

# New Brunswick Drug Plans Formulary

September 2023

## TABLE OF CONTENTS

	Page
Introduction.....	I
New Brunswick Drug Plans.....	I
Exclusions.....	II
Legend.....	III

### Anatomical Therapeutic Chemical (ATC) Classification of Drugs

A	Alimentary Tract and Metabolism	1
B	Blood and Blood Forming Organs	24
C	Cardiovascular System	34
D	Dermatologicals	85
G	Genito Urinary System and Sex Hormones	94
H	Systemic Hormonal Preparations excluding Sex Hormones	105
J	Antiinfectives for Systemic Use	112
L	Antineoplastic and Immunomodulating Agents	137
M	Musculo-Skeletal System	164
N	Nervous System	173
P	Antiparasitic Products, Insecticides and Repellants	244
R	Respiratory System	245
S	Sensory Organs	256
V	Various	264

### Appendices

I-A	Abbreviations of Dosage forms.....	A - 1
I-B	Abbreviations of Routes.....	A - 4
I-C	Abbreviations of Units.....	A - 6
I-D	Abbreviations of Manufacturers' Names.....	A - 8
II	Extemporaneous Preparations.....	A - 10
III	Special Authorization Criteria.....	A - 12
IV	Provisional Benefits.....	A - 120

## New Brunswick Drug Plans Formulary

### Introduction

The Government of New Brunswick provides prescription drug coverage to eligible New Brunswick residents through the New Brunswick Prescription Drug Program, the New Brunswick Drug Plan and other government sponsored plans (collectively known as the New Brunswick Drug Plans). See below for a complete list of plans.

The New Brunswick Drug Plans Formulary is a list of the drugs which are eligible benefits under the NB Drug Plans. The Formulary is updated monthly and all drugs considered for listing as benefits must be reviewed according to the [Drug Review Process](#).

Most drugs listed in the Formulary are "regular" benefits which are reimbursed with no criteria or prior approval requirements. Some drugs are special authorization benefits and have specific criteria that must be met before they are approved for reimbursement (see Formulary Appendix III). The process and forms for submitting special authorization requests is posted on the NB Drug Plans [website](#). Certain drug products are not eligible benefits and are identified on the exclusion list.

### The New Brunswick Drug Plans

Plan	Group Code
<b>The New Brunswick Prescription Drug Program (NBPDP)</b>	
Seniors	A
Correctional Services	C
Social Development Clients	F
Adult Residential Facilities	E
Children in Care of the Minister of Social Development and Special Needs Children	G
Nursing Home Residents	V
Cystic Fibrosis	B
Growth Hormone Deficiency	T
HIV/AIDS	U
Multiple Sclerosis	H
Organ Transplant	R
<b>The New Brunswick Drug Plan</b>	D
<b>Other Government Sponsored Plans</b>	
Public Health Plan	I
Tuberculosis	P
Extra-Mural Program Clients	W
Medical Abortion Program	J

Details regarding the New Brunswick Drug Plans are available on the Government of New Brunswick's [website](#).

## Exclusions

The following classes of products, except those specifically listed in the Formulary, are excluded as benefits under the New Brunswick Drug Plans.

- Drugs not authorized for sale and use in Canada (e.g., drugs obtained through Health Canada's Special Access Program, experimental or investigational drugs)
- Non-prescription drugs<sup>1</sup>
- Natural health products<sup>1</sup> (e.g., vitamins and minerals, herbal remedies, probiotics, homeopathic medicines, traditional medicines)
- Cannabis or cannabis products
- Nutritional supplements and food products
- Weight loss products
- Products for the treatment of erectile/sexual dysfunction, or infertility
- Drugs for the prevention of travel acquired diseases
- Products for esthetic or cosmetic purposes
- Soaps, cleansers, shampoos, antiseptics, or disinfectants
- Diagnostic agents and point-of-care testing kits
- Medical supplies, devices and equipment (e.g., prostheses, first aid supplies, ostomy supplies, diabetes test strips and syringes, etc.)
- Vaccines

<sup>1</sup>To be listed in the Formulary, a non-prescription drug or natural health product must be recommended by an expert advisory committee based on evidence that supports its clinical efficacy and cost effectiveness.

## Legend

**M01<sup>1</sup> ANTIINFLAMMATORY AND ANTIRHEUMATIC PRODUCTS**

**M01A<sup>2</sup> ANTIINFLAMMATORY AND ANTIRHEUMATIC PRODUCTS, NON-STERIODS**

**M01AE<sup>3</sup> PROPIONIC ACID DERIVATIVES**

M01AE02<sup>4</sup> NAPROXEN

ECT<sup>5</sup> OrI<sup>5</sup> 250mg<sup>5</sup>

6

7

8

9

Naproxen EC 02350785 SAS ADEFGVW

Teva-Naprox EC 02243312 TEV ADEFGVW

ECT OrI 375mg

Naprosyn E 02162415 MTP ADEFGVW

Apo-Naproxen EC 02246700 APX ADEFGVW

Naproxen EC 02350793 SAS ADEFGVW

Mylan-Naproxen EC 02243432 MYL ADEFGVW

10 pms-Naproxen EC (Disc/non disp Mar 4/19) 02294702 PMS ADEFGVW

Teva-Naprox EC 02243313 TEV ADEFGVW

<sup>1</sup> Second level ATC, therapeutic subgroup

<sup>2</sup> Third level ATC, pharmacological subgroup

<sup>3</sup> Fourth level ATC, chemical subgroup

<sup>4</sup> Fifth level ATC, chemical substance

<sup>5</sup> Dosage form, route and strength. Strength represents the amount of ingredients present in a solid dosage form (tablet) or in one gram or one millilitre of a product (cream, liquid, etc.)

<sup>6</sup> Brand or manufacturers' product name approved by Health Canada.

<sup>7</sup> Drug Identification Number (DIN)

<sup>8</sup> Manufacturers' identification code. See Appendix I-D for details.

<sup>9</sup> Drug plans for which the product is a benefit. See page II for details. Please note that products marked (SA) are only eligible for coverage under NB Drug Plans through special authorization.

<sup>10</sup> Manufacturer has discontinued this product. It will be deleted from the Formulary on the date indicated.

**A ALIMENTARY TRACT AND METABOLISM****A01 STOMATOLOGICAL PREPARATIONS****A01A STOMATOLOGICAL PREPARATIONS****A01AC CORTICOSTEROIDS FOR LOCAL ORAL TREATMENT**

A01AC01 TRIAMCINOLONE

Pst Den 0.1%

Oracort 01964054 TAR ACDEFGV

**A01AD OTHER AGENTS FOR LOCAL ORAL TREATMENT**

A01AD02 BENZYDAMINE

Liq Buc 0.15%

Odan-Benzylamine 02463105 ODN ACDEFGV

pms-Benzylamine 02239537 PMS ACDEFGV

**A02 DRUGS FOR ACID RELATED DISORDERS****A02A ANTACIDS****A02AD COMBINATIONS AND COMPLEXES OF ALUMINIUM, CALCIUM AND MAGNESIUM COMPOUNDS**

A02AD01 ORDINARY SALT COMBINATIONS

ALUMINIUM / MAGNESIUM

Sus Orl 45.6 mg / 40 mg

Diovol 01966529 CHU G

**A02AH ANTACIDS WITH SODIUM BICARBONATE**

A02AH01 SODIUM BICARBONATE

Tab Orl 500 mg

Jamp-Sodium Bicarbonate 80030520 JPC (SA)

Sandoz Sodium Bicarbonate 80022194 SDZ (SA)

**A02B DRUGS FOR PEPTIC ULCER AND GASTRO-OESOPHAGEAL REFLUX DISEASE (GORD)****A02BA H2-RECEPTOR ANTAGONISTS**

A02BA01 CIMETIDINE

Tab Orl 200 mg

Cimetidine 00584215 AAP ACDEFGV

Tab Orl 300 mg

Cimetidine 00487872 AAP ACDEFGV

A02BA02 RANITIDINE

Liq Orl 15 mg/mL

Apo-Ranitidine 02280833 APX CDEFGVW

Tab Orl 150 mg

Apo-Ranitidine 00733059 APX ACDEFGVW

Jamp-Ranitidine 02463717 JPC ACDEFGVW

Mar-Ranitidine 02443708 MAR ACDEFGVW

Mint-Ranitidine 02526379 MNT ACDEFGVW

pms-Ranitidine 02242453 PMS ACDEFGVW

Ranitidine 02353016 SAS ACDEFGVW

A02BA02 RANITIDINE

Tab Orl 300 mg

Apo-Ranitidine 00733067 APX ACDEFGVW  
 Jamp-Ranitidine 02463725 JPC ACDEFGVW  
 Mar-Ranitidine 02443716 MAR ACDEFGVW  
 Mint-Ranitidine 02526387 MNT ACDEFGVW  
 pms-Ranitidine 02242454 PMS ACDEFGVW

A02BA03 FAMOTIDINE

Tab Orl 20 mg

Jamp Famotidine 02507749 JPC ACDEFGV  
 Teva-Famotidine 02022133 TEV ACDEFGV

Tab Orl 40 mg

Jamp Famotidine 02507757 JPC ACDEFGV  
 Teva-Famotidine 02022141 TEV ACDEFGV

**A02BB PROSTAGLANDINS**

A02BB01 MISOPROSTOL

Tab Orl 100 mcg

Misoprostol 02244022 AAP ACDEFGV

Tab Orl 200 mcg

Misoprostol 02244023 AAP ACDEFGJV

**A02BC PROTON PUMP INHIBITORS**

A02BC01 OMEPRAZOLE

SRC Orl 20 mg

Losec 00846503 AZE ACDEFGV  
 Apo-Omeprazole 02245058 APX ACDEFGV  
 Omeprazole 02348691 SAS ACDEFGV  
 Omeprazole 02411857 SIV ACDEFGV  
 pms-Omeprazole 02320851 PMS ACDEFGV  
 Sandoz Omeprazole 02296446 SDZ ACDEFGV  
 Jamp-Omeprazole 02420198 JPC ACDEFGV  
 Nat-Omeprazole DR 02439549 NAT ACDEFGV  
 Omeprazole 02416549 AHI ACDEFGV  
 Omeprazole Magnesium 02504294 SAS ACDEFGV  
 Teva-Omeprazole 02295415 TEV ACDEFGV

SRT Orl 20 mg

A02BC02 PANTOPRAZOLE  
 PANTOPRAZOLE MAGNESIUM

A02BC02 PANTOPRAZOLE  
PANTOPRAZOLE MAGNESIUM

ECT	Orl	40 mg	Tecta	02267233	TAK	ACDEFGV
			Mylan-Pantoprazole T	02408570	MYL	ACDEFGV
			Pantoprazole Magnesium	02441853	ATS	ACDEFGV
			Pantoprazole T	02466147	SAS	ACDEFGV
			Pantoprazole T	02519534	SIV	ACDEFGV
			Teva-Pantoprazole Magnesium	02440628	TEV	ACDEFGV

PANTOPRAZOLE SODIUM

ECT	Orl	20 mg	Pantoloc	02241804	TAK	ACDEFGV
			Apo-Pantoprazole	02292912	APX	ACDEFGV
			Jamp Pantoprazole Sodium	02392615	JPC	ACDEFGV
			Jamp-Pantoprazole	02408414	JPC	ACDEFGV
			Pantoprazole-20	02428172	SIV	ACDEFGV
			Sandoz Pantoprazole	02301075	SDZ	ACDEFGV
			Teva-Pantoprazole	02285479	TEV	ACDEFGV

ECT	Orl	40 mg	Pantoloc	02229453	TAK	ACDEFGV
			Apo-Pantoprazole	02292920	APX	ACDEFGV
			Auro-Pantoprazole	02415208	ARO	ACDEFGV
			Jamp Pantoprazole Sodium	02392623	JPC	ACDEFGV
			Jamp-Pantoprazole	02357054	JPC	ACDEFGV
			M-Pantoprazole	02467372	MRA	ACDEFGV
			Mar-Pantoprazole	02416565	MAR	ACDEFGV
			Mint-Pantoprazole	02417448	MNT	ACDEFGV
			NRA-Pantoprazole	02471825	NRA	ACDEFGV
			Pantoprazole	02318695	PDL	ACDEFGV
			Pantoprazole	02437945	PMS	ACDEFGV
			Pantoprazole	02370808	SAS	ACDEFGV
			Pantoprazole-40	02428180	SIV	ACDEFGV
			pms-Pantoprazole	02307871	PMS	ACDEFGV
			Sandoz Pantoprazole	02301083	SDZ	ACDEFGV
			Taro-Pantoprazole	02305046	SUN	ACDEFGV
			Teva-Pantoprazole	02285487	TEV	ACDEFGV

