECTION, DIGOXIN, UP TO 0.5 MG	LANOXIN PEDIATRIC 0.1 MG/1 ML	1 ML	AM	IJ	ML		0.5 MG		0.2	01/17/2018	99/99/9999						
70860-0205-50		J9201		10/11/2017	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (SDV, USP,PF,LATEX-FREE) 1 GM	1 EA	VL	IV	EA		200 MG		5	10/11/2017	99/99/9999						
70860-0208-05		J9000		12/15/2017	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,SDV,PF,LATEX-FREE) 2 MG/1 ML	5 ML	VL	IV	ML		10 MG		0.2	12/15/2017	99/99/9999						
70860-0208-25		J9000		12/15/2017	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,SDV,PF,LATEX-FREE) 2 MG/1 ML	25 ML	VL	IV	ML		10 MG		0.2	12/15/2017	99/99/9999						
70860-0208-51		J9000		12/15/2017	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (USP,SDV,PF,LATEX-FREE) 2 MG/1 ML	100 ML	VL	IV	ML		10 MG		0.2	12/15/2017	99/99/9999						
75987-0111-01		J9216		01/15/2018	99/99/9999	INJECTION, INTERFERON, GAMMA 1-B, 3 MILLION UNITS	ACTIMMUNE 2 MILLION IU/0.5 ML	0.5 ML	VL	SC	ML		3000000 U		1.33333	01/15/2018	99/99/9999						
75987-0111-11		J9216		01/15/2018	99/99/9999	INJECTION, INTERFERON, GAMMA 1-B, 3 MILLION UNITS	ACTIMMUNE 2 MILLION IU/0.5 ML	0.5 ML	VL	SC	ML		3000000 U		1.33333	01/15/2018	99/99/9999						
00574-0805-30		J0132		12/27/2012	99/99/9999	INJECTION, ACETYLCYSTEINE, 100 MG	ACETYLCYSTEINE (SDV, 4X30ML,PF) 200 MG/1 ML	30 ML	VL	IV	ML		100 MG		2	12/27/2012	99/99/9999						
00115-9930-78	KO	J7614	KO	01/09/2018	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.31 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.20666	01/09/2018	99/99/9999						
00115-9931-78	KO	J7614	KO	01/09/2018	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 0.63 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.42	01/09/2018	99/99/9999						
00115-9932-78	KO	J7614	KO	01/09/2018	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (PF) 1.25 MG/3 ML	3 ML	VL	IH	ML		0.5 MG		0.83333	01/09/2018	99/99/9999						
00069-0809-01		Q5103		04/01/2018	99/99/9999	INJECTION, INFLIXIMAB-DYBV, BIOSIMILAR, (INFLECTRA), 10 MG	INFLECTRA (SDV,PF) 100 MG	1 EA	VL	IV	EA		10 MG		10	04/01/2018	99/99/9999						
00006-4305-02		Q5104		04/01/2018	99/99/9999	INJECTION, INFLIXIMAB-ABDA, BIOSIMILAR, (RENFLIXIS), 10 MG	RENFLIXIS (PF,LYOPHILIZED) 100 MG	1 EA	VL	IV	EA		10 MG		10	04/01/2018	99/99/9999						
61314-0304-01		Q5101		04/01/2018	99/99/9999	INJECTION, FILGRASTIM-SNDZ, BIOSIMILAR, (ZARXIO), 1 MICROGRAM	ZARXIO (PF) 300 MCG/0.5 ML	0.5 ML	SR	IJ	ML		1 MCG		600	04/01/2018	99/99/9999						
63402-0201-00	KO	J7643	KO	02/16/2018	99/99/9999	GLYCOPYRRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	LONHALA MAGNAIR (STARTER KIT) 25 MCG/1 ML	1 ML	VL	IH	ML		1 MG		0.025	02/16/2018	99/99/9999						
63402-0301-01	KO	J7643	KO	02/16/2018	99/99/9999	GLYCOPYRRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	LONHALA MAGNAIR (REFILL KIT) 25 MCG/1 ML	1 ML	VL	IH	ML		1 MG		0.025	02/16/2018	99/99/9999						
71288-0100-05		J9045		09/15/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	5 ML	VL	IV	ML		50 MG		0.2	09/15/2017	99/99/9999						
71288-0100-15		J9045		09/15/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	15 ML	VL	IV	ML		50 MG		0.2	09/15/2017	99/99/9999						
71288-0100-45		J9045		09/15/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	45 ML	VL	IV	ML		50 MG		0.2	09/15/2017	99/99/9999						
71288-0100-51		J9045		09/15/2017	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF,LATEX-FREE) 10 MG/1 ML	60 ML	VL	IV	ML		50 MG		0.2	09/15/2017	99/99/9999						
00143-9247-01		J1190		01/29/2018	99/99/9999	INJECTION, DEXRAZOXANE HYDROCHLORIDE, PER 250 MG	DEXRAZOXANE (SDV W/DILUENT) 250 MG	1 EA	VL	IV	EA		250 MG		1	01/29/2018	99/99/9999						
00143-9248-01		J1190		01/29/2018	99/99/9999	INJECTION, DEXRAZOXANE HYDROCHLORIDE, PER 250 MG	DEXRAZOXANE (SDV W/DILUENT) 500 MG	1 EA	VL	IV	EA		250 MG		2	01/29/2018	99/99/9999						
00143-9298-10		J2916		02/14/2018	99/99/9999	INJECTION, SODIUM FERRIC GLUCONATE COMPLEX IN SUCROSE INJECTION, 12.5 MG	SODIUM FERRIC GLUCONATE COMPLEX SUCROSE NOVAPLUS (LATEX-FREE) 62.5 MG/5 ML	5 ML	VL	IV	ML		12.5 MG		1	02/14/2018	99/99/9999						
00143-9519-10		J9250		02/13/2018	99/99/9999	METHOTREXATE SODIUM, 5 MG	METHOTREXATE SODIUM (10X2ML SDV,PF) 25 MG/1 ML	2 ML	VL	IJ	ML		5 MG		5	02/13/2018	99/99/9999						
00143-9872-10		J1800		02/12/2018	99/99/9999	INJECTION, PROPRANOLOL HCL, UP TO 1 MG	PROPRANOLOL HCL (10X1ML) 1 MG/1 ML	1 ML	VL	IV	ML		1 MG		1	02/12/2018	99/99/9999						
00378-6961-12		J1595		10/04/2017	99/99/9999	INJECTION, GLATIRAMER ACETATE, 20 MG	GLATIRAMER ACETATE 40 MG/1 ML	1 ML	SR	SC	ML		20 MG		2	10/04/2017	99/99/9999						
00409-3595-01		J0698		01/22/2018	99/99/9999	INJECTION, CEFOTAXIME SODIUM, PER GM	CEFOTAXIME (USP) 1 GM	25 EA	VL	IJ	EA		1 GM		1	01/22/2018	99/99/9999						
00517-1825-10		J2800		01/29/2018	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL 100 MG/1 ML	10 ML	VL	IJ	ML		10 ML		0.1	01/29/2018	99/99/9999						
25021-0245-01		J9171		02/14/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SDV,PF,LATEX-FREE) 20 MG/1 ML	1 ML	VL	IV	ML		1 MG		20	02/14/2018	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
25021-0245-04		J9171		02/14/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SDV,PF,LATEX-FREE) 20 MG/1 ML	4	ML	VL	IV	ML	1	MG	20	02/14/2018	99/99/9999						
25021-0676-50		J2515		01/29/2018	99/99/9999	INJECTION, PENTOBARBITAL SODIUM, PER 50 MG	PENTOBARBITAL SODIUM (MDV,PF,LATEX-FREE) 50 MG/1 ML	50	ML	VL	IJ	ML	50	MG	1	01/29/2018	99/99/9999						
42195-0121-06		J8540		01/31/2018	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	TAPERDEX (6-DAY) 1.5 MG	21	EA	ST	PO	EA	0.25	MG	6	01/31/2018	99/99/9999						
42195-0149-12		J8540		01/31/2018	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	TAPERDEX (12-DAY) 1.5 MG	49	EA	ST	PO	EA	0.25	MG	6	01/31/2018	99/99/9999						
47781-0593-07		J9267		01/23/2018	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV,PF,LATEX-FREE) 6 MG/1 ML	5	ML	VL	IV	ML	1	MG	6	01/23/2018	99/99/9999						
47781-0594-07		J9267		01/23/2018	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV,PF,LATEX-FREE) 6 MG/1 ML	16.7	ML	VL	IV	ML	1	MG	6	01/23/2018	99/99/9999						
47781-0595-07		J9267		01/23/2018	99/99/9999	INJECTION, PACLITAXEL, 1 MG	PACLITAXEL (MDV,PF,LATEX-FREE) 6 MG/1 ML	50	ML	VL	IV	ML	1	MG	6	01/23/2018	99/99/9999						
50742-0401-02		J9206		02/05/2018	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV) 20 MG/1 ML	2	ML	VL	IV	ML	20	MG	1	02/05/2018	99/99/9999						
50742-0402-05		J9206		02/05/2018	99/99/9999	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (SDV) 20 MG/1 ML	5	ML	VL	IV	ML	20	MG	1	02/05/2018	99/99/9999						
50742-0445-05		J9045		01/29/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	5	ML	VL	IV	ML	50	MG	0.2	01/29/2018	99/99/9999						
50742-0446-15		J9045		01/29/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	15	ML	VL	IV	ML	50	MG	0.2	01/29/2018	99/99/9999						
50742-0447-45		J9045		01/29/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	45	ML	VL	IV	ML	50	MG	0.2	01/29/2018	99/99/9999						
50742-0448-60		J9045		01/29/2018	99/99/9999	INJECTION, CARBOPLATIN, 50 MG	CARBOPLATIN (PF) 10 MG/1 ML	60	ML	VL	IV	ML	50	MG	0.2	01/29/2018	99/99/9999						
51991-0933-17		J1630		02/05/2018	99/99/9999	INJECTION, HALOPERIDOL, UP TO 5 MG	HALOPERIDOL (10X1ML) 5 MG/1 ML	1	ML	SR	IM	ML	5	MG	1	02/05/2018	99/99/9999						
55150-0192-20		J0153		02/08/2018	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (SDV,PF,LATEX-FREE) 3 MG/1 ML	20	ML	VL	IV	ML	1	MG	3	02/08/2018	99/99/9999						
55150-0193-30		J0153		02/08/2018	99/99/9999	INJECTION, ADENOSINE, 1 MG (NOT TO BE USED TO REPORT ANY ADENOSINE PHOSPHATE COMPOUNDS)	ADENOSINE (SDV,PF,LATEX-FREE) 3 MG/1 ML	30	ML	VL	IV	ML	1	MG	3	02/08/2018	99/99/9999						
63323-0064-03		J3475		01/30/2018	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (25X2ML,PF) 500 MG/1 ML	2	ML	VL	IJ	ML	500	MG	1	01/30/2018	99/99/9999						
63323-0064-11		J3475		01/30/2018	99/99/9999	INJECTION, MAGNESIUM SULFATE, PER 500 MG	MAGNESIUM SULFATE (25X10ML,PF) 500 MG/1 ML	10	ML	VL	IJ	ML	500	MG	1	01/30/2018	99/99/9999						
63402-0201-00		J7643		02/16/2018	99/99/9999	GLYCOPYRRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	LONHALA MAGNAIR (STARTER KIT) 25 MCG/1 ML	1	ML	VL	IH	ML	1	MG	0.025	02/16/2018	99/99/9999						