A02BC03 LANSOPRAZOLE



A02BC03 LANSOPRAZOLE

SRC Orl 15 mg

Prevacid 02165503 ABB (SA)  
 Apo-Lansoprazole 02293811 APX (SA)  
 Lansoprazole 02433001 PMS (SA)  
 Lansoprazole 02357682 SAS (SA)  
 Lansoprazole 02385767 SIV (SA)  
 Mylan-Lansoprazole 02353830 MYL (SA)  
 Sandoz Lansoprazole 02385643 SDZ (SA)  
 Taro-Lansoprazole 02402610 SUN (SA)  
 Teva-Lansoprazole 02280515 TEV (SA)

SRC Orl 30 mg

Prevacid 02165511 ABB (SA)  
 Apo-Lansoprazole 02293838 APX (SA)  
 Lansoprazole 02433028 PMS (SA)  
 Lansoprazole 02357690 SAS (SA)  
 Lansoprazole 02410389 SIV (SA)  
 Mylan-Lansoprazole 02353849 MYL (SA)  
 Sandoz Lansoprazole 02385651 SDZ (SA)  
 Taro-Lansoprazole 02402629 SUN (SA)  
 Teva-Lansoprazole 02280523 TEV (SA)

SRT Orl 15 mg

Prevacid FasTab 02249464 ABB (SA)

SRT Orl 30 mg

Prevacid FasTab 02249472 ABB (SA)

A02BC04 RABEPRAZOLE

ECT Orl 10 mg

Pariet 02243796 JAN ACDEFGV  
 Jamp Rabeprazole 02415283 JPC ACDEFGV  
 pms-Rabeprazole EC 02310805 PMS ACDEFGV  
 Rabeprazole 02385449 SIV ACDEFGV  
 Rabeprazole EC 02356511 SAS ACDEFGV  
 Sandoz Rabeprazole 02314177 SDZ ACDEFGV  
 Taro-Rabeprazole 02298074 SUN ACDEFGV

ECT Orl 20 mg

Pariet 02243797 JAN ACDEFGV  
 Jamp Rabeprazole 02415291 JPC ACDEFGV  
 pms-Rabeprazole EC 02310813 PMS ACDEFGV  
 Rabeprazole 02385457 SIV ACDEFGV  
 Rabeprazole EC 02356538 SAS ACDEFGV  
 Sandoz Rabeprazole 02314185 SDZ ACDEFGV  
 Taro-Rabeprazole 02298082 SUN ACDEFGV

**A02BX OTHER DRUGS FOR PEPTIC ULCER AND GASTROESOPHAGEAL REFLUX DISEASE (GORD)****A02BX02 SUCRALFATE**

Sus	Orl	1 g / 5 mL	Sulcrate Suspension Plus	02103567	AXC	ACDEFGV
Tab	Orl	1 g	Sulcrate	02100622	AXC	ACDEFGV
			Apo-Sulcralfate	02125250	APX	ACDEFGV
			Teva-Sulcralfate	02045702	TEV	ACDEFGV

**A03 DRUGS FOR FUNCTIONAL GASTROINTESTINAL DISORDERS****A03A DRUGS FOR FUNCTIONAL GASTROINTESTINAL DISORDERS****A03AA SYNTHETIC ANTICHOLINERGICS, ESTERS WITH TERTIARY AMINO GROUP****A03AA05 TRIMEBUTINE**

Tab	Orl	100 mg	Trimebutine	02245663	AAP	ACDEFGV
Tab	Orl	200 mg	Trimebutine	02245664	AAP	ACDEFGV

**A03AA07 DICYCLOVERINE (DICYCLOMINE)**

Cap	Orl	10 mg	Protylol	00287709	PDL	ACDEFGV
Tab	Orl	20 mg	Jamp-Dicyclomine	02366088	JPC	ACDEFGV

**A03AB SYNTHETIC ANTICHOLINERGICS, QUATERNARY AMMONIUM COMPOUNDS****A03AB02 GLYCOPYRRONIUM BROMIDE (GLYCOPYRROLATE)**

Liq	Inj	0.2 mg/mL	Glycopyrrolate	02039508	SDZ	ACDEFGVW
			Glycopyrrolate Injection USP	02473879	STR	ACDEFGVW
Liq	Inj	0.4 mg / 2 mL	Glycopyrrolate Injection USP	02473895	STR	ACDEFGVW
Liq	Inj	4 mg / 20 mL	Glycopyrrolate Injection USP	02473887	STR	ACDEFGVW

**A03AX OTHER DRUGS FOR FUNCTIONAL GASTROINTESTINAL DISORDERS****A03AX04 PINAVERIUM**

Tab	Orl	50 mg	Dicetel	01950592	ABB	ACDEFGV
			Pinaverium	02469677	AAP	ACDEFGV
Tab	Orl	100 mg	Dicetel	02230684	ABB	ACDEFGV
			Pinaverium	02469685	AAP	ACDEFGV

**A03F PROPULSIVES****A03FA PROPULSIVES****A03FA01 METOCLOPRAMIDE**

Liq	Inj	5 mg/mL	Metoclopramide	02185431	SDZ	ACDEFVW
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A03FA01	METOCLOPRAMIDE						
Syr	Orl	1 mg/mL	pms-Metoclopramide	02230433	PMS	ACDEFGVW	
Tab	Orl	5 mg	Mar-Metoclopramide	02517795	MAR	ACDEFGVW	
			pms-Metoclopramide	02230431	PMS	ACDEFGVW	
Tab	Orl	10 mg	pms-Metoclopramide	02230432	PMS	ACDEFGVW	
A03FA03	DOMPERIDONE						
Tab	Orl	10 mg	Apo-Domperidone	02103613	APX	ACDEFGVW	
			Domperidone	02350440	SAS	ACDEFGVW	
			Domperidone	02238341	SIV	ACDEFGVW	
			Jamp-Domperidone	02369206	JPC	ACDEFGVW	
			Mar-Domperidone	02403870	MAR	ACDEFGVW	
			pms-Domperidone	02236466	PMS	ACDEFGVW	
			PRZ-Domperidone	02462834	PRZ	ACDEFGVW	
			Ran-Domperidone	02268078	SUN	ACDEFGVW	
			Teva-Domperidone	01912070	TEV	ACDEFGVW	

**A04 ANTIEMETICS AND ANTINAUSEANTS**

**A04A ANTIEMETICS AND ANTINAUSEANTS**

**A04AA SEROTONIN (5HT3) ANTAGONISTS**

A04AA01	ONDANSETRON						
Liq	Inj	2 mg/mL	Ondansetron Hydrochloride Dihydrate	02274418	SDZ	W (SA)	
			Ondansetron Injection USP	02279436	SDZ	W (SA)	
			Ondansetron Injection USP	02462257	STR	W (SA)	
			Ondansetron Injection USP (PF)	02464578	STR	W (SA)	
Liq	Orl	4 mg / 5 mL	Zofran	02229639	NVR	(SA)	
			Jamp Ondansetron	02490617	JPC	(SA)	
			Ondansetron	02291967	APX	(SA)	
ODT	Slg	4 mg	Zofran ODT	02239372	SDZ	(SA)	
			Athena-Ondansetron ODT	02444674	AHC	(SA)	
			Auro-Ondansetron ODT	02511282	ARO	(SA)	
			Mar-Ondansetron ODT	02514966	MAR	(SA)	
			Mint-Ondansetron ODT	02487330	MNT	(SA)	
			Ondansetron ODT	02519232	JPC	(SA)	
			Ondansetron ODT	02481723	SDZ	(SA)	
			Ondissolve	02389983	TAK	(SA)	
			pms-Ondansetron ODT	02519445	PMS	(SA)	

## A04AA01 ONDANSETRON

ODT Slg 8 mg

Zofran ODT	02239373	SDZ	(SA)
Athena-Ondansetron ODT	02444682	AHC	(SA)
Auro-Ondansetron ODT	02511290	ARO	(SA)
Mar-Ondansetron ODT	02514974	MAR	(SA)
Mint-Ondansetron ODT	02487349	MNT	(SA)
Ondansetron ODT	02519240	JPC	(SA)
Ondansetron ODT	02481731	SDZ	(SA)
Ondissolve	02389991	TAK	(SA)
pms-Ondansetron ODT	02519453	PMS	(SA)

Tab Orl 4 mg

Zofran	02213567	NVR	W (SA)
Apo-Ondansetron	02288184	APX	W (SA)
CCP-Ondansetron	02458810	CCM	W (SA)
Jamp-Ondansetron	02313685	JPC	W (SA)
Mar-Ondansetron	02371731	MAR	W (SA)
Mint-Ondansetron	02305259	MNT	W (SA)
Mylan-Ondansetron	02297868	MYL	W (SA)
Nat-Ondansetron	02417839	NAT	W (SA)
Ondansetron	02421402	SAS	W (SA)
pms-Ondansetron	02258188	PMS	W (SA)
Sandoz Ondansetron	02274310	SDZ	W (SA)
Septa-Ondansetron	02376091	SPT	W (SA)
Teva-Ondansetron	02296349	TEV	W (SA)

Tab Orl 8 mg

Zofran	02213575	NVR	W (SA)
Apo-Ondansetron	02288192	APX	W (SA)
CCP-Ondansetron	02458802	CCM	W (SA)
Jamp-Ondansetron	02313693	JPC	W (SA)
Mar-Ondansetron	02371758	MAR	W (SA)
Mint-Ondansetron	02305267	MNT	W (SA)
Mylan-Ondansetron	02297876	MYL	W (SA)
Nat-Ondansetron	02417847	NAT	W (SA)
Ondansetron	02421410	SAS	W (SA)
pms-Ondansetron	02258196	PMS	W (SA)
Sandoz Ondansetron	02274329	SDZ	W (SA)
Septa-Ondansetron	02376105	SPT	W (SA)
Teva-Ondansetron	02296357	TEV	W (SA)

## A04AA55 PALONOSETRON, COMBINATIONS

A04AA55 PALONOSETRON, COMBINATIONS  
PALONOSETRON / NETUPITANT

Cap Orl 300 mg / 0.5 mg Akynzeo 02468735 KNI (SA)

**A04AD OTHER ANTIEMETICS**

A03BB01 BUTYLSCOPOLAMINE

Liq Inj 20 mg/mL Buscopan 00363839 SNC ACDEFGVW

Hyoscine Butylbromide 02229868 SDZ ACDEFGVW

Tab Orl 10 mg Buscopan 00363812 SNC ACDEFGVW

Accel-Hyoscine 02512335 ACC ACDEFGVW

A04AD01 SCOPOLAMINE

Srd Trd 1.5 mg Transderm-V 80024336 SDZ AEFVW

A04AD11 NABILONE

Cap Orl 0.25 mg Cesamet 02312263 BSL ACDEFVW

pms-Nabilone 02380897 PMS ACDEFVW

Teva-Nabilone 02392925 TEV ACDEFVW

Cap Orl 0.5 mg Cesamet 02256193 BSL ACDEFVW

pms-Nabilone 02380900 PMS ACDEFVW

Teva-Nabilone 02384884 TEV ACDEFVW

Cap Orl 1 mg Cesamet 00548375 BSL ACDEFVW

pms-Nabilone 02380919 PMS ACDEFVW

Teva-Nabilone 02384892 TEV ACDEFVW

A04AD12 APREPITANT

Cap Orl 80 mg Emend 02298791 FRS (SA)

Cap Orl 125 mg Emend 02298805 FRS (SA)

Kit Orl 80 mg, 125 mg Emend-Tri-Pack 02298813 FRS (SA)