63402-0301-01		J7643		02/16/2018	99/99/9999	GLYCOPYRRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	LONHALA MAGNAIR (REFILL KIT) 25 MCG/1 ML	1	ML	VL	IH	ML	1	MG	0.025	02/16/2018	99/99/9999						
66794-0156-01		J0475		02/01/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	GABLOFEN (1X20ML,SINGLE USE) 1 MG/1 ML	20	ML	SR	IN	ML	10	MG	0.1	02/01/2018	99/99/9999						
68001-0338-62		J3370		02/15/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF,LATEX-FREE) 500 MG	10	EA	VL	IV	EA	500	MG	1	02/15/2018	99/99/9999						
68001-0339-64		J3370		02/15/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF,LATEX-FREE) 1 GM	10	EA	VL	IV	EA	500	MG	2	02/15/2018	99/99/9999						
68001-0341-36		J9263		02/15/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	10	ML	VL	IV	ML	0.5	MG	10	02/15/2018	99/99/9999						
68001-0341-37		J9263		02/15/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	20	ML	VL	IV	ML	0.5	MG	10	02/15/2018	99/99/9999						
70720-0951-30		J9202		02/02/2018	99/99/9999	GOSERELIN ACETATE IMPLANT, PER 3.6 MG	ZOLADEX (SAFESYSTEM SRN) 10.8 MG	1	EA	SR	SC	EA	3.6	MG	3	02/02/2018	99/99/9999						
67457-0864-04		J1626		03/21/2018	99/99/9999	INJECTION, GRANISETRON HYDROCHLORIDE, 100 MCG	GRANISETRON HYDROCHLORIDE (1X4ML,MDV,LATEX-FREE) 1 MG/1 ML	4	ML	VL	IV	ML	100	MCG	10	03/21/2018	99/99/9999						
69097-0927-35		J2469		03/23/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL (S.D.V.) 0.05 MG/1 ML	5	ML	VL	IV	ML	25	MCG	2	03/23/2018	99/99/9999						
70720-0950-36		J9202		04/06/2018	99/99/9999	GOSERELIN ACETATE IMPLANT, PER 3.6 MG	ZOLADEX (SAFESYSTEM SRN) 3.6 MG	1	EA	SR	SC	EA	3.6	MG	1	04/06/2018	99/99/9999						
70860-0106-10		J0637		03/01/2018	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LATEX-FREE) 50 MG	1	EA	VL	IV	EA	5	MG	10	03/01/2018	99/99/9999						
70860-0107-10		J0637		03/01/2018	99/99/9999	INJECTION, CASPOFUNGIN ACETATE, 5 MG	CASPOFUNGIN ACETATE (PF,LATEX-FREE) 70 MG	1	EA	VL	IV	EA	5	MG	14	03/01/2018	99/99/9999						
68382-0755-96		None		06/01/2018	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 180 MG	5	EA	BO	PO	EA	20	MG	9	06/01/2018	99/99/9999						
00078-0930-61		J0883		03/14/2018	99/99/9999	INJECTION, ARGATROBAN, 1 MG (FOR NON-ESRD USE)	ARGATROBAN (SINGLE USE VIAL,PF) 100 MG/1 ML	2.5	ML	VL	IV	ML	1	MG	100	03/14/2018	99/99/9999						
00143-9510-01		J9181		02/26/2018	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (USP, MDV) 20 MG/1 ML	5	ML	VL	IV	ML	10	MG	2	02/26/2018	99/99/9999						
00143-9511-01		J9181		02/26/2018	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (USP, MDV) 20 MG/1 ML	25	ML	VL	IV	ML	10	MG	2	02/26/2018	99/99/9999						
00143-9512-01		J9181		02/26/2018	99/99/9999	INJECTION, ETOPOSIDE, 10 MG	ETOPOSIDE (USP, MDV) 20 MG/1 ML	50	ML	VL	IV	ML	10	MG	2	02/26/2018	99/99/9999						
00409-7241-61		J0171		01/01/2018	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE 1 MG/1 ML	1	ML	AM	IJ	ML	0.1	MG	10	01/01/2018	99/99/9999						
00781-3250-89		J1595		02/27/2018	99/99/9999	INJECTION, GLATIRAMER ACETATE, 20 MG	GLATIRAMER ACETATE 20 MG	1	ML	SR	SC	ML	20	MG	2	02/27/2018	99/99/9999						
13533-0631-11		J2790		04/01/2018	99/99/9999	INJECTION, RHO D IMMUNE GLOBULIN, HUMAN, FULL DOSE, 300 MICROGRAMS (1500 IU)	HYPERRHO S/D (PF,LATEX-FREE) 300 MCG	10	EA	SR	IM	EA	300	MCG	1	04/01/2018	99/99/9999						
43066-0001-01		J9171		02/23/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (1X2ML,MDV) 10 MG/1 ML	2	ML	VL	IV	ML	1	MG	10	02/23/2018	99/99/9999						
43066-0006-01		J9171		02/23/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (1X8ML,MDV) 10 MG/1 ML	8	ML	VL	IV	ML	1	MG	10	02/23/2018	99/99/9999						
43066-0010-01		J9171		02/23/2018	99/99/9999	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (1X2ML,MDV) 10 MG/1 ML	16	ML	VL	IV	ML	1	MG	10	02/23/2018	99/99/9999						
43066-0014-01		J9263		02/23/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	10	ML	VL	IV	ML	0.5	MG	10	02/23/2018	99/99/9999						
43066-0018-01		J9263		02/23/2018	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	20	ML	VL	IV	ML	0.5	MG	10	02/23/2018	99/99/9999						
47781-0597-91		J3370		04/01/2017	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PF,LATEX-FREE) 1 GM	10	EA	VL	IV	EA	500	MG	2	04/01/2017	99/99/9999						
51224-0012-10		J2760		03/15/2018	99/99/9999	INJECTION, PHENTOLAMINE MESYLATE, UP TO 5 MG	PHENTOLAMINE MESYLATE (LYOPHILIZED) 5 MG	10	EA	VL	IJ	EA	5	MG	1	03/15/2018	99/99/9999						
60505-6114-00		J9201		02/23/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 38 MG/1 ML	26.3	ML	VL	IV	ML	200	MG	0.19	02/23/2018	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
60505-6115-02		J9201		02/23/2018	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE 38 MG/1 ML MELPHALAN HYDROCHLORIDE (W/10ML DILUENT) 50 MG	52.6	ML	VL	IV	ML	200	MG	0.19	02/23/2018	99/99/9999							
63323-0760-20		J9245		02/21/2018	99/99/9999	INJECTION, MELPHALAN HYDROCHLORIDE, 50 MG	GAMMAPLEX 10% (INNER PACK NDC,PF) 100 MG/1 ML	1	EA	VL	IV	EA	50	MG	1	02/21/2018	99/99/9999							
64208-8235-01		J1557		04/01/2017	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), 500 MG	GAMMAPLEX 10% (INNER PACK NDC,PF) 100 MG/1 ML	50	ML	VL	IV	ML	500	MG	0.2	04/01/2017	99/99/9999							
64208-8235-02		J1557		04/01/2017	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), 500 MG	GAMMAPLEX 10% (INNER PACK NDC,PF) 100 MG/1 ML	100	ML	VL	IV	ML	500	MG	0.2	04/01/2017	99/99/9999							
64208-8235-03		J1557		04/01/2017	99/99/9999	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NON-LYOPHILIZED (E.G., LIQUID), 500 MG	GAMMAPLEX 10% (INNER PACK NDC,PF) 100 MG/1 ML	200	ML	VL	IV	ML	500	MG	0.2	04/01/2017	99/99/9999							
66794-0155-02		J0475		04/01/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	GABLOFEN (1X20ML,SINGLE USE) 0.5 MG/1 ML	20	ML	VL	IN	ML	10	MG	0.05	04/01/2018	99/99/9999							
66794-0156-02		J0475		04/01/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	GABLOFEN (1X20ML,SINGLE USE) 1 MG/1 ML	20	ML	VL	IN	ML	10	MG	0.1	04/01/2018	99/99/9999							
66794-0157-02		J0475		04/01/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	GABLOFEN (1X20ML,SINGLE USE) 2 MG/1 ML	20	ML	VL	IN	ML	10	MG	0.2	04/01/2018	99/99/9999							
67457-0374-99		J1644		03/16/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X1ML) 5000 U/1 ML	1	ML	VL	IJ	ML	1000	U	5	03/16/2018	99/99/9999							
67457-0384-99		J1644		03/16/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X30ML) 1000 U/1 ML	30	ML	VL	IJ	ML	1000	U	1	03/16/2018	99/99/9999							
67457-0385-99		J1644		03/16/2018	99/99/9999	INJECTION, HEPARIN SODIUM, PER 1000 UNITS	HEPARIN SODIUM (MDV,25X10ML) 1000 U/1 ML	10	ML	VL	IJ	ML	1000	U	1	03/16/2018	99/99/9999							
67457-0513-05		J9120		01/01/2018	99/99/9999	INJECTION, DACTINOMYCIN, 0.5 MG	DACTINOMYCIN (PF,LYOPHILIZED) 0.5 MG	1	EA	VL	IV	EA	0.5	MG	1	01/01/2018	99/99/9999							
67457-0518-05		J9280		02/28/2018	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MITOMYCIN (PF,LYOPHILIZED) 5 MG	1	EA	VL	IV	EA	5	MG	1	02/28/2018	99/99/9999							
67457-0519-20		J9280		02/28/2018	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MITOMYCIN (SDV,PF,LYOPHILIZED) 20 MG	1	EA	VL	IV	EA	5	MG	4	02/28/2018	99/99/9999							
68001-0345-26		Q2050		04/02/2018	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, LIPOSOMAL, NOT OTHERWISE SPECIFIED, 10 MG	DOXORUBICIN HCL LIPOSOME 2 MG/1 ML	25	ML	VL	IV	ML	10	MG	0.2	04/02/2018	99/99/9999							
68001-0345-36		Q2050		04/02/2018	99/99/9999	INJECTION, DOXORUBICIN HYDROCHLORIDE, LIPOSOMAL, NOT OTHERWISE SPECIFIED, 10 MG	DOXORUBICIN HCL LIPOSOME 2 MG/1 ML	10	ML	VL	IV	ML	10	MG	0.2	04/02/2018	99/99/9999							
69448-0001-05		J9280		09/25/2017	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MUTAMYCIN 5 MG	1	EA	VL	IV	EA	5	MG	1	09/25/2017	99/99/9999							
69448-0002-11		J9280		09/25/2017	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MUTAMYCIN 20 MG	1	EA	VL	IV	EA	5	MG	4	09/25/2017	99/99/9999							
69448-0003-38		J9280		09/25/2017	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MUTAMYCIN 40 MG	1	EA	VL	IV	EA	5	MG	8	09/25/2017	99/99/9999							