N05CM05 SCOPOLAMINE

Liq Inj 0.4 mg/mL Scopolamine Hydrobromide 02242810 OMG ACDEFVW

Liq Inj 0.6 mg/mL Scopolamine Hydrobromide 02242811 OMG ACDEFVW

**A05 BILE AND LIVER THERAPY**

**A05A BILE THERAPY**

**A05AA BILE ACID PREPERATIONS****A05AA02 URSODEOXYCHOLIC ACID (URSODIOL)**

Tab	Orl	250 mg	GLN-Ursodiol	02426900	GLM	ACDEFGV
			Jamp-Ursodiol	02472392	JPC	ACDEFGV
			pms-Ursodiol C	02273497	PMS	ACDEFGV
			Ursodiol C	02515520	SAS	ACDEFGV

Tab	Orl	500 mg	Urso DS (Disc/non disp Mar 6/24)	02245894	AXC	ACDEFGV
			GLN-Ursodiol	02426919	GLM	ACDEFGV
			Jamp-Ursodiol	02472406	JPC	ACDEFGV
			pms-Ursodiol C	02273500	PMS	ACDEFGV
			Ursodiol C	02515539	SAS	ACDEFGV

**A05AA04 OBETICHOLIC ACID**

Tab	Orl	5 mg	Ocaliva	02463121	ADZ	(SA)
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Tab	Orl	10 mg	Ocaliva	02463148	ADZ	(SA)
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**A06 DRUGS FOR CONSTIPATION****A06A DRUGS FOR CONSTIPATION****A06AD OSMOTICALLY ACTING LAXATIVES****A06AD11 LACTULOSE**

Syr	Orl	667 mg	Jamp-Lactulose	02295881	JPC	(SA)
			Lactulose	02412268	SAS	(SA)
			pms-Lactulose	00703486	PMS	(SA)
			pms-Lactulose-pharma	02469391	PMS	(SA)
			ratio-Lactulose	00854409	TEV	(SA)

**A07 ANTIDIARRHEALS, INTESTINAL ANTIINFLAMMATORY/ANTIINFECTIVE AGENTS****A07A INTESTINAL ANTIINFECTIVES****A07AA ANTIBIOTICS****A07AA02 NYSTATIN**

Sus	Orl	100 000 IU/mL	Jamp-Nystatin	02433443	JPC	ACDEFGVW
			pms-Nystatin Suspension	00792667	PMS	ACDEFGVW
			Teva-Nystatin	02194201	TEV	ACDEFGVW

**A07AA11 RIFAXIMIN**

Tab	Orl	550 mg	Zaxine	02410702	SAX	(SA)
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**A07AA12 FIDAXOMICIN**

Tab	Orl	200 mg	Dificid	02387174	FRS	W (SA)
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**A07D ANTIPROPULSIVES****A07DA ANTIPROPULSIVES**

A07DA01 DIPHENOXYLATE

DIPHENOXYLATE / ATROPINE

Tab Orl 2.5 mg / 0.025 mg

Lomotil 00036323 PFI ACDEFGV

A07DA03 LOPERAMIDE

Liq Orl 0.2 mg/mL

pms-Loperamide Hydrochloride 02016095 PMS AEFVG

Tab Orl 2 mg

pms-Loperamide 02228351 PMS AEFVG

Teva-Loperamide 02132591 TEV AEFVG

**A07E INTESTINAL ANTIINFLAMMATORY AGENTS****A07EA CORTICOSTEROIDS ACTING LOCALLY**

A07EA06 BUDESONIDE

Cap Orl 3 mg

Entocort 02229293 AZE ACDEFGV

Enm Rt 2.3 mg

Entocort 02052431 AZE ACDEFGV

**A07EC AMINOSALICYLIC ACID AND SIMILAR AGENTS**

A07EC01 SULFASALAZINE

ECT Orl 500 mg

Salazopyrin EN 02064472 PFI ACDEFGV

pms-Sulfasalazine EC 00598488 PMS ACDEFGV

Tab Orl 500 mg

Salazopyrin 02064480 PFI ACDEFGV

pms-Sulfasalazine 00598461 PMS ACDEFGV

A07EC02 MESALAZINE

Aer Rt 1 g

Mezera 02474026 AVI ACDEFGV

ECT Orl 400 mg

Teva-5-ASA 02171929 TEV ACDEFGV

ECT Orl 500 mg

Salofalk 02112787 ABV ACDEFGV

ERT Orl 500 mg

Pentasa 02099683 FEI ACDEFGV

ERT Orl 1 000 mg

Pentasa 02399466 FEI ACDEFGV

Sup Rt 500 mg

Salofalk 02112760 ABV ACDEFGV

A07EC02		MESALAZINE					
Sup	Rt	1 g		Salofalk	02242146	ABV	ACDEFGV
				Mezera	02474018	AVI	ACDEFGV
				Pentasa	02153564	FEI	ACDEFGV
Susp	Rt	1 g / 100 mL		Pentasa	02153521	FEI	ACDEFGV
Susp	Rt	2 g/60 mL		Salofalk	02112795	ABV	ACDEFGV
Susp	Rt	4 g / 100 mL		Pentasa	02153556	FEI	ACDEFGV
Susp	Rt	4 g / 60 mL		Salofalk	02112809	ABV	ACDEFGV
Tab	Orl	1.2 g		Mezavant	02297558	TAK	ACDEFGV

A07EC03		OLSALAZINE					
Cap	Orl	250 mg		Dipentum	02063808	SLP	ACDEFGV

**A07F ANTIDIARRHEAL MICROORGANISMS**

**A07FA ANTIDIARRHEAL MICROORGANISMS**

A07FA01		LACTIC ACID PRODUCING ORGANISMS					
Cap	Orl	1 B		Bacid	80017987	ERF	AEFGV

**A09 DIGESTIVES, INCLUDING ENZYMES**

**A09A DIGESTIVES, INCLUDING ENZYMES**

**A09AA ENZYME PREPARATIONS**

A09AA02		MULTIENZYMES (LIPASE, PROTEASE, ETC)					
Cap	Orl	35 000 U / 10 000 U / 40 000 U		Cotazym	00263818	ORG	ABCDEFGV
ECC	Orl	4 200 U / 10 000 U / 17 500 U		Pancrease MT 4	00789445	VVS	ABCDEFGV
ECC	Orl	6 000 U / 19 000 U / 30 000 U		Creon 6 Minimicrospheres	02415194	BGP	ABCDEFGV
ECC	Orl	10 000 U / 730 U / 11 200 U		Creon 10 Minimicrospheres	02200104	BGP	ABCDEFGV
ECC	Orl	10 500 U / 25 000 U / 43 750 U		Pancrease MT 10	00789437	VVS	ABCDEFGV
ECC	Orl	10 800 U / 45 000 U / 42 000 U		Cotazym ECS 8	00502790	ORG	ABCDEFGV
ECC	Orl	16 800 U / 40 000 U / 70 000 U		Pancrease MT 16	00789429	VVS	ABCDEFGV



A09AA02 MULTIENZYMES (LIPASE, PROTEASE, ETC)

ECC	Orl	25 000 U / 1 600 U / 25 500 U	Creon 25 Minimicrospheres	01985205	BGP	ABCDEFGV
ECC	Orl	25 000 U / 100 000 U / 100 000 U	Cotazym ECS 20	00821373	ORG	ABCDEFGV
ECC	Orl	35 000 U / 2 240 U / 35 700 U	Creon 35 Minimicrospheres	02494639	BGP	ABCDEFGV
Gran	Orl	5 000 U / 5 100 U / 320 U	Creon Minimicrospheres Micro	02445158	BGP	ABCDEFGV
Tab	Orl	10 440 U / 39 150 U / 39 150 U	Viokace	02230019	ARN	ABCDEFGV
Tab	Orl	20 880 U / 78 300 U / 78 300 U	Viokace	02241933	ARN	ABCDEFGV

**A10 DRUGS USED IN DIABETES**

**A10A INSULINS AND ANALOGUES**

**A10AB INSULINS AND ANALOGUES FOR INJECTION, FAST-ACTING**

A10AB01 INSULIN (HUMAN)

Liq	Inj	100 U/mL	Humulin R	00586714	LIL	ACDEFGV
			Humulin R (cartridge)	01959220	LIL	ACDEFGV
			Novolin GE Toronto	02024233	NNO	ACDEFGV
			Novolin GE Toronto (penfill)	02024284	NNO	ACDEFGV

A10AB04 INSULIN LISPRO

Liq	Inj	100 U/mL	Admelog	02469901	SAV	ACDEFGV
			Admelog (cartridge)	02469898	SAV	ACDEFGV
			Admelog (SoloSTAR)	02469871	SAV	ACDEFGV

A10AB05 INSULIN ASPART

Liq	Inj	100 U/mL	Kirsty (prefilled pen)	02520974	BGP	ACDEFGV
			NovoRapid	02245397	NNO	ACDEFGV
			Trurapi (cartridge)	02506564	SAV	ACDEFGV
			Trurapi (SoloSTAR)	02506572	SAV	ACDEFGV

A10AB06 INSULIN GLULISINE

Liq	Inj	100 U/mL	Apidra	02279460	SAV	ACDEFGV
			Apidra (cartridge)	02279479	SAV	ACDEFGV
			Apidra Solostar	02294346	SAV	ACDEFGV

**A10AC INSULINS AND ANALOGUES FOR INJECTION, INTERMEDIATE-ACTING**

A10AC01 INSULIN (HUMAN)

A10AC01 INSULIN (HUMAN)

Sus Inj 100 U/mL

Humulin N 00587737 LIL ACDEFGV  
 Humulin N (cartridge) 01959239 LIL ACDEFGV  
 Humulin N (KwikPen) 02403447 LIL ACDEFGV  
 Novolin GE NPH 02024225 NNO ACDEFGV  
 Novolin GE NPH (penfill) 02024268 NNO ACDEFGV

Sus Inj 500 U/mL

Entuzity (KwikPen) 02466864 LIL ACDEFGV

**A10AD INSULINS AND ANALOGUES FOR INJECTION, INTERMEDIATE-ACTING AND FAST-ACTING**

A10AD01 INSULIN (HUMAN)

Sus Inj 30 U / 70 U

Humulin 30/70 00795879 LIL ACDEFGV  
 Humulin 30/70 (cartridge) 01959212 LIL ACDEFGV  
 Novolin GE 30/70 02024217 NNO ACDEFGV  
 Novolin GE 30/70 (penfill) 02025248 NNO ACDEFGV

Sus Inj 40 U / 60 U

Novolin GE 40/60 (penfill) (Disc/non disp Mar 31/24) 02024314 NNO ACDEFGV

Sus Inj 50 U / 50 U

Novolin GE 50/50 (penfill) (Disc/non disp Mar 31/24) 02024322 NNO ACDEFGV

**A10AE INSULINS AND ANALOGUES FOR INJECTION, LONG-ACTING**

A10AE04 INSULIN GLARGINE

Liq Inj 100 U/mL

Basaglar cartridge 02444844 LIL ACDEFGV  
 Basaglar KwikPen 02461528 LIL ACDEFGV  
 Semglee (prefilled pen) 02526441 BGP ACDEFGV

A10AE05 INSULIN DETEMIR

Liq Inj 100 U/mL

Levemir FlexTouch 02412829 NNO (SA)  
 Levemir Penfill Cartridge 02271842 NNO (SA)

A10AE06 INSULIN DEGLUDEC

Liq Inj 100 U/mL

Tresiba Flextouch 02467879 NNO ACDEFGV

Liq Inj 200 U/mL

Tresiba Flextouch 02467887 NNO ACDEFGV

**A10B BLOOD GLUCOSE LOWERING DRUGS, EXCLUDING INSULINS**

**A10BA BIGUANIDES**

A10BA02 METFORMIN

A10BA02 METFORMIN

Tab Orl 500 mg

Glucophage 02099233 SAV ACDEFGV  
 Act Metformin 02257726 TEV ACDEFGV  
 Auro-Metformin 02438275 ARO ACDEFGV  
 Jamp-Metformin 02380196 JPC ACDEFGV  
 Mar-Metformin 02378620 MAR ACDEFGV  
 Metformin 02353377 SAS ACDEFGV  
 Metformin FC 02385341 SIV ACDEFGV  
 pms-Metformin 02223562 PMS ACDEFGV  
 pmsc-Metformin 02520303 PMS ACDEFGV  
 Pro-Metformin 02314908 PDL ACDEFGV  
 PRZ-Metformin 02531895 PRZ ACDEFGV  
 Sandoz Metformin FC 02246820 SDZ ACDEFGV