70121-1076-05		J1940		04/19/2017	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/1 ML	10	ML	VL	IJ	ML	20	MG	0.5	04/19/2017	99/99/9999							
70121-1163-05		J1940		04/19/2017	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/1 ML	2	ML	VL	IJ	ML	20	MG	0.5	04/19/2017	99/99/9999							
70121-1164-05		J1940		04/19/2017	99/99/9999	INJECTION, FUROSEMIDE, UP TO 20 MG	FUROSEMIDE (SDV) 10 MG/1 ML	4	ML	VL	IJ	ML	20	MG	0.5	04/19/2017	99/99/9999							
70860-0209-10		J9209		01/10/2018	99/99/9999	INJECTION, MESNA, 200 MG	MESNA 100 MG/1 ML	10	ML	VL	IV	ML	200	MG	0.5	01/10/2018	99/99/9999							
70860-0700-02		J1885		03/01/2018	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (PF,LATEX-FREE) 15 MG/1 ML	1	ML	VL	IJ	ML	15	MG	1	03/01/2018	99/99/9999							
70860-0701-03		J1885		03/01/2018	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (PF,LATEX-FREE) 30 MG/1 ML	1	ML	VL	IM	ML	15	MG	2	03/01/2018	99/99/9999							
70860-0701-04		J1885		03/01/2018	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (PF,LATEX-FREE) 30 MG/1 ML	2	ML	VL	IM	ML	15	MG	2	03/01/2018	99/99/9999							
68382-0756-96		None		06/01/2018	99/99/9999	TEMOZOLOMIDE, 250 MG, ORAL	TEMOZOLOMIDE (HARD GELATIN) 250 MG	5	EA	BO	PO	EA	250	MG	1	06/01/2018	99/99/9999							
69097-0537-37		J1071		06/19/2018	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (USP,MDV) 200 MG/1 ML	10	ML	VL	IM	ML	1	MG	200	06/19/2018	99/99/9999							
70069-0064-01		J2795		07/02/2018	99/99/9999	INJECTION, ROPIVACAINE HYDROCHLORIDE, 1 MG	ROPIVACAINE HCL (PF,LATEX-FREE) 5 MG/1 ML	30	ML	VL	IJ	ML	1	MG	5	07/02/2018	99/99/9999							
70801-0003-01		Q9993		07/01/2018	12/31/2018	INJECTION, TRIAMCINOLONE ACETONIDE, PRESERVATIVE-FREE, EXTENDED-RELEASE, MICROSPHERE FORMULATION, 1 MG	ZILRETTA (W/DILUENT) 32 MG	1	EA	VL	IJ	EA	1	MG	32	07/01/2018	12/31/2018							
70842-0140-03		J2407		06/25/2018	99/99/9999	INJECTION, ORITAVANCIN, 10 MG	ORBACTIV (PF,LYOPHILIZED) 400 MG	3	EA	VL	IV	EA	10	MG	40	06/25/2018	99/99/9999							
70860-0118-99		J0290		06/25/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (PHARMACY BULK,USP,PF) 10 GM	1	EA	VL	IJ	EA	500	MG	20	06/25/2018	99/99/9999							
71930-0017-30		Q0162		07/18/2018	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HCL (FILM-COATED) 4 MG	30	EA	BO	PO	EA	1	MG	4	07/18/2018	99/99/9999							
71930-0018-30		Q0162		07/18/2018	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON (FILM-COATED) 8 MG	30	EA	BO	PO	EA	1	MG	8	07/18/2018	99/99/9999							
00517-1767-01		J1729		06/22/2018	99/99/9999	INJECTION, HYDROXYPROGESTERONE CAPROATE, NOT OTHERWISE SPECIFIED, 10 MG	HYDROXYPROGESTERONE CAPROATE (PF) 250 MG/1 ML	1	ML	VL	IM	ML	10	MG	25	06/22/2018	99/99/9999							
12496-0100-01		Q9991		07/01/2018	99/99/9999	INJECTION, BUPRENORPHINE EXTENDED-RELEASE (SUBLOCADE), LESS THAN OR EQUAL TO 100 MG	SUBLOCADE 100 MG/0.5 ML	0.5	ML	SR	SC	ML	100	MG	2	07/01/2018	99/99/9999							
12496-0300-01		Q9992		07/01/2018	99/99/9999	INJECTION, BUPRENORPHINE EXTENDED-RELEASE (SUBLOCADE), GREATER THAN 100 MG	SUBLOCADE 100 MG/0.5 ML	1.5	ML	SR	SC	ML	100	MG	2	07/01/2018	99/99/9999							
16714-0834-01		J2469		08/08/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL 0.05 MG/1 ML	5	ML	VL	IV	ML	25	MCG	2	08/08/2018	99/99/9999							
43598-2050-11		J9340		05/08/2018	99/99/9999	INJECTION, THIOTEPA, 15 MG	THIOTEPA (SDV,LYOPHILIZED) 15 MG	1	EA	VL	IJ	EA	15	MG	1	05/08/2018	99/99/9999							
52652-2001-06		None		07/31/2018	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	XATMEP 2.5 MG/1 ML	60	ML	BO	PO	ML	2.5	MG	1	07/31/2018	99/99/9999							
55111-0694-07		J2469		03/23/2018	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL 0.05 MG/1 ML	5	ML	VL	IV	ML	25	MCG	2	03/23/2018	99/99/9999							
55150-0282-20		J1335		06/27/2018	99/99/9999	INJECTION, ERTAPENEM SODIUM, 500 MG	ERTAPENEM (LATEX-FREE,LYOPHILIZED) 1 GM	10	EA	VL	IJ	EA	500	MG	2	06/27/2018	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3		
63323-0852-25	J1170			06/19/2018	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (PF,LATEX-FREE) 1 MG/1 ML	1 ML	VL	IJ	ML		4 MG	0.25	06/19/2018	99/99/9999									
63323-0853-25	J1170			06/19/2018	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (PF,LATEX-FREE) 2 MG/1 ML	1 ML	VL	IJ	ML		4 MG	0.5	06/19/2018	99/99/9999									
63323-0854-10	J1170			06/19/2018	99/99/9999	INJECTION, HYDROMORPHONE, UP TO 4 MG	HYDROMORPHONE HCL (PF,LATEX-FREE) 4 MG/1 ML	1 ML	VL	IJ	ML		4 MG	1	06/19/2018	99/99/9999									
65757-0500-03	J1942			07/02/2018	99/99/9999	INJECTION, ARIPIRAZOLE LAUROXIL, 1 MG	ARISTADA INITIO (LATEX-FREE) 675 MG/2.4 ML	2.4 ML	SR	IM	ML		1 MG	281.25	07/02/2018	99/99/9999									
66993-0038-83	J1729			07/02/2018	99/99/9999	INJECTION, HYDROXYPROGESTERONE CAPROATE, NOT OTHERWISE SPECIFIED, 10 MG	HYDROXYPROGESTERONE CAPROATE (PF) 250 MG/1 ML	1 ML	VL	IM	ML		10 MG	25	07/02/2018	99/99/9999									
69097-0536-37	J1071			06/19/2018	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (USP,MDV) 100 MG/1 ML	10 ML	VL	IM	ML		1 MG	100	06/19/2018	99/99/9999									
69097-0537-31	J1071			06/19/2018	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (USP,SDV) 200 MG/1 ML	1 ML	VL	IM	ML		1 MG	200	06/19/2018	99/99/9999									
00009-3475-01	J1040			01/07/1992	99/99/9999	INJECTION, METHYLPREDNISOLONE ACETATE, 80 MG	DEPO-MEDROL (S.D.V.) 80 MG/1 ML	1 ML	VL	IJ	ML		80 MG	1	01/07/1992	99/99/9999									
00310-1730-30	J3490			11/14/2017	99/99/9999	UNCLASSIFIED DRUGS	FASENRA (PF) 30 MG/1 ML	1 ML	SR	SC	ML		1 MG	1	11/14/2017	99/99/9999									
00409-3795-19	J1885			01/06/2006	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE (INNER PACK,LATEX-FREE) 30 MG/1 ML	1 ML	VL	IJ	EA		15 MG	2	01/06/2006	99/99/9999									
70860-0113-15	J0290			08/01/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (USP,PF,LATEX-FREE) 500 MG	10 EA	VL	IJ	EA		500 MG	1	08/01/2018	99/99/9999									
70860-0114-15	J0290			08/01/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (PF,LATEX-FREE) 1 GM	10 EA	VL	IJ	EA		500 MG	2	08/01/2018	99/99/9999									
70860-0115-26	J0290			07/31/2018	99/99/9999	INJECTION, AMPICILLIN SODIUM, 500 MG	AMPICILLIN (PF,LATEX-FREE) 2 GM	10 EA	VL	IJ	EA		500 MG	4	07/31/2018	99/99/9999									
71274-0350-02	J0596			04/01/2018	99/99/9999	INJECTION, C1 ESTERASE INHIBITOR (RECOMBINANT), RUCONEST, 10 UNITS	RUCONEST (PF) 2100 IU	1 EA	BX	IV	EA		10 U	210	04/01/2018	99/99/9999									
76075-0103-01	J9047			08/21/2018	99/99/9999	INJECTION, CARFILZOMIB, 1 MG	KYPROLIS (LYOPHILIZED) 10 MG	1 EA	VL	IV	EA		1 MG	10	08/21/2018	99/99/9999									
63275-5100-04	J3010			06/01/2015	99/99/9999	INJECTION, FENTANYL CITRATE, 0.1 MG	FENTANYL CITRATE (USP)	25 GM	BO	NA	GM		0.1 MG	10000	06/01/2015	99/99/9999									
00069-1308-10	J0885			05/22/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 10000 U/1 ML	1 ML	VL	IJ	ML		1000 U	10	05/22/2018	12/31/2018									
00069-1309-04	J0885			05/22/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 40000 U/1 ML	1 ML	VL	IJ	ML		1000 U	40	05/22/2018	12/31/2018									
59353-0002-01	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 2000 U/1 ML	1 ML	VL	IJ	ML		1000 U	2	05/25/2018	12/31/2018									
59353-0002-10	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 2000 U/1 ML	1 ML	VL	IJ	ML		1000 U	2	05/25/2018	12/31/2018									
59353-0003-01	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 3000 U/1 ML	1 ML	VL	IJ	ML		1000 U	3	05/25/2018	12/31/2018									
59353-0003-10	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 3000 U/1 ML	1 ML	VL	IJ	ML		1000 U	3	05/25/2018	12/31/2018									
59353-0003-10	Q5106			01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 3000 U/1 ML	1 ML	VL	IJ	ML		1000 U	3	01/01/2019	99/99/9999									
59353-0004-01	Q5106			01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 4000 U/1 ML	1 ML	VL	IJ	ML		1000 U	4	01/01/2019	99/99/9999									
59353-0004-10	Q5106			01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 4000 U/1 ML	1 ML	VL	IJ	ML		1000 U	4	01/01/2019	99/99/9999									
59353-0010-01	Q5106			01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 10000 U/1 ML	1 ML	VL	IJ	ML		1000 U	10	01/01/2019	99/99/9999									