Tab Orl 850 mg

Glucophage 02162849 SAV ACDEFGV  
 Act Metformin 02257734 TEV ACDEFGV  
 Auro-Metformin 02438283 ARO ACDEFGV  
 Jamp-Metformin 02380218 JPC ACDEFGV  
 Mar-Metformin 02378639 MAR ACDEFGV  
 Metformin 02353385 SAS ACDEFGV  
 Metformin FC 02385368 SIV ACDEFGV  
 pms-Metformin 02242589 PMS ACDEFGV  
 pmsc-Metformin 02520311 PMS ACDEFGV  
 Pro-Metformin 02314894 PDL ACDEFGV  
 PRZ-Metformin 02531909 PRZ ACDEFGV  
 Sandoz Metformin FC 02246821 SDZ ACDEFGV

Tab Orl 1000 mg

PRZ-Metformin 02534673 PRZ ACDEFGV

**A10BB SULFONAMIDES, UREA DERIVATIVES**

A10BB01 GLIBENCLAMIDE (GLYBURIDE)

Tab Orl 2.5 mg

Apo-Glyburide 01913654 APX ACDEFGV  
 Glyburide 02350459 SAS ACDEFGV  
 Teva-Glyburide 01913670 TEV ACDEFGV

Tab Orl 5 mg

Apo-Glyburide 01913662 APX ACDEFGV  
 Glyburide 02350467 SAS ACDEFGV  
 Teva-Glyburide 01913689 TEV ACDEFGV

A10BB09 GLICLAZIDE

A10BB09 GLICLAZIDE

ERT	Orl	30 mg	Diamicon MR	02242987	SEV	ACDEFGV
			Apo-Gliclazide MR	02297795	APX	ACDEFGV
			Gliclazide MR	02524856	SAS	ACDEFGV
			Jamp-Gliclazide MR	02429764	JPC	ACDEFGV
			Mint-Gliclazide MR	02423286	MNT	ACDEFGV
			Mylan-Gliclazide MR	02438658	MYL	ACDEFGV
			Sandoz Gliclazide MR	02461323	SDZ	ACDEFGV
			Taro-Gliclazide MR	02463571	SUN	ACDEFGV

ERT	Orl	60 mg	Diamicon MR	02356422	SEV	ACDEFGV
			Apo-Gliclazide MR	02407124	APX	ACDEFGV
			Gliclazide MR	02524864	SAS	ACDEFGV
			Mint-Gliclazide MR	02423294	MNT	ACDEFGV
			Sandoz Gliclazide MR	02461331	SDZ	ACDEFGV
			Taro-Gliclazide MR	02439328	SUN	ACDEFGV

Tab	Orl	80 mg	Apo-Gliclazide	02245247	APX	ACDEFGV
			Gliclazide	02287072	SAS	ACDEFGV
			Teva-Gliclazide	02238103	TEV	ACDEFGV

**A10BD COMBINATIONS OF ORAL BLOOD GLUCOSE LOWERING DRUGS**

A10BD07 METFORMIN AND SITAGLIPTIN

ERT	Orl	500 mg / 50 mg	Janumet XR	02416786	FRS	ACDEFGV
			Apo-Sitagliptin/Metformin XR	02506270	APX	ACDEFGV
			Sandoz Sitagliptin-Metformin XR	02529106	SDZ	ACDEFGV

ERT	Orl	1 000 mg / 50 mg	Janumet XR	02416794	FRS	ACDEFGV
			Apo-Sitagliptin/Metformin XR	02506289	APX	ACDEFGV
			Sandoz Sitagliptin-Metformin XR	02529114	SDZ	ACDEFGV

ERT	Orl	1 000 mg / 100 mg	Janumet XR	02416808	FRS	ACDEFGV
			Apo-Sitagliptin/Metformin XR	02506297	APX	ACDEFGV
			Sandoz Sitagliptin-Metformin XR	02529122	SDZ	ACDEFGV

Tab	Orl	500 mg / 50 mg	Janumet	02333856	FRS	ACDEFGV
			Apo-Sitagliptin Malate/Metformin HCl	02509415	APX	ACDEFGV
			Sandoz Sitagliptin-Metformin	02503956	SDZ	ACDEFGV

A10BD07 METFORMIN AND SITAGLIPTIN

Tab	Orl	850 mg / 50 mg	Janumet	02333864	FRS	ACDEFGV
			Apo-Sitagliptin Malate/Metformin HCl	02509423	APX	ACDEFGV
			Sandoz Sitagliptin-Metformin	02503964	SDZ	ACDEFGV
Tab	Orl	1 000 mg / 50 mg	Janumet	02333872	FRS	ACDEFGV
			Apo-Sitagliptin Malate/Metformin HCl	02509431	APX	ACDEFGV
			Sandoz Sitagliptin-Metformin	02503972	SDZ	ACDEFGV

A10BD10 METFORMIN AND SAXAGLIPTIN

Tab	Orl	500 mg / 2.5 mg	Komboglyze	02389169	AZE	(SA)
Tab	Orl	850 mg / 2.5 mg	Komboglyze	02389177	AZE	(SA)
Tab	Orl	1 000 mg / 2.5 mg	Komboglyze	02389185	AZE	(SA)

A10BD11 METFORMIN AND LINAGLIPTIN

Tab	Orl	500 mg / 2.5 mg	Jentaduetto	02403250	BOE	(SA)
Tab	Orl	850 mg / 2.5 mg	Jentaduetto	02403269	BOE	(SA)
Tab	Orl	1 000 mg / 2.5 mg	Jentaduetto	02403277	BOE	(SA)

A10BD15 METFORMIN AND DAPAGLIFLOZIN

Tab	Orl	850 mg / 5 mg	XigDuo	02449935	AZE	(SA)
			Auro-Dapagliflozin/Metformin	02533073	ARO	(SA)
Tab	Orl	1 000 mg / 5 mg	XigDuo	02449943	AZE	(SA)
			Auro-Dapagliflozin/Metformin	02533081	ARO	(SA)

A10BD20 METFORMIN AND EMPAGLIFLOZIN

Tab	Orl	500 mg / 5 mg	Synjardy	02456575	BOE	(SA)
Tab	Orl	500 mg / 12.5 mg	Synjardy	02456605	BOE	(SA)
Tab	Orl	850 mg / 5 mg	Synjardy	02456583	BOE	(SA)
Tab	Orl	850 mg / 12.5 mg	Synjardy	02456613	BOE	(SA)
Tab	Orl	1000 mg / 5 mg	Synjardy	02456591	BOE	(SA)
Tab	Orl	1000 mg / 12.5 mg	Synjardy	02456621	BOE	(SA)

**A10BF ALPHA GLUCOSIDASE INHIBITORS**

A10BF01 ACARBOSE

Tab Orl 50 mg

Glucobay (Disc/non disp Apr 18/24) 02190885 BAY ACDEFGV

Acarbose Tablets 02493780 STD ACDEFGV

Mar-Acarbose 02494078 MAR ACDEFGV

Tab Orl 100 mg

Glucobay (Disc/non disp Apr 18/24) 02190893 BAY ACDEFGV

Acarbose Tablets 02493799 STD ACDEFGV

Mar-Acarbose 02494086 MAR ACDEFGV

**A10BG THIAZOLINEDIONES**

A10BG03 PIOGLITAZONE

Tab Orl 15 mg

Ach-Pioglitazone 02391600 AHI ACDEFGV

Act Pioglitazone 02302861 TEV ACDEFGV

Apo-Pioglitazone 02302942 APX ACDEFGV

Jamp-Pioglitazone 02397307 JPC ACDEFGV

Mint-Pioglitazone 02326477 MNT ACDEFGV

pms-Pioglitazone 02303124 PMS ACDEFGV

Tab Orl 30 mg

Ach-Pioglitazone 02339587 AHI ACDEFGV

Act Pioglitazone 02302888 TEV ACDEFGV

Apo-Pioglitazone 02302950 APX ACDEFGV

Jamp-Pioglitazone 02365529 JPC ACDEFGV

Mint-Pioglitazone 02326485 MNT ACDEFGV

pms-Pioglitazone 02303132 PMS ACDEFGV

Tab Orl 45 mg

Ach-Pioglitazone 02339595 AHI ACDEFGV

Act Pioglitazone 02302896 TEV ACDEFGV

Apo-Pioglitazone 02302977 APX ACDEFGV

Jamp-Pioglitazone 02365537 JPC ACDEFGV

Mint-Pioglitazone 02326493 MNT ACDEFGV

pms-Pioglitazone 02303140 PMS ACDEFGV

**A10BH DIPEPTIDYL PEPTIDASE 4 (DPP-4) INHIBITORS**

A10BH01 SITAGLIPTIN

## A10BH01 SITAGLIPTIN

Tab Orl 25 mg

Januvia	02388839	FRS	ACDEFGV
ACH-Sitagliptin	02512475	AHI	ACDEFGV
Apo-Sitagliptin Malate	02508656	APX	ACDEFGV
Auro-Sitagliptin	02529866	ARO	ACDEFGV
Jamp Sitagliptin	02534134	JPC	ACDEFGV
Sandoz Sitagliptin	02504049	SDZ	ACDEFGV
Sitagliptin	02529033	SIV	ACDEFGV
Taro-Sitagliptin Fumarate	02531631	TAR	ACDEFGV

Tab Orl 50 mg

Januvia	02388847	FRS	ACDEFGV
ACH-Sitagliptin	02512483	AHI	ACDEFGV
Apo-Sitagliptin Malate	02508664	APX	ACDEFGV
Auro-Sitagliptin	02529874	ARO	ACDEFGV
Jamp Sitagliptin	02534142	JPC	ACDEFGV
Sandoz Sitagliptin	02504057	SDZ	ACDEFGV
Sitagliptin	02529041	SIV	ACDEFGV
Taro-Sitagliptin Fumarate	02531658	TAR	ACDEFGV

Tab Orl 100 mg

Januvia	02303922	FRS	ACDEFGV
ACH-Sitagliptin	02512491	AHI	ACDEFGV
Apo-Sitagliptin Malate	02508672	APX	ACDEFGV
Auro-Sitagliptin	02529882	ARO	ACDEFGV
Jamp Sitagliptin	02534150	JPC	ACDEFGV
Sandoz Sitagliptin	02504065	SDZ	ACDEFGV
Sitagliptin	02529068	SIV	ACDEFGV
Taro-Sitagliptin Fumarate	02531666	TAR	ACDEFGV

## A10BH03 SAXAGLIPTIN

Tab Orl 2.5 mg

Onglyza	02375842	AZE	(SA)
Apo-Saxagliptin	02507471	APX	(SA)
Sandoz Saxagliptin	02468603	SDZ	(SA)

Tab Orl 5 mg

Onglyza	02333554	AZE	(SA)
Apo-Saxagliptin	02507498	APX	(SA)
Sandoz Saxagliptin	02468611	SDZ	(SA)

## A10BH05 LINAGLIPTIN

Tab Orl 5 mg

Trajenta	02370921	BOE	ACDEFGV
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## A10BJ GLUCAGON-LIKE PEPTIDE-1 (GLP-1) ANALOGUES

A10BJ03	LIXISENATIDE						
	Liq	SC	0.05 mg/mL		Adlyxine	02464276	SAV (SA)
	Liq	SC	0.1 mg/mL		Adlyxine	02464284	SAV (SA)
A10BJ06	SEMAGLUTIDE						
	Liq	SC	2 mg / 1.5 mL		Ozempic (autoinjector)	02471477	NNO (SA)
	Liq	SC	4 mg / 3 mL		Ozempic (autoinjector)	02471469	NNO (SA)

**A10BK SODIUM-GLUCOSE CO-TRANSPORTER 2 (SGLT2) INHIBITORS**

A10BK01	DAPAGLIFLOZIN						
	Tab	Orl	5 mg		Forxiga	02435462	AZE ACDEFGV
					Apo-Dapagliflozin	02527189	APX ACDEFGV
					Auro-Dapagliflozin	02531402	ARO ACDEFGV
					GLN-Dapagliflozin	02519852	GLM ACDEFGV
					Jamp Dapagliflozin	02531364	JPC ACDEFGV
					M-Dapagliflozin	02535297	MRA ACDEFGV
					pms-Dapagliflozin	02531550	PMS ACDEFGV
					Sandoz Dapagliflozin	02518732	SDZ ACDEFGV
	Tab	Orl	10 mg		Forxiga	02435470	AZE ACDEFGV
					Apo-Dapagliflozin	02527197	APX ACDEFGV
					Auro-Dapagliflozin	02531410	ARO ACDEFGV
					GLN-Dapagliflozin	02519860	GLM ACDEFGV
					Jamp Dapagliflozin	02531372	JPC ACDEFGV
					M-Dapagliflozin	02535300	MRA ACDEFGV
					pms-Dapagliflozin	02531569	PMS ACDEFGV
					Sandoz Dapagliflozin	02518740	SDZ ACDEFGV
A10BK02	CANAGLIFLOZIN						
	Tab	Orl	100 mg		Invokana	02425483	JAN (SA)
	Tab	Orl	300 mg		Invokana	02425491	JAN (SA)
A10BK03	EMPAGLIFLOZIN						
	Tab	Orl	10 mg		Jardiance	02443937	BOE (SA)
	Tab	Orl	25 mg		Jardiance	02443945	BOE (SA)

**A10BX OTHER BLOOD GLUCOSE LOWERING DRUGS, EXCL INSULINS**

A10BX02 REPAGLINIDE



A10BX02 REPAGLINIDE

Tab Orl 0.5 mg

Gluconorm 02239924 NNO ACDEFGV  
 Act Repaglinide 02321475 TEV ACDEFGV  
 Auro-Repaglinide 02424258 ARO ACDEFGV  
 Jamp-Repaglinide 02354926 JPC ACDEFGV  
 Sandoz Repaglinide 02357453 SDZ ACDEFGV

Tab Orl 1 mg

Gluconorm 02239925 NNO ACDEFGV  
 Act Repaglinide 02321483 TEV ACDEFGV  
 Auro-Repaglinide 02424266 ARO ACDEFGV  
 Jamp-Repaglinide 02354934 JPC ACDEFGV  
 Sandoz Repaglinide 02357461 SDZ ACDEFGV

Tab Orl 2 mg

Gluconorm 02239926 NNO ACDEFGV  
 Act Repaglinide 02321491 TEV ACDEFGV  
 Auro-Repaglinide 02424274 ARO ACDEFGV  
 Jamp-Repaglinide 02354942 JPC ACDEFGV  
 Sandoz Repaglinide 02357488 SDZ ACDEFGV

**A11 VITAMINS**

**A11C VITAMIN A AND D, INCLUDING COMBINATIONS OF THE TWO**

**A11CC VITAMIN D AND ANALOGUES**

A11CC01 ERGOCALCIFEROL

Cap Orl 50 000 IU

D-Forte 02237450 SDZ ACDEFGV

A11CC03 ALFACALCIDOL

Cap Orl 0.25 mcg

One-Alpha 00474517 XPI ACDEFGV  
 Sandoz Alfacalcidol 02533316 SDZ ACDEFGV

Cap Orl 1 mcg

One-Alpha 00474525 XPI ACDEFGV  
 Sandoz Alfacalcidol 02533324 SDZ ACDEFGV

Liq Orl 2 mcg/mL

One-Alpha 02240329 XPI ACDEFGV

A11CC04 CALCITRIOL

Cap Orl 0.25 mcg

Rocaltrol 00481823 SLP ACDEFGV  
 Calcitriol-Odan 02431637 ODN ACDEFGV  
 pms-Calcitriol 02495899 PMS ACDEFGV  
 Taro-Calcitriol 02485710 TAR ACDEFGV

A11CC04    CALCITRIOL

Cap    Orl    0.5 mcg

Rocaltrol    00481815    SLP    ACDEFGV  
Calcitriol-Odan    02431645    ODN    ACDEFGV  
pms-Calcitriol    02495902    PMS    ACDEFGV  
Taro-Calcitriol    02485729    TAR    ACDEFGV

A11CC05    CHOLECALCIFEROL

Tab    Orl    1 000 IU

Vitamin D    80000436    JAM    EF-18G

**A11E        VITAMIN B-COMPLEX, INCLUDING COMBINATIONS**

**A11EB       VITAMIN B-COMPLEX WITH VITAMIN C**

A11EB99    VITAMIN B-COMPLEX WITH VITAMIN C

Tab    Orl    100 mg

Replavite    80007498    WNP    (SA)

**A11H        OTHER PLAIN VITAMIN PREPARATIONS**

**A11HA       OTHER PLAIN VITAMIN PREPARATIONS**

A11HA03    TOCOPHEROL (VIT E)

Cap    Orl    100 IU

Vitamin E    00189227    EXZ    BEF-18G  
Vitamin E Natural    00122823    JAM    BEF-18G

Cap    Orl    200 IU

Vitamin E    00189235    EXZ    BEF-18G  
Vitamin E Natural    00122831    JAM    BEF-18G

Cap    Orl    400 IU

Vitamin E    00266108    CCM    BEF-18G  
Vitamin E    02040816    CCM    BEF-18G  
Vitamin E Natural    00122858    JAM    BEF-18G  
Vitamin E Natural    00201995    WAM    BEF-18G  
Vitamin E Synthetic    00274259    WAM    BEF-18G

Dps    Orl    50 IU

Aquasol E    02162075    CLC    BEF-18G

**A12        MINERAL SUPPLEMENTS**

**A12B       POTASSIUM**

**A12BA       POTASSIUM**

A12BA01    POTASSIUM CHLORIDE

Liq    Orl    100 mg/mL

Jamp-Potassium Chloride    80024835    JPC    ACDEFGV  
Odan Potassium Chloride    80046782    ODN    ACDEFGV  
pms-Potassium Chloride    02238604    PMS    ACDEFGV

SRC    Orl    600 mg

Micro-K    02042304    PAL    ACDEFGV  
Jamp-Potassium Chloride ER    80062704    JPC    ACDEFGV

A12BA01 POTASSIUM CHLORIDE

SRT	Orl	600 mg	Jamp-K8	80013005	JPC	ACDEFGV
			M-K8 L.A.	80035346	MRA	ACDEFGV
			Sandoz K 8	02246734	SDZ	ACDEFGV
SRT	Orl	1 500 mg	Jamp-K20	80013007	JPC	ACDEFGV
			Odan K-20	80004415	ODN	ACDEFGV
			Sandoz K 20	02242261	SDZ	ACDEFGV

A12BA02 POTASSIUM CITRATE

ERT	Orl	540 mg	Urocit-K	01914022	PAL	ACDEFGV
ERT	Orl	1 080 mg	Urocit-K	02353997	PAL	ACDEFGV
Evt	Orl	975 mg	K-Lyte	02085992	WLS	ACDEFGV
			Jamp-K Effervescent	80033602	JPC	ACDEFGV

**A12C OTHER MINERAL SUPPLEMENTS**

**A12CC MAGNESIUM**

A12CC99 MAGNESIUM GLUCOHEPTONATE

Liq	Orl	100 mg/mL	Rougier Magnesium	00026697	ROG	ACDEFGV
			Jamp Magnesium	80009357	JPC	ACDEFGV

**A12CD FLUORIDE**

A12CD01 SODIUM FLUORIDE

Dps	Orl	5.56 mg/mL	Fluor-a-Day	00610100	PDP	EF-18G
Tab	Orl	2.21 mg	Fluor-a-Day	00575569	PDP	EF-18G

**A16 OTHER ALIMENTARY TRACT AND METABOLISM PRODUCTS**

**A16A OTHER ALIMENTARY TRACT AND METABOLISM PRODUTS**

**A16AA AMINO ACIDS AND DERIVATIVES**

A16AA01 LEVOCARNITINE

Liq	Orl	100 mg/mL	Carnitor	02144336	LBI	(SA)
			Odan-Levocarnitine	02492105	ODN	(SA)
Tab	Orl	330 mg	Carnitor	02144328	LBI	(SA)

A16AA04 MERCAPTAMINE (CYSTEAMINE)

CDR	Orl	25 mg	Procysbi	02464705	HRZ	(SA)
CDR	Orl	75 mg	Procysbi	02464713	HRZ	(SA)

**A16AB ENZYMES**

A16AB07	ALGLUCOSIDASE ALFA								
Pws	IV	50 mg			Myozyme	02284863	GZM	(SA)	
A16AB10	VELAGLUCERASE ALFA								
Pws	IV	400 units			VPRIV	02357119	PAL	(SA)	
A16AB11	TALIGLUCERASE ALFA								
Pws	IV	200 units/vial			Elelyso	02425637	PFI	(SA)	
A16AB12	ELOSULFASE ALFA								
Liq	IV	5 mg / 5 mL			Vimizim	02427184	BMR	(SA)	
A16AB13	ASFOTASE ALFA								
Liq	SC	18 mg / 0.45 mL			Strensiq	02444615	ALX	(SA)	
Liq	SC	28 mg / 0.7 mL			Strensiq	02444623	ALX	(SA)	
Liq	SC	40 mg/mL			Strensiq	02444631	ALX	(SA)	
Liq	SC	80 mg / 0.8 mL			Strensiq	02444658	ALX	(SA)	
A16AB14	SEBELIPASE ALFA								
Liq	IV	2 mg/mL			Kanuma	02469596	ALX	(SA)	
A16AB17	CERLIPONASE ALFA								
Liq	IVR	150 mg / 5 mL			Brineura	02484013	BMR	(SA)	

**A16AX VARIOUS ALIMENTARY TRACT AND METABOLISM PRODUCTS**

A16AX03	SODIUM PHENYLBUTYRATE								
Gran	Orl	483 mg/g			Pheburane	02436663	MDU	(SA)	
A16AX04	NITISINONE								
Cap	Orl	2 mg			Orfadin	02459698	BVT	(SA)	
					MDK-Nitisinone	02457717	MDK	(SA)	
Cap	Orl	5 mg			Orfadin	02459701	BVT	(SA)	
					MDK-Nitisinone	02457725	MDK	(SA)	
Cap	Orl	10 mg			Orfadin	02459728	BVT	(SA)	
					MDK-Nitisinone	02457733	MDK	(SA)	

A16AX04	NITISINONE							
Cap	Orl	20 mg				Orfadin	02459736	BVT (SA)
						MDK-Nitisinone	02470055	MDK (SA)
A16AX07	SAPROPTERIN							
Pws	Orl	100 mg				Kuvan	02482207	BMR (SA)
Pws	Orl	500 mg				Kuvan	02482215	BMR (SA)
Tab	Orl	100 mg				Kuvan	02350580	BMR (SA)
A16AX08	TEDUGLUTIDE							
Pws	SC	5 mg				Revestive	02445727	TAK (SA)
A16AX09	GLYCEROL PHENYLBUTYRATE							
Liq	Orl	1.1 g/mL				Ravicti	02453304	HRZ (SA)
A16AX12	TRIENTINE							
Cap	Orl	250 mg				Mar-Trientine	02504855	MAR (SA)
						Waymade-Trientine	02515067	WMD (SA)
A16AX14	MIGALASTAT							
Cap	Orl	123 mg				Galafold	02468042	AMT (SA)

**B BLOOD AND BLOOD FORMING ORGANS**

**B01 ANTITHROMBOTIC AGENTS**

**B01A ANTITHROMBOTIC AGENTS**

**B01AA VITAMIN K ANTAGONISTS**

B01AA03	WARFARIN							
Tab	Orl	1 mg				Apo-Warfarin	02242924	APX ACDEFGV
						Taro-Warfarin	02242680	TAR ACDEFGV
Tab	Orl	2 mg				Apo-Warfarin	02242925	APX ACDEFGV
						Taro-Warfarin	02242681	TAR ACDEFGV
Tab	Orl	2.5 mg				Apo-Warfarin	02242926	APX ACDEFGV
						Taro-Warfarin	02242682	TAR ACDEFGV
Tab	Orl	3 mg				Apo-Warfarin	02245618	APX ACDEFGV
						Taro-Warfarin	02242683	TAR ACDEFGV