59353-0010-10	Q5106			01/01/2019	99/99/9999	INJECTION, EPOETIN ALFA, BIOSIMILAR, (RETACRIT) (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 10000 U/1 ML	1 ML	VL	IJ	ML		1000 U	10	01/01/2019	99/99/9999									
62064-0122-02	J1746			01/01/2019	99/99/9999	INJECTION, IBALIZUMAB-IUYK, 10 MG	TROGARZO (PF) 150 MG/1 ML	1.33 ML	VL	IV	ML		10 MG	15	01/01/2019	99/99/9999									
63459-0918-59	J1447			09/04/2018	99/99/9999	INJECTION, TBO-FILGRASTIM, 1 MICROGRAM	GRANIX (PF) 300 MCG/1 ML	1 ML	VL	SC	ML		1 MCG	300	09/04/2018	99/99/9999									
63459-0920-59	J1447			09/04/2018	99/99/9999	INJECTION, TBO-FILGRASTIM, 1 MICROGRAM	GRANIX (PF) 480 MCG/1.6 ML	1.6 ML	VL	SC	ML		1 MCG	300	09/04/2018	99/99/9999									
66621-0790-02	J0841			01/01/2019	99/99/9999	INJECTION, CROTALIDAE IMMUNE F(AB)2 (EQUINE), 120 MG	ANAVIP (LYOPHILIZED) (10ML VL) 24 MG/1 ML	1 EA	VL	IV	EA		120 MG	2	01/01/2019	99/99/9999									
67457-0879-05	J3030			11/06/2018	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	SUMATRIPTAN SUCCINATE (PREFILLED,PF,LATEX-FREE) 6 MG/0.5 ML	0.5 ML	SR	SC	ML		6 MG	2	11/06/2018	99/99/9999									
67457-0880-05	J3030			11/06/2018	99/99/9999	INJECTION, SUMATRIPTAN SUCCINATE, 6 MG (CODE MAY BE USED FOR MEDICARE WHEN DRUG ADMINISTERED UNDER THE DIRECT SUPERVISION OF A PHYSICIAN, NOT FOR USE WHEN DRUG IS SELF ADMINISTERED)	SUMATRIPTAN SUCCINATE (5X0.5ML,SDV,PF) 6 MG/0.5 ML	0.5 ML	VL	SC	ML		6 MG	2	11/06/2018	99/99/9999									
68001-0370-27	J9070			11/05/2018	99/99/9999	CYCLOPHOSPHAMIDE, 100 MG	CYCLOPHOSPHAMIDE (SDV,USP,PF) 500 MG	1 EA	VL	IV	EA		100 MG	5	11/05/2018	99/99/9999									
68001-0371-32	J9070			11/05/2018	99/99/9999	CYCLOPHOSPHAMIDE, 100 MG	CYCLOPHOSPHAMIDE (SDV,USP,PF) 1 GM	1 EA	VL	IV	EA		100 MG	10	11/05/2018	99/99/9999									
68001-0372-32	J9070			11/05/2018	99/99/9999	CYCLOPHOSPHAMIDE, 100 MG	CYCLOPHOSPHAMIDE (SDV,USP,PF) 2 GM	1 EA	VL	IV	EA		100 MG	20	11/05/2018	99/99/9999									
69639-0102-01	J1454			01/01/2019	99/99/9999	INJECTION, FOSNETUPITANT 235 MG AND PALONOSETRON 0.25 MG	AKYNZEO (SDV,PF,LYOPHILIZED) 235 MG-0.25 MG	1 EA	VL	IV	EA		235.25 MG	1	01/01/2019	99/99/9999									
69656-0102-10	J2797			01/01/2019	99/99/9999	INJECTION, ROLAPITANT, 0.5 MG	VARUBI (SDV) 1.8 MG/1 ML	92.5 ML	VL	IV	ML		0.5 MG	3.6	01/01/2019	99/99/9999									
69794-0001-01	J3397			01/01/2019	99/99/9999	INJECTION, VESTRONIDASE ALFA-VJBK, 1 MG	MEPSEVII (PF) 2 MG/1 ML	5 ML	VL	IV	ML		1 MG	2	01/01/2019	99/99/9999									
69794-0102-01	J3490			04/17/2018	12/31/2018	UNCLASSIFIED DRUGS	CRYSVITA (PF) 10 MG/1 ML	1 ML	VL	SC	ML		1 MG	1	04/17/2018	12/31/2018									
69794-0203-01	J3490			04/17/2018	12/31/2018	UNCLASSIFIED DRUGS	CRYSVITA (PF) 20 MG/1 ML	1 ML	VL	SC	ML		1 MG	1	04/17/2018	12/31/2018									
69794-0304-01	J3490			04/17/2018	12/31/2018	UNCLASSIFIED DRUGS	CRYSVITA (PF) 30 MG/1 ML	1 ML	VL	SC	ML		1 MG	1	04/17/2018	12/31/2018									
69918-0720-10	J9017			11/13/2018	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (10X10 SDV,PF) 1 MG/1 ML	10 ML	VL	IV	ML		1 MG	1	11/13/2018	99/99/9999									
70594-0046-02	J3370			11/06/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (USP,LATEX-FREE) 1 GM	10 EA	VL	IV	EA		500 MG	2	11/06/2018	99/99/9999									
70594-0048-01	J3370			12/14/2018	99/99/9999	INJECTION, VANCOMYCIN HCL, 500 MG	VANCOMYCIN HCL (PHARMACY BULK PACKAGE) 10 GM	1 EA	VL	IV	EA		500 MG	20	12/14/2018	99/99/9999									
70860-0778-02	J0780			11/02/2018	99/99/9999	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE (LATEX-FREE) 5 MG/1 ML	2 ML	VL	IJ	ML		10 MG	0.5	11/02/2018	99/99/9999									

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Units of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3	
70860-0778-10	J0780			11/01/2018	99/99/9999	INJECTION, PROCHLORPERAZINE, UP TO 10 MG	PROCHLORPERAZINE EDISYLATE (MDV,LATEX-FREE) 5 MG/1 ML	10	ML	VL	IJ	ML	10	MG	0.5	11/01/2018	99/99/9999							
72485-0201-01	J9025			10/25/2018	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (SDV) 100 MG	1	EA	VL	IJ	EA	1	MG	100	10/25/2018	99/99/9999							
00143-9739-10	J7512			06/11/2013	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	1000	EA	BO	PO	EA	1	MG	10	06/11/2013	99/99/9999							
51079-0721-20	J7517			06/01/2009	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (10 X 10,HARD GELATIN) 250 MG	100	EA	ST	PO	EA	250	MG	1	06/01/2009	99/99/9999							
60505-0687-04	J2543			09/21/2009	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SDV) 3 GM-0.375 GM	10	EA	VL	IV	EA	1.125	GM	3	09/21/2009	99/99/9999							
60505-0688-04	J2543			09/21/2009	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SDV) 4 GM-0.5 GM	10	EA	VL	IV	EA	1.125	GM	4	09/21/2009	99/99/9999							
00003-0371-13	J0485			06/23/2011	99/99/9999	INJECTION, BELATACEPT, 1 MG	NULOJIX 250 MG	1	EA	VL	IV	EA	1	MG	250	06/23/2011	99/99/9999							
50090-3418-02	None			06/08/2018	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	100	EA	BO	PO	EA	2.5	MG	1	06/08/2018	99/99/9999							
50090-3418-09	None			06/08/2018	99/99/9999	METHOTREXATE, 2.5 MG, ORAL	METHOTREXATE SODIUM 2.5 MG	36	EA	BO	PO	EA	2.5	MG	1	06/08/2018	99/99/9999							
51407-0095-60	None			08/08/2018	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE 150 MG	60	EA	BO	PO	EA	150	MG	1	08/08/2018	99/99/9999							
51407-0096-12	None			08/08/2018	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE 500 MG	120	EA	BO	PO	EA	500	MG	1	08/08/2018	99/99/9999							
69097-0948-08	None			08/01/2018	99/99/9999	CAPECITABINE, 500 MG, ORAL	CAPECITABINE (FILM COATED) 500 MG	120	EA	BO	PO	EA	500	MG	1	08/01/2018	99/99/9999							
69097-0949-03	None			08/01/2018	99/99/9999	CAPECITABINE, 150 MG, ORAL	CAPECITABINE (FILM COATED) 150 MG	60	EA	BO	PO	EA	150	MG	1	08/01/2018	99/99/9999							
59353-0004-01	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 4000 U/1 ML	1	ML	VL	IJ	ML	1000	U	4	05/25/2018	12/31/2018							
59353-0004-10	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 4000 U/1 ML	1	ML	VL	IJ	ML	1000	U	4	05/25/2018	12/31/2018							
59353-0010-01	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 10000 U/1 ML	1	ML	VL	IJ	ML	1000	U	10	05/25/2018	12/31/2018							
59353-0010-10	J0885			05/25/2018	12/31/2018	INJECTION, EPOETIN ALFA, (FOR NON-ESRD USE), 1000 UNITS	RETACRIT (PF) 10000 U/1 ML	1	ML	VL	IJ	ML	1000	U	10	05/25/2018	12/31/2018							
60505-0773-00	J2543			09/21/2009	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (PHARMACY BULK PACKAGE) 36 GM-4.5 GM	1	EA	BO	IV	EA	1.125	GM	36	09/21/2009	99/99/9999							
60505-6076-04	J0456			09/02/2010	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN (MONOHYDRATE;SINGLE-DOSE) 500 MG	10	EA	VL	IV	EA	500	MG	1	09/02/2010	99/99/9999							
60505-6093-05	J0690			09/10/2012	05/31/2018	INJECTION, CEFZOLIN SODIUM, 500 MG	CEFZOLIN NOVAPLUS (USP) 1 GM	25	EA	VL	IJ	EA	500	MG	2	09/10/2012	05/31/2018							
60505-6102-04	J0696			11/22/2013	99/99/9999	INJECTION, CEFTRIAXONE SODIUM, PER 250 MG	CEFTRIAXONE NOVAPLUS (CRYSTALLINE) 2 GM	10	EA	VL	IJ	EA	250	MG	8	11/22/2013	99/99/9999							
00093-5986-27	J0171			11/27/2018	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	EPINEPHRINE (USP) 0.3 MG/0.3 ML	2	EA	PG	IJ	EA	0.1	MG	3	11/27/2018	99/99/9999							
72627-2100-01	J1071			12/10/2018	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (MDV) 200 MG/1 ML	30	ML	VL	IM	ML	1	MG	200	12/10/2018	99/99/9999							
67457-0631-10	J1327			12/13/2018	99/99/9999	INJECTION, EPTIFIBATIDE, 5 MG	EPTIFIBATIDE 0.75 MG/1 ML	100	ML	VL	IV	ML	5	MG	0.15	12/13/2018	99/99/9999							
70710-1478-01	J1451			12/07/2018	99/99/9999	INJECTION, FOMEPIZOLE, 15 MG	FOMEPIZOLE (1X1.5ML,PF) 1 GM/1 ML	1.5	ML	VL	IV	ML	15	MG	66.666666	12/07/2018	99/99/9999							
51224-0013-10	J1953			12/10/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (10X5ML,SINGLE-USE) 100 MG/1 ML	5	ML	VL	IV	ML	10	MG	10	12/10/2018	99/99/9999							
51224-0013-25	J1953			12/10/2018	99/99/9999	INJECTION, LEVETIRACETAM, 10 MG	LEVETIRACETAM (SINGLE-USE) 100 MG/1 ML	5	ML	VL	IV	ML	10	MG	10	12/10/2018	99/99/9999							
43063-0911-21	J7512			11/30/2018	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 20 MG	21	EA	BO	PO	EA	1	MG	20	11/30/2018	99/99/9999							
50436-1730-05	J7512			11/01/2018	99/99/9999	PREDNISONE, IMMEDIATE RELEASE OR DELAYED RELEASE, ORAL, 1 MG	PREDNISONE 10 MG	21	EA	BO	PO	EA	1	MG	10	11/01/2018	99/99/9999							
00093-4145-56	J7614			12/14/2018	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (6X5,PF) 0.31 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.20666	12/14/2018	99/99/9999							
00093-4148-56	J7614			12/14/2018	99/99/9999	LEVABUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVABUTEROL (6X5,PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	12/14/2018	99/99/9999							
60429-0846-60	J8499			11/12/2018	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	VALGANCICLOVIR HYDROCHLORIDE (FILM-COATED) 450 MG	60	EA	BO	PO	EA	1	MG	1	11/12/2018	99/99/9999							
69097-0277-03	J8499			12/12/2018	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	EXEMESTANE (FILM COATED) 25 MG	30	EA	BO	PO	EA	1	MG	1	12/12/2018	99/99/9999							
68382-0383-06	J8999			11/08/2018	99/99/9999	PRESCRIPTION DRUG, ORAL, CHEMOTHERAPEUTIC, NOS	ARSENIC TRIOXIDE (SDV,PF,LATEX-FREE) 1 MG/1 ML	10	ML	VL	IV	ML	1	MG	1	11/15/2018	99/99/9999							
50742-0438-10	J9017			11/15/2018	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (SDV,PF,LATEX-FREE) 1 MG/1 ML	10	ML	VL	IV	ML	1	MG	1	11/15/2018	99/99/9999							
68382-0997-10	J9017			12/11/2018	99/99/9999	INJECTION, ARSENIC TRIOXIDE, 1 MG	ARSENIC TRIOXIDE (SDV,PF,LATEX-FREE) 1 MG/1 ML	10	ML	VL	IV	ML	1	MG	1	12/11/2018	99/99/9999							
70121-1482-02	J9050			11/15/2018	99/99/9999	INJECTION, CARMUSTINE, 100 MG	CARMUSTINE (SDV,LYPHILIZED) 100 MG	1	EA	VL	IV	EA	100	MG	1	11/15/2018	99/99/9999							
55700-0705-06	Q0144			11/30/2018	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN 250 MG	6	EA	BO	PO	EA	1000	MG	0.25	11/30/2018	99/99/9999							
50436-1880-01	Q0162			12/04/2018	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON HYDROCHLORIDE 8 MG	30	EA	BO	PO	EA	1	MG	8	12/04/2018	99/99/9999							
43063-0742-15	Q0164			11/06/2018	99/99/9999	PROCHLORPERAZINE MALEATE, 5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROCHLORPERAZINE MALEATE 10 MG	15	EA	BO	PO	EA	5	MG	2	11/06/2018	99/99/9999							
43063-0874-20	Q0169			12/05/2018	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 25 MG	20	EA	BO	PO	EA	12.5	MG	2	12/05/2018	99/99/9999							
43063-0876-04	Q0169			12/05/2018	99/99/9999	PROMETHAZINE HYDROCHLORIDE, 12.5 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	PROMETHAZINE HCL 50 MG	4	EA	BO	PO	EA	12.5	MG	4	12/05/2018	99/99/9999							

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items in NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
00115-1804-02		Q0177		12/03/2018	99/99/9999	HYDROXYZINE PAMOATE, 25 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	HYDROXYZINE PAMOATE 50 MG	500	EA	BO	PO	EA	25	MG	2	12/03/2018	99/99/9999						
67877-0225-05		J7517		03/20/2012	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (FILM-COATED) 500 mg	500	EA	BO	PO	EA	250	MG	2	03/20/2012	99/99/9999						
59762-4538-02		J1050		09/17/2012	99/99/9999	INJECTION, MEDROXYPROGESTERONE ACETATE, 1 MG	MEDROXYPROGESTERONE ACETATE (1X1ML) strength 150 mg/1 ml	1	ML	SY	IM	ML	1	MG	150	09/17/2012	99/99/9999						
57902-0249-01		J9019		11/01/2017	99/99/9999	INJECTION, ASPARAGINASE (ERWINAZE), 1000 IU	ERWINAZE (SDV, LYOPHILIZED POWDER) 10000 iu	1	EA	VL	IJ	EA	1000	IU	10	11/01/2017	99/99/9999						
57902-0249-05		J9019		11/01/2017	99/99/9999	INJECTION, ASPARAGINASE (ERWINAZE), 1000 IU	ERWINAZE (LYOPHILIZED POWDER) 10000 iu	1	EA	VL	IJ	EA	1000	IU	10	11/01/2017	99/99/9999						
00409-3815-12		J2274		01/01/2015	99/99/9999	INJECTION, MORPHINE SULFATE, PRESERVATIVE-FREE FOR EPIDURAL OR INTRATHECAL USE, 10MG	MORPHINE SULFATE (5X10ML, LATEX-FREE) 1 MG/ML	10	ML	VL	IJ	ML	10	MG	0.1	01/01/2015	99/99/9999						
00093-4145-56	KO	J7614	KO	12/14/2018	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X5, PF) 0.31 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.20666	12/14/2018	99/99/9999						
00093-4148-56	KO	J7614	KO	12/14/2018	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X5, PF) 1.25 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.83333	12/14/2018	99/99/9999						
00023-3921-02		J0585		01/01/2010	99/99/9999	INJECTION, ONABOTULINUMTOXINA, 1 UNIT	BOTOX (SINGLE USE) 200 u	1	EA	VL	IJ	EA	1	U	200	01/01/2010	99/99/9999						
49502-0806-93		J7699		12/14/2018	99/99/9999	NOC DRUGS, INHALATION SOLUTION ADMINISTERED THROUGH DME	YUPELRI 175 mcg/3 ml	3	ML	VL	IH	ML	1	EA	1	12/14/2018	99/99/9999						
43975-0308-10		None		03/26/2018	99/99/9999	CYCLOPHOSPHAMIDE, 50 MG, ORAL	CYCLOPHOSPHAMIDE 50 mg	100	EA	BO	PO	EA	50	MG	1	03/26/2018	99/99/9999						
50242-0051-21		J9312		01/01/2019	99/99/9999	INJECTION, RITUXIMAB, 10 MG	RITUXAN (S.D.V., PF) 10 MG/ML	10	ML	VL	IV	ML	10	MG	1	01/01/2019	99/99/9999						
50242-0053-06		J9312		01/01/2019	99/99/9999	INJECTION, RITUXIMAB, 10 MG	RITUXAN (S.D.V., PF) 10 MG/ML	50	ML	VL	IV	ML	10	MG	1	01/01/2019	99/99/9999						
00085-0566-05		J0702		01/01/2002	02/28/2018	INJECTION, BETAMETHASONE ACETATE 3MG AND BETAMETHASONE SODIUM PHOSPHATE 3MG	CELESTONE SOLUSPAN (M.D.V.) 3 MG/ML	5	ML	VL	IJ	ML	3	MG	1	01/01/2002	02/28/2018						
00006-3845-71		J1335		04/16/2007	07/31/2018	INJECTION, ERTAPENEM SODIUM, 500 MG	INVANZ (SD ADD-VANTAGE) 1 GM	1	EA	VL	IJ	EA	500	MG	2	04/16/2007	07/31/2018						
00703-5040-01		J9000		01/01/2002	01/08/2019	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (M.D.V. POLYMER) 2 MG/ML	100	ML	VL	IV	EA	10	MG	0.2	01/01/2002	01/08/2019						
00703-5046-01		J9000		01/01/2002	01/08/2019	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (S.D.V. POLYMER) 2 MG/ML	25	ML	VL	IV	ML	10	MG	0.2	01/01/2002	01/08/2019						
00703-5043-03		J9000		01/01/2002	01/08/2019	INJECTION, DOXORUBICIN HYDROCHLORIDE, 10 MG	DOXORUBICIN HCL (S.D.V. POLYMER) 2 MG/ML	5	ML	VL	IV	ML	10	MG	0.2	01/01/2002	01/08/2019						
25021-0237-06		J9185		01/01/2015	10/03/2018	INJECTION, FLUDARABINE PHOSPHATE, 50 MG	FLUDARABINE PHOSPHATE (USP, SINGLE-DOSE, PF) 50 MG	1	EA	VL	IV	EA	50	MG	1	01/01/2015	10/03/2018						
25021-0155-15		J2185		03/27/2017	09/04/2018	INJECTION, MEROPENEM, 100 MG	MEROPENEM (PF, LATEX-FREE) 500 MG	10	EA	VL	IV	EA	100	MG	5	03/27/2017	09/04/2018						
25021-0156-30		J2185		03/27/2017	09/04/2018	INJECTION, MEROPENEM, 100 MG	MEROPENEM (PF, LATEX-FREE) 1 GM	10	EA	VL	IV	EA	100	MG	10	03/27/2017	09/04/2018						
00781-3442-20		J0171		01/16/2019	99/99/9999	INJECTION, ADRENALIN, EPINEPHRINE, 0.1 MG	AMPICILLIN-SULBACTAM (USP, PF, LATEX-FREE) 0.3 MG/0.3 ML	2	EA	SY	IJ	EA	0.1	MG	3	01/16/2019	99/99/9999						
71288-0005-20		J0295		01/07/2019	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM (USP, PF, LATEX-FREE) 1 GM/0.5 GM	10	EA	VL	IJ	EA	1.5	GM	1	01/07/2019	99/99/9999						
71288-0006-30		J0295		01/07/2019	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM (USP, PF, LATEX-FREE) 2 GM-1 GM	10	EA	VL	IJ	EA	1.5	GM	2	01/07/2019	99/99/9999						
71288-0007-75		J0295		01/07/2019	99/99/9999	INJECTION, AMPICILLIN SODIUM/SULBACTAM SODIUM, PER 1.5 GM	AMPICILLIN-SULBACTAM (PHARMACY BULK PACKAGE) 10 GM-5 GM	1	EA	BO	IV	EA	1.5	GM	10	01/07/2019	99/99/9999						
70436-0019-82		J0456		12/17/2018	99/99/9999	INJECTION, AZITHROMYCIN, 500 MG	AZITHROMYCIN (LYOPHILIZED) 500 MG	10	EA	VL	IV	EA	500	MG	1	12/17/2018	99/99/9999						
67457-0562-20		J0475		12/21/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (SDV) 0.5 MG/1 ML	20	ML	VL	IN	ML	10	MG	0.05	12/21/2018	99/99/9999						
67457-0563-20		J0475		12/21/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (SDV) 1 MG/1 ML	20	ML	VL	IN	ML	10	MG	0.1	12/21/2018	99/99/9999						
67457-0564-20		J0475		12/21/2018	99/99/9999	INJECTION, BACLOFEN, 10 MG	BACLOFEN (SDV) 1 MG/1 ML	20	ML	VL	IN	ML	10	MG	0.1	12/21/2018	99/99/9999						
00591-2416-30		J0604		01/02/2019	99/99/9999	CINACALCET, ORAL, 1 MG, (FOR ESRD ON DIALYSIS)	CINACALCET HYDROCHLORIDE (FILM-COATED) 30 MG	30	EA	BO	PO	EA	1	MG	30	01/02/2019	99/99/9999						
00591-2417-30		J0604		01/02/2019	99/99/9999	CINACALCET, ORAL, 1 MG, (FOR ESRD ON DIALYSIS)	CINACALCET HYDROCHLORIDE (FILM-COATED) 60 MG	30	EA	BO	PO	EA	1	MG	60	01/02/2019	99/99/9999						
00591-2418-30		J0604		01/02/2019	99/99/9999	CINACALCET, ORAL, 1 MG, (FOR ESRD ON DIALYSIS)	CINACALCET HYDROCHLORIDE (FILM-COATED) 90 MG	30	EA	BO	PO	EA	1	MG	90	01/02/2019	99/99/9999						