**B01AA03 WARFARIN**

Tab	Orl	4 mg	Apo-Warfarin	02242927	APX	ACDEFGV
			Taro-Warfarin	02242684	TAR	ACDEFGV
Tab	Orl	5 mg	Apo-Warfarin	02242928	APX	ACDEFGV
			Taro-Warfarin	02242685	TAR	ACDEFGV
Tab	Orl	6 mg	Taro-Warfarin	02242686	TAR	ACDEFGV
Tab	Orl	10 mg	Apo-Warfarin	02242929	APX	ACDEFGV
			Taro-Warfarin	02242687	TAR	ACDEFGV

**B01AB HEPARIN GROUP****B01AB01 HEPARIN**

Liq	Inj	100 IU/mL	Heparin	00727520	LEO	ACDEFGVW
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**B01AB04 DALTEPARIN**

Liq	Inj	2 500 IU / 0.2 mL	Fragmin (prefilled syringe)	02132621	PFI	W (SA)
Liq	Inj	3 500 IU / 0.28 mL	Fragmin (prefilled syringe)	02430789	PFI	W (SA)
Liq	Inj	5 000 IU / 0.2 mL	Fragmin (prefilled syringe)	02132648	PFI	W (SA)
Liq	Inj	7 500 IU / 0.3 mL	Fragmin (prefilled syringe)	02352648	PFI	W (SA)
Liq	Inj	10 000 IU / 0.4 mL	Fragmin (prefilled syringe)	02352656	PFI	W (SA)
Liq	Inj	10 000 IU/mL	Fragmin (ampoule)	02132664	PFI	W (SA)
Liq	Inj	12 500 IU / 0.5 mL	Fragmin (prefilled syringe)	02352664	PFI	W (SA)
Liq	Inj	15 000 IU / 0.6 mL	Fragmin (prefilled syringe)	02352672	PFI	W (SA)
Liq	Inj	16 500 IU / 0.66 mL	Fragmin (prefilled syringe)	02494582	PFI	W (SA)
Liq	Inj	18 000 IU / 0.72 mL	Fragmin (prefilled syringe)	02352680	PFI	W (SA)
Liq	Inj	25 000 IU/mL	Fragmin (multi-dose vial)	02231171	PFI	W (SA)

**B01AB05 ENOXAPARIN**

B01AB05		ENOXAPARIN						
Liq	Inj	30 mg / 0.3 mL		Elonox (prefilled syringe)	02532247	FKB	ACDEFGVW	
				Inclunox (prefilled syringe)	02507501	SDZ	ACDEFGVW	
				Noromby (prefilled syringe)	02506459	JNO	ACDEFGVW	
				Redesca (prefilled syringe)	02509075	VAL	ACDEFGVW	
Liq	Inj	40 mg / 0.4 mL		Elonox (prefilled syringe)	02532255	FKB	ACDEFGVW	
				Inclunox (prefilled syringe)	02507528	SDZ	ACDEFGVW	
				Noromby (prefilled syringe)	02506467	JNO	ACDEFGVW	
				Redesca (prefilled syringe)	02509083	VAL	ACDEFGVW	
Liq	Inj	60 mg / 0.6 mL		Elonox (prefilled syringe)	02532263	FKB	ACDEFGVW	
				Inclunox (prefilled syringe)	02507536	SDZ	ACDEFGVW	
				Noromby (prefilled syringe)	02506475	JNO	ACDEFGVW	
				Redesca (prefilled syringe)	02509091	VAL	ACDEFGVW	
Liq	Inj	80 mg / 0.8 mL		Elonox (prefilled syringe)	02532271	FKB	ACDEFGVW	
				Inclunox (prefilled syringe)	02507544	SDZ	ACDEFGVW	
				Noromby (prefilled syringe)	02506483	JNO	ACDEFGVW	
				Redesca (prefilled syringe)	02509105	VAL	ACDEFGVW	
Liq	Inj	100 mg/mL		Elonox (prefilled syringe)	02532298	FKB	ACDEFGVW	
				Inclunox (prefilled syringe)	02507552	SDZ	ACDEFGVW	
				Noromby (prefilled syringe)	02506491	JNO	ACDEFGVW	
				Redesca (prefilled syringe)	02509113	VAL	ACDEFGVW	
Liq	Inj	120 mg / 0.8 mL		Elonox HP (prefilled syringe)	02532301	FKB	ACDEFGVW	
				Inclunox HP (prefilled syringe)	02507560	SDZ	ACDEFGVW	
				Noromby HP (prefilled syringe)	02506505	JNO	ACDEFGVW	
				Redesca HP (prefilled syringe)	02509148	VAL	ACDEFGVW	
Liq	Inj	150 mg/mL		Elonox HP (prefilled syringe)	02532328	FKB	ACDEFGVW	
				Inclunox HP (prefilled syringe)	02507579	SDZ	ACDEFGVW	
				Noromby HP (prefilled syringe)	02506513	JNO	ACDEFGVW	
				Redesca HP (prefilled syringe)	02509156	VAL	ACDEFGVW	
Liq	Inj	300 mg / 3 mL		Redesca (multi-dose vial)	02509121	VAL	ACDEFGVW	
B01AB06		NADROPARIN						
Liq	Inj	2 850 IU / 0.3 mL		Fraxiparin (prefilled syringe)	02236913	APN	W (SA)	
Liq	Inj	3 800 IU / 0.4 mL		Fraxiparin (prefilled syringe)	02450623	APN	W (SA)	

**B01AB06 NADROPARIN**

Liq	Inj	5 700 IU / 0.6 mL	Fraxiparin (prefilled syringe)	02450631	APN	W (SA)
Liq	Inj	9 500 IU/mL	Fraxiparin (prefilled syringe)	02450658	APN	W (SA)
Liq	Inj	11 400 IU / 0.6 mL	Fraxiparin Forte (prefilled syringe)	02450674	APN	W (SA)
Liq	Inj	15 200 IU / 0.8 mL	Fraxiparin Forte (prefilled syringe)	02450666	APN	W (SA)
Liq	Inj	19 000 IU/mL	Fraxiparin Forte (prefilled syringe)	02240114	APN	W (SA)

**B01AB10 TINZAPARIN**

Liq	Inj	2 500 IU / 0.25 mL	Innohep (prefilled syringe)	02229755	LEO	W (SA)
Liq	Inj	3 500 IU / 0.35 mL	Innohep (prefilled syringe)	02358158	LEO	W (SA)
Liq	Inj	4 500 IU / 0.45 mL	Innohep (prefilled syringe)	02358166	LEO	W (SA)
Liq	Inj	8 000 IU / 0.4 mL	Innohep (prefilled syringe)	02429462	LEO	W (SA)
Liq	Inj	10 000 IU / 0.5 mL	Innohep (prefilled syringe)	02231478	LEO	W (SA)
Liq	Inj	12 000 IU / 0.6 mL	Innohep (prefilled syringe)	02429470	LEO	W (SA)
Liq	Inj	14 000 IU / 0.7 mL	Innohep (prefilled syringe)	02358174	LEO	W (SA)
Liq	Inj	16 000 IU / 0.8 mL	Innohep (prefilled syringe)	02429489	LEO	W (SA)
Liq	Inj	18 000 IU / 0.9 mL	Innohep (prefilled syringe)	02358182	LEO	W (SA)
Liq	Inj	20 000 IU/2 mL	Innohep (multi-dose vial)	02167840	LEO	W (SA)
Liq	Inj	40 000 IU / 2 mL	Innohep (multi-dose vial)	02229515	LEO	W (SA)

**B01AC PLATELET AGGREGATION INHIBITORS EXCLUDING HEPARIN****B01AC04 CLOPIDOGREL**



## B01AC04 CLOPIDOGREL

Tab Orl 75 mg

Plavix	02238682	SAV	ACDEFV
Apo-Clopidogrel	02252767	APX	ACDEFV
Auro-Clopidogrel	02416387	ARO	ACDEFV
Clopidogrel	02394820	PDL	ACDEFV
Clopidogrel	02400553	SAS	ACDEFV
Clopidogrel	02385813	SIV	ACDEFV
Jamp-Clopidogrel	02415550	JPC	ACDEFV
M-Clopidogrel	02502283	MRA	ACDEFV
Mar-Clopidogrel	02422255	MAR	ACDEFV
Mint-Clopidogrel	02408910	MNT	ACDEFV
NRA-Clopidogrel	02482037	NRA	ACDEFV
pms-Clopidogrel	02348004	PMS	ACDEFV
Taro-Clopidogrel	02379813	SUN	ACDEFV
Teva-Clopidogrel	02293161	TEV	ACDEFV

## B01AC05 TICLOPIDINE

Tab Orl 250 mg

Ticlopidine 02237701 AAP ACDEFV

## B01AC09 EPOPROSTENOL

Pws IV 0.5 mg

Caripul 02397447 JAN (SA)

Flolan 02230845 GSK (SA)

Pws IV 1.5 mg

Caripul 02397455 JAN (SA)

Flolan 02230848 GSK (SA)

## B01AC21 TREPROSTINIL

Liq SC 1 mg/mL

Remodulin 02246552 UTC (SA)

Liq SC 2.5 mg/mL

Remodulin 02246553 UTC (SA)

Liq SC 5 mg/mL

Remodulin 02246554 UTC (SA)

Liq SC 10 mg/mL

Remodulin 02246555 UTC (SA)

## B01AC22 PRASUGREL

Tab Orl 10 mg

Jamp Prasugrel 02502429 JPC (SA)

## B01AC24 TICAGRELOR

Tab Orl 60 mg

Apo-Ticagrelor 02482622 APX (SA)

M-Ticagrelor 02529750 MRA (SA)

Taro-Ticagrelor 02492571 TAR (SA)

**B01AC24 TICAGRELOR**

Tab Orl 90 mg

Brilinta 02368544 AZE (SA)

Apo-Ticagrelor 02482630 APX (SA)

M-Ticagrelor 02529769 MRA (SA)

Taro-Ticagrelor 02492598 TAR (SA)

**B01AC27 SELEXIPAG**

Tab Orl 200 mcg

Uptravi 02451158 JAN (SA)

Tab Orl 400 mcg

Uptravi 02451166 JAN (SA)

Tab Orl 600 mcg

Uptravi 02451174 JAN (SA)

Tab Orl 800 mcg

Uptravi 02451182 JAN (SA)

Tab Orl 1 000 mcg

Uptravi 02451190 JAN (SA)

Tab Orl 1 200 mcg

Uptravi 02451204 JAN (SA)

Tab Orl 1 400 mcg

Uptravi 02451212 JAN (SA)

Tab Orl 1 600 mcg

Uptravi 02451220 JAN (SA)

**B01AC30 COMBINATIONS**

DIPYRIDAMOLE / ACETYLSALICYLIC ACID

Cap Orl 200 mg / 25 mg

Taro-Dipyridamole/ASA 02471051 TAR (SA)

**B01AD ENZYMES****B01AD02 ALTEPLASE**

Pws Isl 2 mg

Cathflo 02245859 HLR (SA)

**B01AE DIRECT THROMBIN INHIBITORS****B01AE07 DABIGATRAN ETEXILATE**

Cap Orl 110 mg

Pradaxa 02312441 BOE (SA)

Apo-Dabigatran 02468905 APX (SA)

Cap Orl 150 mg

Pradaxa 02358808 BOE (SA)

Apo-Dabigatran 02468913 APX (SA)

**B01AF DIRECT FACTOR XA INHIBITORS****B01AF01 RIVAROXABAN**

Tab Orl 2.5 mg

Xarelto 02480808 BAY (SA)

B01AF01	RIVAROXABAN					
Tab	Orl	10 mg	Xarelto	02316986	BAY	(SA)
Tab	Orl	15 mg	Xarelto	02378604	BAY	ACDEFGV
Tab	Orl	20 mg	Xarelto	02378612	BAY	ACDEFGV
B01AF02	APIXABAN					
Tab	Orl	2.5 mg	Eliquis	02377233	BRI	ACDEFGV
			ACH-Apixaban	02487713	AHI	ACDEFGV
			Apixaban	02530708	SIV	ACDEFGV
			Apo-Apixaban	02487381	APX	ACDEFGV
			Auro-Apixaban	02486806	ARO	ACDEFGV
			Jamp Apixaban	02528924	JPC	ACDEFGV
			M-Apixaban	02529009	MRA	ACDEFGV
			Mar-Apixaban	02492369	MAR	ACDEFGV
			Mint-Apixaban	02495430	MNT	ACDEFGV
			Nat-Apixaban	02492814	NAT	ACDEFGV
			Sandoz Apixaban	02489228	SDZ	ACDEFGV
			Taro-Apixaban	02510464	SUN	ACDEFGV
Tab	Orl	5 mg	Eliquis	02397714	BRI	ACDEFGV
			ACH-Apixaban	02487721	AHI	ACDEFGV
			Apixaban	02530716	SIV	ACDEFGV
			Apo-Apixaban	02487403	APX	ACDEFGV
			Auro-Apixaban	02486814	ARO	ACDEFGV
			Jamp Apixaban	02528932	JPC	ACDEFGV
			M-Apixaban	02529017	MRA	ACDEFGV
			Mar-Apixaban	02492377	MAR	ACDEFGV
			Mint-Apixaban	02495449	MNT	ACDEFGV
			Nat-Apixaban	02492822	NAT	ACDEFGV
			Sandoz Apixaban	02489236	SDZ	ACDEFGV
			Taro-Apixaban	02510472	SUN	ACDEFGV
B01AF03	EDOXABAN					
Tab	Orl	15 mg	Lixiana	02458640	SEV	ACDEFGV
Tab	Orl	30 mg	Lixiana	02458659	SEV	ACDEFGV
Tab	Orl	60 mg	Lixiana	02458667	SEV	ACDEFGV

**B02 ANTIHAEMORRHAGICS****B02A ANTIFIBRINOLYTICS****B02AA AMINO ACIDS**

## B02AA02 TRANEXAMIC ACID

Tab Orl 500 mg

Cyklokapron 02064405 PFI ACDEFGV

GD-Tranexamic Acid 02409097 GMD ACDEFGV

Mar-Tranexamic Acid 02496232 MAR ACDEFGV

Tranexamic Acid 02519194 JPC ACDEFGV

Tranexamic Acid 02401231 STR ACDEFGV

**B02B VITAMIN K AND OTHER HEMOSTATICS****B02BA VITAMIN K**

## B02BA01 PHYTOMENADIONE

Liq IM 1 mg / 0.5 mL

Vitamin K 00781878 SDZ ACDEFGVW

Liq IM 10 mg/mL

Vitamin K 00804312 SDZ ACDEFGVW

**B03 ANTIANAEMIC PREPARATIONS****B03A IRON PREPARATIONS****B03AA IRON BIVALENT, ORAL PREPARATIONS**

## B03AA02 FERROUS FUMARATE

Cap Orl 300 mg

Palafer 01923420 BSH AEEFGV

Jamp-Fer 80024232 JPC AEEFGV

Sandoz-Fer 02237556 SDZ AEEFGV

Sus Orl 60 mg/mL

Palafer 01923439 BSH AEEFGV

Tab Orl 300 mg

Ferrous Fumarate 00031089 WAM AEEFGV

## B03AA03 FERROUS GLUCONATE

Tab Orl 300 mg

Ferrous Gluconate 00031097 JPC AEEFGV

Ferrous Gluconate 00582727 VTH AEEFGV

Novo-Ferrogluc 80000435 NUT AEEFGV

## B03AA07 FERROUS SULPHATE

Dps Orl 125 mg/mL

pms-Ferrous Sulfate 00816035 PMS AEEFGV

Liq Orl 75 mg/mL

Fer-In-Sol 00762954 MJO AEEFGV

Ferodan 02237385 ODN AEEFGV

Jamp Ferrous Sulfate 80008309 JPC AEEFGV

Liq Orl 150 mg / 5 mL

Jamp-Ferrous Sulfate 80008295 JPC AEEFGV

**B03AA07 FERROUS SULPHATE**

Syr Orl 150 mg / 5 mL

Fer-In-Sol 00017884 MJO AEFVG

Ferodan 00758469 ODN AEFVG

Tab Orl 300 mg

Ferrous Sulfate 00031100 JPC AEFVG

Ferrous Sulfate SC 00346918 CCM AEFVG

pms-Ferrous Sulfate 00586323 PMS AEFVG

**B03AC IRON TRIVALENT, PARENTERAL PREPARATIONS****B03AC02 SACCHARATED IRON OXIDE**

## IRON SUCROSE

Liq IV 20 mg/mL

Venofer 02243716 FRE ACDEFGV

pms-Iron Sucrose 02502917 PMS ACDEFGV

**B03AC07 FERRIC SODIUM GLUCONATE COMPLEX**

Liq Inj 12.5 mg/mL

Ferrelecit 02243333 SAV (SA)

**B03AC99 FERRIC DERISOMALTOSE**

Liq Inj 100 mg/mL

Monoferric 02477777 PFI (SA)

**B03B VITAMIN B12 AND FOLIC ACID****B03BA VITAMIN B12 (CYANOCOBALAMIN AND DERIVATIVES)****B03BA01 CYANOCOBALAMIN**

Liq Inj 1 000 mcg/mL

Cyanocobalamin 01987003 STR ACDEFGV

Vitamin B12 00521515 SDZ ACDEFGV

**B03BB FOLIC ACID AND DERIVATIVES****B03BB01 FOLIC ACID**

Tab Orl 5 mg

Jamp-Folic 02366061 JPC ACDEFGV

Sandoz-Folic 02285673 SDZ ACDEFGV

**B03X OTHER ANTIANEMIC PREPARATIONS****B03XA OTHER ANTIANEMIC PREPARATIONS****B03XA01 ERYTHROPOIETIN (EPOETIN ALFA)**

Liq Inj 1 000 IU / 0.5 mL

Eprex 02231583 JAN W (SA)

Liq Inj 2 000 IU / 0.5 mL

Eprex 02231584 JAN W (SA)

Liq Inj 3 000 IU / 0.3 mL

Eprex 02231585 JAN W (SA)

Liq Inj 4 000 IU / 0.4 mL

Eprex 02231586 JAN W (SA)

B03XA01 ERYTHROPOIETIN (EPOETIN ALFA)

Liq	Inj	5 000 IU / 0.5 mL	Eprex	02243400	JAN	W (SA)
Liq	Inj	6 000 IU / 0.6 mL	Eprex	02243401	JAN	W (SA)
Liq	Inj	8 000 IU / 0.8 mL	Eprex	02243403	JAN	W (SA)
Liq	Inj	10 000 IU/mL	Eprex	02231587	JAN	W (SA)
Liq	Inj	20 000 IU / 0.5 mL	Eprex	02243239	JAN	W (SA)
Liq	Inj	30 000 IU / 0.75 mL	Eprex	02288680	JAN	W (SA)
Liq	Inj	40 000 IU/mL	Eprex	02240722	JAN	W (SA)

B03XA02 DARBEPOETIN ALFA

Liq	Inj	10 mcg / 0.4 mL	Aranesp	02392313	AGA	W (SA)
Liq	Inj	20 mcg / 0.5 mL	Aranesp	02392321	AGA	W (SA)
Liq	Inj	30 mcg / 0.3 mL	Aranesp	02392348	AGA	W (SA)
Liq	Inj	40 mcg / 0.4 mL	Aranesp	02391740	AGA	W (SA)
Liq	Inj	50 mcg / 0.5 mL	Aranesp	02391759	AGA	W (SA)
Liq	Inj	60 mcg / 0.3 mL	Aranesp	02392356	AGA	W (SA)
Liq	Inj	80 mcg / 0.4 mL	Aranesp	02391767	AGA	W (SA)
Liq	Inj	100 mcg / 0.5 mL	Aranesp	02391775	AGA	W (SA)
Liq	Inj	130 mcg / 0.65 mL	Aranesp	02391783	AGA	W (SA)
Liq	Inj	150 mcg / 0.3 mL	Aranesp	02391791	AGA	W (SA)
Liq	Inj	200 mcg / 0.4 mL	Aranesp	02391805	AGA	W (SA)
Liq	Inj	300 mcg / 0.6 mL	Aranesp	02391821	AGA	W (SA)
Liq	Inj	500 mcg / 1 mL	Aranesp	02392364	AGA	W (SA)

B03XA06 LUSPATERCEPT

Pws	SC	25 mg	Reblozyl	02505541	CEL	(SA)
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B03XA06 LUSPATERCEPT

Pws SC 75 mg

Reblozyl 02505568 CEL (SA)

**B06 OTHER HEMATOLOGICAL AGENTS**

**B06A OTHER HEMATOLOGICAL AGENTS**

**B06AC DRUGS USED IN HEREDITARY ANGIOEDEMA**

B06AC02 ICATIBANT

Liq SC 30 mg / 3 mL

Firazyr 02425696 TAK (SA)

B06AC05 LANADELUMAB

Liq SC 300 mg / 2 mL

Takhzyro 02480948 SHI (SA)

Takhzyro 02505614 SHI (SA)

**C CARDIOVASCULAR SYSTEM**

**C01 CARDIAC THERAPY**

**C01A CARDIAC GLYCOSIDES**

**C01AA DIGITALIS GLYCOSIDES**

C01AA05 DIGOXIN

Liq Orl 0.05 mg/mL

pms-Digoxin 02242320 PMS ACDEFGV

Tab Orl 0.0625 mg

Jamp-Digoxin 02498502 JPC ACDEFGV

pms-Digoxin 02335700 PMS ACDEFGV

Tab Orl 0.125 mg

Jamp-Digoxin 02498510 JPC ACDEFGV

pms-Digoxin 02335719 PMS ACDEFGV

**C01B ANTIARRHYTHMICS, CLASS I AND III**

**C01BA ANTIARRHYTHMICS, CLASS IA**

C01BA03 DISOPYRAMIDE

Cap Orl 100 mg

Rythmodan 02224801 XPI ACDEFGV

**C01BB ANTIARRHYTHMICS, CLASS IB**

C01BB02 MEXILETINE

Cap Orl 100 mg

Teva-Mexiletine 02230359 TEV ACDEFGV

Cap Orl 200 mg

Teva-Mexiletine 02230360 TEV ACDEFGV

**C01BC ANTIARRHYTHMICS, CLASS IC**

C01BC03 PROPAFENONE

C01BC03 PROPAFENONE

Tab Orl 150 mg

Rythmol 00603708 BGP ACDEFGV  
Apo-Propafenone 02243324 APX ACDEFGV  
Mylan-Propafenone 02457172 MYL ACDEFGV  
Propafenone 02343053 SAS ACDEFGV

Tab Orl 300 mg

Rythmol 00603716 BGP ACDEFGV  
Apo-Propafenone 02243325 APX ACDEFGV  
Mylan-Propafenone 02457164 MYL ACDEFGV  
Propafenone 02343061 SAS ACDEFGV

C01BC04 FLECAINIDE

Tab Orl 50 mg

Apo-Flecainide 02275538 APX ACDEFGV  
Auro-Flecainide 02459957 ARO ACDEFGV  
Jamp-Flecainide 02493705 JPC ACDEFGV  
Mar-Flecainide 02476177 MAR ACDEFGV

Tab Orl 100 mg

Apo-Flecainide 02275546 APX ACDEFGV  
Auro-Flecainide 02459965 ARO ACDEFGV  
Jamp-Flecainide 02493713 JPC ACDEFGV  
Mar-Flecainide 02476185 MAR ACDEFGV

**C01BD ANTIARRHYTHMICS, CLASS III**

C01BD01 AMIODARONE

Tab Orl 100 mg

pms-Amiodarone 02292173 PMS ACDEFGV

Tab Orl 200 mg

Amiodarone 02364336 SAS ACDEFGV  
Amiodarone 02385465 SIV ACDEFGV  
Apo-Amiodarone 02246194 APX ACDEFGV  
Jamp Amiodarone 02531844 JPC ACDEFGV  
pms-Amiodarone 02242472 PMS ACDEFGV  
Sandoz Amiodarone 02243836 SDZ ACDEFGV  
Teva-Amiodarone 02239835 TEV ACDEFGV