67457-0530-35		J0640		01/02/2019	99/99/9999	INJECTION, LEUCOVORIN CALCIUM, PER 50 MG	LEUCOVORIN CALCIUM (PF, LYOPHILIZED) 350 MG	1	EA	VL	IJ	EA	50	MG	7	01/02/2019	99/99/9999						
71288-0008-15		J0692		01/07/2019	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (SDV, PF, LATEX-FREE) 1 GM	10	EA	VL	IJ	EA	500	MG	2	01/07/2019	99/99/9999						
71288-0009-20		J0692		01/07/2019	99/99/9999	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (SDV, PF, LATEX-FREE) 2 GM	10	EA	VL	IJ	EA	500	MG	4	01/07/2019	99/99/9999						
70594-0023-04		J0770		01/16/2019	99/99/9999	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE 150 MG	12	EA	VL	IJ	EA	150	MG	1	01/16/2019	99/99/9999						
70594-0034-01		J0878		01/15/2019	99/99/9999	INJECTION, DAPTOMYCIN, 1 MG	DAPTOMYCIN (SDV, PF, LYOPHILIZED) 500 MG	1	EA	VL	IV	EA	1	MG	500	01/15/2019	99/99/9999						
64980-0467-99		J1071		01/14/2019	99/99/9999	INJECTION, TESTOSTERONE CYPIONATE, 1 MG	TESTOSTERONE CYPIONATE (SDV) 200 MG/1 ML	1	ML	VL	IM	ML	1	MG	200	01/14/2019	99/99/9999						
70436-0089-55		J1570		01/10/2019	99/99/9999	INJECTION, GANCICLOVIR SODIUM, 500 MG	GANCICLOVIR (USP, LYOPHILIZED) 500 MG	25	EA	VL	IV	EA	500	MG	1	01/10/2019	99/99/9999						
60505-0791-04		J1650		01/16/2019	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 30 MG/0.3 ML	0.3	ML	SY	IJ	ML	10	MG	10	01/16/2019	99/99/9999						
60505-0792-04		J1650		01/16/2019	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 40 MG/0.4 ML	0.4	ML	SY	IJ	ML	10	MG	10	01/16/2019	99/99/9999						
60505-0793-04		J1650		01/16/2019	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 60 MG/0.6 ML	0.6	ML	SY	IJ	ML	10	MG	10	01/16/2019	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
60505-0794-04		J1650		01/16/2019	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 80 MG/0.8 ML	0.8 ML	SY	IJ	ML		10 MG		10	01/16/2019	99/99/9999						
60505-0795-04		J1650		01/16/2019	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 100 MG/1 ML	1 ML	SY	IJ	ML		10 MG		10	01/16/2019	99/99/9999						
60505-0796-04		J1650		01/16/2019	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 120 MG/0.8 ML	0.8 ML	SY	IJ	ML		10 MG		15	01/16/2019	99/99/9999						
60505-0798-04		J1650		01/16/2019	99/99/9999	INJECTION, ENOXAPARIN SODIUM, 10 MG	ENOXAPARIN SODIUM (PF) 150 MG/1 ML	1 ML	SY	IJ	ML		10 MG		15	01/16/2019	99/99/9999						
00023-6082-10		J1750		01/01/2019	99/99/9999	INJECTION, IRON DEXTRAN, 50 MG	INFED (S.D.V.) 50 MG/1 ML	2 ML	VL	IJ	ML		50 MG		1	01/01/2019	99/99/9999						
00024-5926-05		J1817		01/28/2019	99/99/9999	INSULIN FOR ADMINISTRATION THROUGH DME (I.E., INSULIN PUMP) PER 50 UNITS	ADMELOG 100 U/1 ML	3 ML	VL	IJ	ML		50 UNITS		2	01/28/2019	99/99/9999						
00338-0072-25		J1885		01/30/2019	99/99/9999	INJECTION, KETOROLAC TROMETHAMINE, PER 15 MG	KETOROLAC TROMETHAMINE 30 MG/1 ML	1 ML	VL	IJ	ML		15 MG		2	01/30/2019	99/99/9999						
00143-9326-10		J2260		01/14/2019	99/99/9999	INJECTION, MILRINONE LACTATE, 5 MG	PREMIERPRO RX MILRINONE LACTATE (PF) 1 MG/1 ML	20 ML	VL	IV	ML		5 MG		0.2	01/14/2019	99/99/9999						
00781-3415-75		J2469		01/08/2019	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	PALONOSETRON HCL NOVAPLUS (SDV) 0.05 MG/1 ML	5 ML	VL	IV	ML		25 MCG		2	01/08/2019	99/99/9999						
67457-0379-25		J2501		12/21/2018	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL 0.002 MG/1 ML	1 ML	VL	IV	ML		1 MCG		2	12/21/2018	99/99/9999						
67457-0380-25		J2501		12/21/2018	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL 0.005 MG/1 ML	1 ML	VL	IV	ML		1 MCG		5	12/21/2018	99/99/9999						
67457-0389-25		J2501		12/21/2018	99/99/9999	INJECTION, PARICALCITOL, 1 MCG	PARICALCITOL 0.005 MG/1 ML	2 ML	VL	IV	ML		1 MCG		5	12/21/2018	99/99/9999						
25021-0162-68		J2700		01/22/2019	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN NOVAPLUS (USP,PF,LATEX-FREE) 2 GM	10 EA	VL	IJ	EA		250 MG		8	01/22/2019	99/99/9999						
25021-0163-68		J2700		01/22/2019	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN NOVAPLUS (PHARMACY BULK PACKAGE) 10 GM	10 EA	BO	IV	EA		250 MG		40	01/22/2019	99/99/9999						
70655-0103-95		J2700		01/02/2019	99/99/9999	INJECTION, OXACILLIN SODIUM, UP TO 250 MG	OXACILLIN 10 GM	10 EA	VL	IV	EA		250 MG		40	01/02/2019	99/99/9999						
70121-1478-07		J2710		12/20/2018	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 0.5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		1	12/20/2018	99/99/9999						
70121-1479-07		J2710		12/20/2018	99/99/9999	INJECTION, NEOSTIGMINE METHYLSULFATE, UP TO 0.5 MG	NEOSTIGMINE METHYLSULFATE (LATEX-FREE) 1 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		2	12/20/2018	99/99/9999						
63323-0778-10		J2800		01/11/2019	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (LATEX-FREE) 100 MG/1 ML	10 ML	VL	IJ	ML		10 ML		0.1	01/11/2019	99/99/9999						
71288-0716-10		J2800		01/21/2019	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (PF,LATEX-FREE) 100 MG/1 ML	10 ML	VL	IJ	ML		10 ML		0.1	01/21/2019	99/99/9999						
70121-1049-02		J3301		01/11/2019	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE 40 MG/1 ML	1 ML	VL	IJ	ML		10 MG		4	01/11/2019	99/99/9999						
70121-1651-01		J3301		12/28/2018	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE NOVAPLUS 40 MG/1 ML	1 ML	VL	IJ	ML		10 MG		4	12/28/2018	99/99/9999						
70121-1651-05		J3301		12/28/2018	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE NOVAPLUS 40 MG/1 ML	1 ML	VL	IJ	ML		10 MG		4	12/28/2018	99/99/9999						
70121-1652-01		J3301		12/28/2018	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE NOVAPLUS 40 MG/1 ML	5 ML	VL	IJ	ML		10 MG		4	12/28/2018	99/99/9999						
70121-1653-01		J3301		12/28/2018	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	TRIAMCINOLONE ACETONIDE NOVAPLUS 40 MG/1 ML	10 ML	VL	IJ	ML		10 MG		4	12/28/2018	99/99/9999						
70121-1654-01		J3301		12/28/2018	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	PREMIERPRO RX TRIAMCINOLONE ACETONIDE 40 MG/1 ML	5 ML	VL	IJ	ML		10 MG		4	12/28/2018	99/99/9999						
70121-1655-01		J3301		12/28/2018	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	PREMIERPRO RX TRIAMCINOLONE ACETONIDE 40 MG/1 ML	10 ML	VL	IJ	ML		10 MG		4	12/28/2018	99/99/9999						
70121-1657-01		J3301		12/28/2018	99/99/9999	INJECTION, TRIAMCINOLONE ACETONIDE, NOT OTHERWISE SPECIFIED, 10 MG	PREMIERPRO RX TRIAMCINOLONE ACETONIDE 40 MG/1 ML	1 ML	VL	IJ	ML		10 MG		4	12/28/2018	99/99/9999						
69680-0112-25		J3420		01/02/2019	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (MDV) 1000 MCG/1 ML	1 ML	VL	IJ	ML		1000 MCG		1	01/02/2019	99/99/9999						
69680-0113-99		J3420		01/02/2019	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (MDV) 1000 MCG/1 ML	10 ML	VL	IJ	ML		1000 MCG		1	01/02/2019	99/99/9999						
70436-0029-80		J3465		01/10/2019	99/99/9999	INJECTION, VORICONAZOLE, 10 MG	VORICONAZOLE (PF,LATEX-FREE) 200 MG	1 EA	VL	IV	EA		10 MG		20	01/10/2019	99/99/9999						
54288-0100-01		J3489		01/09/2019	99/99/9999	INJECTION, ZOLEDRONIC ACID, 1 MG	ZOLEDRONIC ACID (SINGLE-USE,LATEX-FREE) 4 MG/5 ML	5 ML	VL	IV	ML		1 MG		0.8	01/09/2019	99/99/9999						
70594-0026-02		J3490		01/07/2019	99/99/9999	UNCLASSIFIED DRUGS	BACITRACIN (LYOPHILIZED) 50000 U	10 EA	VL	IM	EA		1 EA		1	01/07/2019	99/99/9999						
00338-9147-30		J7060		01/28/2019	99/99/9999	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (MINI-BAG PLUS) 5%	100 ML	FC	IV	ML		500 ML		0.002	01/28/2019	99/99/9999						
00904-6785-04		J7518		12/24/2018	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (3X10) 180 MG	30 EA	BX	PO	EA		180 MG		1	12/24/2018	99/99/9999						
00904-6785-61		J7518		12/24/2018	99/99/9999	MYCOPHENOLIC ACID, ORAL, 180 MG	MYCOPHENOLIC ACID (10X10) 180 MG	100 EA	BX	PO	EA		180 MG		1	12/24/2018	99/99/9999						
60687-0395-83		J7613		12/26/2018	99/99/9999	ALBUTEROL INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3 ML	PC	IH	ML		1 MG		0.83	12/26/2018	99/99/9999						
60687-0405-83		J7620		12/26/2018	99/99/9999	ALBUTEROL UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE 3 MG/3 ML-0.5 MG/3 ML	3 ML	PC	IH	ML		3 MG		0.333333	12/26/2018	99/99/9999						
00093-6815-55		J7626		01/11/2019	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.25 MG/2 ML	2 ML	PC	IH	ML		0.5 MG		0.25	01/11/2019	99/99/9999						
00093-6816-55		J7626		01/11/2019	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.5 MG/2 ML	2 ML	PC	IH	ML		0.5 MG		0.5	01/11/2019	99/99/9999						
55150-0292-01		J7643		01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (SDV,LATEX-FREE) 0.2 MG/1 ML	1 ML	VL	IJ	ML		1 MG		0.2	01/08/2019	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
55150-0293-02		J7643		01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (SDV,LATEX-FREE) 0.2 MG/1 ML	2	ML	VL	IJ	ML	1	MG	0.2	01/08/2019	99/99/9999						
55150-0294-05		J7643		01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (MDV,LATEX-FREE) 0.2 MG/1 ML	5	ML	VL	IJ	ML	1	MG	0.2	01/08/2019	99/99/9999						
55150-0295-20		J7643		01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (MDV,LATEX-FREE) 0.2 MG/1 ML	20	ML	VL	IJ	ML	1	MG	0.2	01/08/2019	99/99/9999						
60687-0394-83		J7644		12/26/2018	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	12/26/2018	99/99/9999						
42195-0151-10		J8540		01/07/2019	99/99/9999	DEXAMETHASONE, ORAL, 0.25 MG	DEXAMETHASONE (USP) 1.5 MG	100	EA	BO	PO	EA	0.25	MG	6	01/07/2019	99/99/9999						
16729-0306-10		J9025		01/01/2019	99/99/9999	INJECTION, AZACITIDINE, 1 MG	AZACITIDINE (PF,LYOPHILIZED) 100 MG	1	EA	VL	IJ	EA	1	MG	100	01/01/2019	99/99/9999						
71288-0113-10		J9201		02/04/2019	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 200 MG	1	EA	VL	IV	EA	200	MG	1	02/04/2019	99/99/9999						
71288-0114-50		J9201		02/04/2019	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 1 GM	1	EA	VL	IV	EA	200	MG	5	02/04/2019	99/99/9999						
45963-0614-81		J9206		01/17/2019	99/99/9999	INJECTION, IRINOTECAN, 20 MG	PREMIERPRO RX IRINOTECAN HCL (PF,LATEX-FREE) 20 MG/1 ML	2	ML	VL	IV	ML	20	MG	1	01/17/2019	99/99/9999						
00143-9279-01		J9280		01/14/2019	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MITOMYCIN 20 MG	1	EA	VL	IV	EA	5	MG	4	01/14/2019	99/99/9999						
00143-9280-01		J9280		01/14/2019	99/99/9999	INJECTION, MITOMYCIN, 5 MG	MITOMYCIN 40 MG	1	EA	VL	IV	EA	5	MG	8	01/14/2019	99/99/9999						
59651-0007-15		Q0144		12/19/2018	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY BANANA) 100 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.02	12/19/2018	99/99/9999						
59651-0008-15		Q0144		12/19/2018	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY BANANA) 200 MG/5 ML	15	ML	BO	PO	ML	1	GM	0.04	12/19/2018	99/99/9999						
59651-0008-23		Q0144		12/19/2018	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY BANANA) 200 MG/5 ML	22.5	ML	BO	PO	ML	1	GM	0.04	12/19/2018	99/99/9999						
59651-0008-30		Q0144		12/19/2018	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (CHERRY BANANA) 200 MG/5 ML	30	ML	BO	PO	ML	1	GM	0.04	12/19/2018	99/99/9999						
65862-0642-90		Q0144		01/03/2019	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (3X3,FILM-COATED) 500 MG	9	EA	BX	PO	EA	1	GM	0.5	01/03/2019	99/99/9999						
60687-0395-83	KO	J7613	KO	12/26/2018	99/99/9999	ALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 1 MG	ALBUTEROL SULFATE 0.083%	3	ML	PC	IH	ML	1	MG	0.83	12/26/2018	99/99/9999						
60687-0405-83	KO	J7620	KO	12/26/2018	99/99/9999	ALBUTEROL, UP TO 2.5 MG AND IPRATROPIUM BROMIDE, UP TO 0.5 MG, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME	IPRATROPIUM BROMIDE-ALBUTEROL SULFATE 3 MG/3 ML-0.5 MG/3 ML	3	ML	PC	IH	ML	3	MG	0.333333	12/26/2018	99/99/9999						
00093-6815-55	KO	J7626	KO	01/11/2019	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.25 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	0.25	01/11/2019	99/99/9999						
00093-6816-55	KO	J7626	KO	01/11/2019	99/99/9999	BUDESONIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, UP TO 0.5 MG	BUDESONIDE (30X2ML,MICRONIZED) 0.5 MG/2 ML	2	ML	PC	IH	ML	0.5	MG	0.5	01/11/2019	99/99/9999						
55150-0292-01	KO	J7643	KO	01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (SDV,LATEX-FREE) 0.2 MG/1 ML	1	ML	VL	IJ	ML	1	MG	0.2	01/08/2019	99/99/9999						
55150-0293-02	KO	J7643	KO	01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (SDV,LATEX-FREE) 0.2 MG/1 ML	2	ML	VL	IJ	ML	1	MG	0.2	01/08/2019	99/99/9999						
55150-0294-05	KO	J7643	KO	01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (MDV,LATEX-FREE) 0.2 MG/1 ML	5	ML	VL	IJ	ML	1	MG	0.2	01/08/2019	99/99/9999						
55150-0295-20	KO	J7643	KO	01/08/2019	99/99/9999	GLYCOPYRROLATE, INHALATION SOLUTION, COMPOUNDED PRODUCT, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	GLYCOPYRROLATE (MDV,LATEX-FREE) 0.2 MG/1 ML	20	ML	VL	IJ	ML	1	MG	0.2	01/08/2019	99/99/9999						
60687-0394-83	KO	J7644	KO	12/26/2018	99/99/9999	IPRATROPIUM BROMIDE, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE FORM, PER MILLIGRAM	IPRATROPIUM BROMIDE (30X2.5ML,PF) 0.02%	2.5	ML	PC	IH	ML	1	MG	0.2	12/26/2018	99/99/9999						
00264-1290-55		J7799		01/01/2002	01/31/2018	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE HYPERTONIC (GLASS W/SOLID STOPPER) 70%	1000	ML	GC	IV	ML	1	EA	1	01/01/2002	01/31/2018						
57896-0782-01		Q0163		08/01/2002	01/24/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GERIDRYL (CAPLET) 25 MG	100	EA	NA	PO	EA	50	MG	0.5	08/01/2002	01/24/2014						
57896-0781-01		Q0163		08/01/2002	01/24/2014	DIPHENHYDRAMINE HYDROCHLORIDE, 50 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT TIME OF CHEMOTHERAPY TREATMENT NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	GERIDRYL 25 MG	100	EA	NA	PO	EA	50	MG	0.5	08/01/2002	01/24/2014						
00264-1510-36		J7060		01/01/2002	08/31/2017	5% DEXTROSE/WATER (500 ML = 1 UNIT)	DEXTROSE (100 ML PAB) 5%	25	ML	FC	IV	ML	500	ML	0.002	01/01/2002	08/31/2017						
00264-7612-20		J7799		01/01/2002	03/31/2019	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	DEXTROSE/SODIUM CHLORIDE (EXCEL) 5%-0.45%	250	ML	FC	IV	ML	1	EA	1	01/01/2002	03/31/2019						
00264-7578-20		J7799		01/01/2002	03/31/2019	NOC DRUGS, OTHER THAN INHALATION DRUGS, ADMINISTERED THROUGH DME	MANNITOL (EXCEL) 20%	250	ML	FC	IV	ML	1	EA	1	01/01/2002	03/31/2019						
00703-4434-11		J9206		02/28/2008	05/02/2018	INJECTION, IRINOTECAN, 20 MG	IRINOTECAN HYDROCHLORIDE (1X5ML,SINGLE DOSE) 20 MG/ML	5	ML	VL	IV	ML	20	MG	1	02/28/2008	05/02/2018						
60505-0686-04		J2543		09/21/2009	02/20/2019	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SDV) 2 GM-0.25 GM	10	EA	VL	IV	EA	1.125	GM	2	09/21/2009	02/20/2019						
60505-0681-04		J0692		06/19/2007	02/04/2019	INJECTION, CEFEPIME HYDROCHLORIDE, 500 MG	CEFEPIME (USP) 2 GM	10	EA	VL	IJ	EA	500	MG	4	06/19/2007	02/04/2019						
00093-4146-64	KO	J7614	KO	04/29/2013	02/15/2019	COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X4,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	04/29/2013	02/15/2019						
00093-4146-64		J7614		04/29/2013	02/15/2019	COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X4,PF) 0.63 MG/3 ML	3	ML	PC	IH	ML	0.5	MG	0.42	04/29/2013	02/15/2019						
42367-0121-21		J9171		01/29/2016	09/30/2018	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (AF) 20 MG/1 ML	1	ML	VL	IV	ML	1	MG	20	01/29/2016	09/30/2018						
42367-0121-25		J9171		01/29/2016	09/30/2018	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (AF) 20 MG/1 ML	4	ML	VL	IV	ML	1	MG	20	01/29/2016	09/30/2018						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
42367-0121-29		J9171		01/29/2016	09/30/2018	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (AF) 20 MG/1 ML COLISTIMETHATE (LYOPHILIZED CAKE) 150 MG	8 ML	VL	IV	ML		1 MG		20	01/29/2016	09/30/2018						
39822-0617-02	J0770			07/01/2016	02/08/2019	INJECTION, COLISTIMETHATE SODIUM, UP TO 150 MG	COLISTIMETHATE (LYOPHILIZED CAKE) 150 MG	12 EA	VL	IJ	EA		150 MG		1	07/01/2016	02/08/2019						
39822-2120-01	J9171			05/05/2017	02/22/2019	INJECTION, DOCETAXEL, 1 MG	DOCETAXEL (SDV) 20 MG/1 ML	1 ML	VL	IV	ML		1 MG		20	05/05/2017	02/22/2019						
00409-1112-01	J0594			02/28/2019	99/99/9999	INJECTION, BUSULFAN, 1 MG	BUSULFAN (8X10ML,SINGLE-USE) 6 MG/1 ML	10 ML	VL	IV	ML		1 MG		6	02/28/2019	99/99/9999						
16714-0890-01	J0641			03/14/2019	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (PF) 10 MG/1 ML	17.5 ML	VL	IV	ML		0.5 MG		20	03/14/2019	99/99/9999						
16714-0915-01	J0641			03/14/2019	99/99/9999	INJECTION, LEVOLEUCOVORIN CALCIUM, 0.5 MG	LEVOLEUCOVORIN CALCIUM (PF) 10 MG/1 ML	25 ML	VL	IV	ML		0.5 MG		20	03/14/2019	99/99/9999						
51754-5060-01	J0702			02/04/2019	99/99/9999	INJECTION, BETAMETHASONE ACETATE 3 MG AND BETAMETHASONE SODIUM PHOSPHATE 3 MG	BETAMETHASONE ACETATE-BETAMETHASONE SODIUM PHOSPH (MDV) 3 MG/1 ML-3 MG/1 ML	5 ML	VL	IJ	ML		6 MG		1	02/04/2019	99/99/9999						
55150-0241-10	J0883			02/07/2019	99/99/9999	INJECTION, ARGATROBAN, 1 MG (FOR NON-ESRD USE)	ARGATROBAN (LATEX-FREE) 1 MG/1 ML PALONOSETRON HCL (PF,LATEX-FREE) 0.05 MG/1 ML	50 ML	VL	IV	ML		1 MG		1	02/07/2019	99/99/9999						
55150-0186-05	J2469			02/07/2019	99/99/9999	INJECTION, PALONOSETRON HCL, 25 MCG	ARGATROBAN (LATEX-FREE) 1 MG/1 ML PALONOSETRON HCL (PF,LATEX-FREE) 0.05 MG/1 ML	5 ML	VL	IV	ML		25 MCG		2	02/07/2019	99/99/9999						
60505-6156-00	J2543			02/15/2019	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 2 GM-0.25 GM	1 EA	VL	IV	EA		1.125 GM		2	02/15/2019	99/99/9999						
60505-6156-04	J2543			02/15/2019	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 2 GM-0.25 GM	10 EA	VL	IV	EA		1.125 GM		2	02/15/2019	99/99/9999						
60505-6157-00	J2543			02/15/2019	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 3 GM-0.375 GM	1 EA	VL	IV	EA		1.125 GM		3	02/15/2019	99/99/9999						
60505-6157-04	J2543			02/15/2019	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 3 GM-0.375 GM	10 EA	VL	IV	EA		1.125 GM		3	02/15/2019	99/99/9999						
60505-6159-00	J2543			02/15/2019	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 4 GM-0.5 GM	1 EA	VL	IV	EA		1.125 GM		4	02/15/2019	99/99/9999						
60505-6159-04	J2543			02/15/2019	99/99/9999	INJECTION, PIPERACILLIN SODIUM/TAZOBACTAM SODIUM, 1 GRAM/0.125 GRAMS (1.125 GRAMS)	PIPERACILLIN AND TAZOBACTAM (SINGLE DOSE,PF) 4 GM-0.5 GM	10 EA	VL	IV	EA		1.125 GM		4	02/15/2019	99/99/9999						
70860-0653-10	J2800			01/02/2019	99/99/9999	INJECTION, METHOCARBAMOL, UP TO 10 ML	METHOCARBAMOL (PF,LATEX-FREE) 100 MG/1 ML	10 ML	VL	IJ	ML		10 ML		0.1	01/02/2019	99/99/9999						
70069-0171-10	J3420			02/15/2019	99/99/9999	INJECTION, VITAMIN B-12 CYANOCOBALAMIN, UP TO 1000 MCG	CYANOCOBALAMIN (MDV) 1000 MCG/1 ML	30 ML	VL	IJ	ML		1000 MCG		1	02/15/2019	99/99/9999						
55566-1000-01	J3490			02/14/2019	99/99/9999	UNCLASSIFIED DRUGS	GANIRELIX ACETATE 250 MCG/0.5 ML SODIUM CHLORIDE (RFID TAGGED,PF) 0.9%	0.5 ML	SR	SC	ML		1 EA		1	02/14/2019	99/99/9999						
00019-1188-27	A4217			01/08/2019	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (PF) 0.9%	125 ML	SR	IJ	ML		500 ML		0.002	01/08/2019	99/99/9999						
00019-1188-75	A4216			01/08/2019	99/99/9999	STERILE WATER,SALINE AND/OR DEXTROSE, DILUENT/FLUSH, 10 ML	SODIUM CHLORIDE (PF) 0.9%	50 ML	SR	IJ	ML		10 ML		0.1	01/08/2019	99/99/9999						
00019-1188-81	A4217			01/08/2019	99/99/9999	STERILE WATER/SALINE, 500 ML	SODIUM CHLORIDE (PF) 0.9%	125 ML	SR	IJ	ML		500 ML		0.002	01/08/2019	99/99/9999						
66689-0307-08	J7517			02/15/2019	99/99/9999	MYCOPHENOLATE MOFETIL, ORAL, 250 MG	MYCOPHENOLATE MOFETIL (BANANA) 200 MG/1 ML	175 ML	BO	PO	ML		250 MG		0.8	02/15/2019	99/99/9999						
66689-0347-02	J7520			02/01/2019	99/99/9999	SIROLIMUS, ORAL, 1 MG	SIROLIMUS 1 MG/1 ML	60 ML	BO	PO	ML		1 MG		1	02/01/2019	99/99/9999						
00093-4146-56	J7614			02/15/2019	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X5,PF) 0.63 MG/3 ML	3 ML	PC	IH	ML		0.5 MG		0.42	02/15/2019	99/99/9999						
00093-4146-56	J7614	KO		02/15/2019	99/99/9999	LEVALBUTEROL, INHALATION SOLUTION, FDA-APPROVED FINAL PRODUCT, NON-COMPOUNDED, ADMINISTERED THROUGH DME, UNIT DOSE, 0.5 MG	LEVALBUTEROL (6X5,PF) 0.63 MG/3 ML	3 ML	PC	IH	ML		0.5 MG		0.42	02/15/2019	99/99/9999						
42291-0017-01	J8499			01/21/2019	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	100 EA	BO	PO	EA		1 MG		1	01/21/2019	99/99/9999						
42291-0017-50	J8499			01/21/2019	99/99/9999	PRESCRIPTION DRUG, ORAL, NON CHEMOTHERAPEUTIC, NOS	ACYCLOVIR 200 MG	500 EA	BO	PO	EA		1 MG		1	01/21/2019	99/99/9999						
25021-0239-05	J9201			02/19/2019	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 38 MG/1 ML	5.26 ML	VL	IV	ML		200 MG		0.19	02/19/2019	99/99/9999						
25021-0239-26	J9201			02/19/2019	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 38 MG/1 ML	26.3 ML	VL	IV	ML		200 MG		0.19	02/19/2019	99/99/9999						
25021-0239-52	J9201			02/19/2019	99/99/9999	INJECTION, GEMCITABINE HYDROCHLORIDE, 200 MG	GEMCITABINE (PF,LATEX-FREE) 38 MG/1 ML	52.6 ML	VL	IV	ML		200 MG		0.19	02/19/2019	99/99/9999						
10019-0927-01	J9208			01/18/2019	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE NOVAPLUS 1 GM	1 EA	VL	IV	EA		1 GM		1	01/18/2019	99/99/9999						
10019-0929-03	J9208			01/18/2019	99/99/9999	INJECTION, IFOSFAMIDE, 1 GRAM	IFOSFAMIDE NOVAPLUS 3 GM	1 EA	VL	IV	EA		1 GM		3	01/18/2019	99/99/9999						
10019-0951-05	J9209			01/18/2019	99/99/9999	INJECTION, MESNA, 200 MG	MESNA NOVAPLUS (MDV) 100 MG/1 ML	10 ML	VL	IV	ML		200 MG		0.5	01/18/2019	99/99/9999						
50742-0405-10	J9263			02/20/2019	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	10 ML	VL	IV	ML		0.5 MG		10	02/20/2019	99/99/9999						
50742-0406-20	J9263			02/20/2019	99/99/9999	INJECTION, OXALIPLATIN, 0.5 MG	OXALIPLATIN (PF) 5 MG/1 ML	20 ML	VL	IV	ML		0.5 MG		10	02/20/2019	99/99/9999						
00904-6708-06	Q0144			02/25/2019	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (5X10,FILM-COATED) 250 MG	50 EA	BX	PO	EA		1 GM		0.25	02/25/2019	99/99/9999						
00904-6708-61	Q0144			02/25/2019	99/99/9999	AZITHROMYCIN DIHYDRATE, ORAL, CAPSULES/POWDER, 1 GRAM	AZITHROMYCIN (10X10,FILM-COATED) 250 MG	100 EA	BX	PO	EA		1 GM		0.25	02/25/2019	99/99/9999						
60687-0252-86	Q0162			01/28/2019	99/99/9999	ONDANSETRON 1 MG, ORAL, FDA APPROVED PRESCRIPTION ANTI-EMETIC, FOR USE AS A COMPLETE THERAPEUTIC SUBSTITUTE FOR AN IV ANTI-EMETIC AT THE TIME OF CHEMOTHERAPY TREATMENT, NOT TO EXCEED A 48 HOUR DOSAGE REGIMEN	ONDANSETRON 4 MG/5 ML	5 ML	CP	PO	ML		1 MG		0.8	01/28/2019	99/99/9999						
59923-0703-05	None			01/25/2019	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	5 EA	BO	PO	EA		5 MG		1	01/25/2019	99/99/9999						
59923-0704-14	None			01/25/2019	99/99/9999	TEMODAR, 5 MG, ORAL	TEMOZOLOMIDE 5 MG	14 EA	BO	PO	EA		5 MG		1	01/25/2019	99/99/9999						
59923-0713-05	None			01/25/2019	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE 250 MG	5 EA	BO	PO	EA		250 MG		1	01/25/2019	99/99/9999						
59923-0707-05	None			01/25/2019	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	5 EA	BO	PO	EA		100 MG		1	01/25/2019	99/99/9999						
59923-0708-14	None			01/25/2019	99/99/9999	TEMODAR, 100 MG, ORAL	TEMOZOLOMIDE 100 MG	14 EA	BO	PO	EA		100 MG		1	01/25/2019	99/99/9999						
59923-0705-05	None			01/25/2019	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	5 EA	BO	PO	EA		20 MG		1	01/25/2019	99/99/9999						
59923-0706-14	None			01/25/2019	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 20 MG	14 EA	BO	PO	EA		20 MG		1	01/25/2019	99/99/9999						

NDC	NDC Mod	HCPCS	HCPCS Mod	Relationship Start Date	Relationship End Date	HCPCS Description	NDC Label	Number of Items In NDC Package	NDC Package Measure	NDC Package Type	Route of Administration	Billing Units	HCPCS Amount #1	HCPCS Measure #1	CF	Start Date #1	End Date #1	Prior Start Date #2	Prior End Date #2	Prior Conversion Factor #2	Prior Start Date #3	Prior End Date #3	Prior Conversion Factor #3
59923-0709-05		None		01/25/2019	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	5 EA	BO	PO	EA	20 MG		7	01/25/2019	99/99/9999							
59923-0710-14		None		01/25/2019	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 140 MG	14 EA	BO	PO	EA	20 MG		7	01/25/2019	99/99/9999							
59923-0711-05		None		01/25/2019	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	5 EA	BO	PO	EA	20 MG		9	01/25/2019	99/99/9999							
59923-0712-14		None		01/25/2019	99/99/9999	TEMOZOLOMIDE, 20 MG, ORAL	TEMOZOLOMIDE 180 MG	14 EA	BO	PO	EA	20 MG		9	01/25/2019	99/99/9999							
69097-0516-07		None		01/28/2019	99/99/9999	CYCLOPHOSPHAMIDE, 25 MG, ORAL	CYCLOPHOSPHAMIDE (HARD GELATIN) 25 MG	100 EA	PC	PO	EA	25 MG		1	01/28/2019	99/99/9999							
69097-0517-07		None		01/28/2019	99/99/9999	CYCLOPHOSPHAMIDE, 50 MG, ORAL	CYCLOPHOSPHAMIDE (HARD GELATIN) 50 MG	100 EA	PC	PO	EA	50 MG		1	01/28/2019	99/99/9999							