**C01C CARDIAC STIMULANTS EXCLUDING CARDIAC GLYCOSIDES**

**C01CA ADRENERGIC AND DOPAMINERGIC AGENTS**

C01CA17 MIDODRINE

Tab Orl 2.5 mg

Apo-Midodrine 02278677 APX ACDEFGV  
Jamp Midodrine 02517701 JPC ACDEFGV  
Mar-Midodrine 02473984 MAR ACDEFGV  
Midodrine 02533200 SAS ACDEFGV



**C01CA17 MIDODRINE**

Tab Orl 5 mg

Apo-Midodrine	02278685	APX	ACDEFGV
Jamp Midodrine	02517728	JPC	ACDEFGV
Mar-Midodrine	02473992	MAR	ACDEFGV
Midodrine	02533219	SAS	ACDEFGV

**C01CA24 EPINEPHRINE**

Liq Inj 0.15 mg

Allerject	02382059	KLO	ACDEFGV
EpiPen Jr	00578657	PFI	ACDEFGV

Liq Inj 0.3 mg

Allerject	02382067	KLO	ACDEFGV
Emerade	02458446	BSL	ACDEFGV
EpiPen	00509558	PFI	ACDEFGV

Liq Inj 0.5 mg

Emerade	02458454	BSL	ACDEFGV
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Liq Inj 1 mg/mL

Adrenalin	00155357	ERF	ACDEFGV
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**C01D VASODILATORS USED IN CARDIAC DISEASES****C01DA ORGANIC NITRATES****C01DA02 NITROGLYCERIN (GLYCERYL TRINITRATE)**

Aem Slg 0.4 mg

Nitrolingual	02231441	SAV	ACDEFGV
Mylan-Nitro SL	02243588	MYL	ACDEFGV
Rho-Nitro	02238998	SDZ	ACDEFGV

Pth Trd 0.2 mg/hr

Nitro-Dur	01911910	RCH	ACDEFV
Trinipatch	02230732	PAL	ACDEFV
Mylan-Nitro Patch	02407442	MYL	ACDEFV

Pth Trd 0.4 mg/hr

Nitro-Dur	01911902	RCH	ACDEFV
Trinipatch	02230733	PAL	ACDEFV
Mylan-Nitro Patch	02407450	MYL	ACDEFV

Pth Trd 0.6 mg/hr

Nitro-Dur	01911929	RCH	ACDEFV
Trinipatch	02230734	PAL	ACDEFV
Mylan-Nitro Patch	02407469	MYL	ACDEFV

Pth Trd 0.8 mg/hr

Nitro-Dur	02011271	RCH	ACDEFV
Mylan-Nitro Patch	02407477	MYL	ACDEFV

Slit Slg 0.3 mg

Nitrostat	00037613	UJC	ACDEFGV
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C01DA02	NITROGLYCERIN (GLYCERYL TRINITRATE)						
Slit	Slg	0.6 mg		Nitrostat	00037621	UJC	ACDEFGV
C01DA08	ISOSORBIDE DINITRATE						
Tab	Orl	10 mg		ISDN	00441686	AAP	ACDEFGV
Tab	Orl	30 mg		ISDN	00441694	AAP	ACDEFGV
C01DA14	ISOSORBIDE MONONITRATE						
SRT	Orl	60 mg		Imdur	02126559	JNO	ACDEFGV
				Apo-ISMN	02272830	APX	ACDEFGV
				pms-ISMN	02301288	PMS	ACDEFGV

**C01E OTHER CARDIAC PREPARATIONS**

**C01EB OTHER CARDIAC PRODUCTS**

C01EB17	IVABRADINE						
Tab	Orl	5 mg		Lancora	02459973	SEV	(SA)
Tab	Orl	7.5 mg		Lancora	02459981	SEV	(SA)

**C02 ANTIHYPERTENSIVES**

**C02A ANTIADRENERGIC AGENTS, CENTRALLY ACTING**

**C02AB METHYLDOPA**

C02AB02	METHYLDOPA (RACEMIC)						
Tab	Orl	125 mg		Methyldopa	00360252	AAP	ACDEFGV
Tab	Orl	250 mg		Methyldopa	00360260	AAP	ACDEFGV
Tab	Orl	500 mg		Methyldopa	00426830	AAP	ACDEFGV

**C02AC IMIDAZOLINE RECEPTOR AGONISTS**

C02AC01	CLONIDINE						
Tab	Orl	0.025 mg		Jamp Clonidine	02528207	JPC	ACDEFGV
				Mar-Clonidine	02524198	MAR	ACDEFGV
				Mint-Clonidine	02534738	MNT	ACDEFGV
				Sandoz Clonidine	02516217	SDZ	ACDEFGV
				Teva-Clonidine	02304163	TEV	ACDEFGV
Tab	Orl	0.1 mg		Mint-Clonidine	02462192	MNT	ACDEFGV
				Sandoz Clonidine	02515784	SDZ	ACDEFGV
				Teva-Clonidine	02046121	TEV	ACDEFGV

C02AC01 CLONIDINE

Tab Orl 0.2 mg

Mint-Clonidine 02462206 MNT ACDEFGV  
Sandoz Clonidine 02515792 SDZ ACDEFGV  
Teva-Clonidine 02046148 TEV ACDEFGV

**C02C ANTIADRENERGIC AGENTS, PERIPHERALLY ACTING**

**C02CA ALPHA-ADRENOCEPTOR ANTAGONISTS**

C02CA01 PRAZOSIN

Tab Orl 1 mg

Teva-Prazin 01934198 TEV ACDEFGV

Tab Orl 2 mg

Teva-Prazin 01934201 TEV ACDEFGV

Tab Orl 5 mg

Teva-Prazin 01934228 TEV ACDEFGV

C02CA04 DOXAZOSIN

Tab Orl 1 mg

Apo-Doxazosin 02240588 APX ACDEFGV  
Jamp-Doxazosin 02489937 JPC ACDEFGV  
Teva-Doxazosin 02242728 TEV ACDEFGV

Tab Orl 2 mg

Apo-Doxazosin 02240589 APX ACDEFGV  
Jamp-Doxazosin 02489945 JPC ACDEFGV  
Teva-Doxazosin 02242729 TEV ACDEFGV

Tab Orl 4 mg

Apo-Doxazosin 02240590 APX ACDEFGV  
Jamp-Doxazosin 02489953 JPC ACDEFGV  
Teva-Doxazosin 02242730 TEV ACDEFGV

**C02D ARTERIOLAR SMOOTH MUSCLE, AGENTS ACTING ON**

**C02DB HYDRAZINOPHTHALAZINE DERIVATIVES**

C02DB02 HYDRALAZINE

Tab Orl 10 mg

Apo-Hydralazine 00441619 APX ACDEFGV  
Jamp-Hydralazine 02457865 JPC ACDEFGV  
Mint-Hydralazine 02468778 MNT ACDEFGV

Tab Orl 25 mg

Apo-Hydralazine 00441627 APX ACDEFGV  
Jamp-Hydralazine 02457873 JPC ACDEFGV  
Mint-Hydralazine 02468786 MNT ACDEFGV

Tab Orl 50 mg

Apo-Hydralazine 00441635 APX ACDEFGV  
Jamp-Hydralazine 02457881 JPC ACDEFGV  
Mint-Hydralazine 02468794 MNT ACDEFGV

**C02DC PYRIMIDINE DERIVATIVES**

C02DC01 MINOXIDIL

Tab Orl 2.5 mg

Loniten 00514497 PFI ACDEFGV

Tab Orl 10 mg

Loniten 00514500 PFI ACDEFGV

**C02K OTHER ANTIHYPERTENSIVES****C02KX ANTIHYPERTENSIVES FOR PULMONARY ARTERIAL HYPERTENSION**

C02KX01 BOSENTAN

Tab Orl 62.5 mg

Tracleer 02244981 JAN (SA)

Nat-Bosentan 02467984 NAT (SA)

pms-Bosentan 02383012 PMS (SA)

Taro-Bosentan 02483130 TAR (SA)

Tab Orl 125 mg

Tracleer 02244982 JAN (SA)

Nat-Bosentan 02467992 NAT (SA)

pms-Bosentan 02383020 PMS (SA)

Taro-Bosentan 02483149 TAR (SA)

C02KX02 AMBRISENTAN

Tab Orl 5 mg

Volibris 02307065 GSK (SA)

Apo-Ambrisentan 02475375 APX (SA)

Jamp Ambrisentan 02521938 JPC (SA)

Sandoz Ambrisentan 02526875 SDZ (SA)

Tab Orl 10 mg

Volibris 02307073 GSK (SA)

Apo-Ambrisentan 02475383 APX (SA)

Jamp Ambrisentan 02521946 JPC (SA)

Sandoz Ambrisentan 02526883 SDZ (SA)

C02KX04 MACITENTAN

Tab Orl 10 mg

Opsumit 02415690 JAN (SA)

C02KX05 RIOCIGUAT

Tab Orl 0.5 mg

Adempas 02412764 BAY (SA)

Tab Orl 1 mg

Adempas 02412772 BAY (SA)

Tab Orl 1.5 mg

Adempas 02412799 BAY (SA)

Tab Orl 2 mg

Adempas 02412802 BAY (SA)

C02KX05	RIOCIGUAT								
Tab	Orl	2.5 mg				Adempas	02412810	BAY	(SA)

C02KX99	SILDENAFIL								
Tab	Orl	20 mg				Revatio	02279401	UJC	(SA)
						Jamp Sildenafil R	02469669	JPC	(SA)
						pms-Sildenafil R	02412179	PMS	(SA)
						Teva-Sildenafil R	02319500	TEV	(SA)

**C03 DIURETICS**

**C03A LOW-CEILING DIURETICS, THIAZIDES**

**C03AA THIAZIDES, PLAIN**

C03AA03	HYDROCHLOROTHIAZIDE								
Tab	Orl	12.5 mg				Apo-Hydro	02327856	APX	ACDEFGV
						Mint-Hydrochlorothiazide	02425947	MNT	ACDEFGV
						pms-Hydrochlorothiazide	02274086	PMS	ACDEFGV
Tab	Orl	25 mg				Apo-Hydro	00326844	APX	ACDEFGV
						Hydrochlorothiazide	02360594	SAS	ACDEFGV
						Mint-Hydrochlorothiazide	02426196	MNT	ACDEFGV
						pms-Hydrochlorothiazide	02247386	PMS	ACDEFGV
						Teva-Hydrochlorothiazide	00021474	TEV	ACDEFGV
Tab	Orl	50 mg				Apo-Hydro	00312800	APX	ACDEFGV
						Hydrochlorothiazide	02360608	SAS	ACDEFGV
						pms-Hydrochlorothiazide	02247387	PMS	ACDEFGV
						Teva-Hydrazide	00021482	TEV	ACDEFGV

**C03B LOW-CEILING DIURETICS, EXCLUDING THIAZIDES**

**C03BA SULFONAMIDES, PLAIN**

C03BA04	CHLORTHALIDONE								
Tab	Orl	50 mg				Chlorthalidone	00360279	AAP	ACDEFGV
						Jamp Chlorthalidone	02523817	JPC	ACDEFGV
C03BA08	METOLAZONE								
Tab	Orl	2.5 mg				Zaroxolyn	00888400	SAV	ACDEFGV
C03BA11	INDAPAMIDE								
Tab	Orl	1.25 mg				Apo-Indapamide	02245246	APX	ACDEFGV
						Mylan-Indapamide	02240067	MYL	ACDEFGV

C03BA11 INDAPAMIDE

Tab Orl 2.5 mg

Apo-Indapamide 02223678 APX ACDEFGV

Mylan-Indapamide 02153483 MYL ACDEFGV

**C03C HIGH-CEILING DIURETICS**

**C03CA SULFONAMIDES, PLAIN**

C03CA01 FUROSEMIDE

Liq Inj 10 mg/mL

Furosemide 00527033 SDZ ACDEFGVW

Furosemide 02382539 SDZ ACDEFGVW

Furosemide Injection USP 02461404 STR ACDEFGVW

Liq Orl 10 mg/mL

Lasix 02224720 SAV ACDEFGVW

Tab Orl 20 mg

Apo-Furosemide 00396788 APX ACDEFGVW

Furosemide 02351420 SAS ACDEFGVW

Mint-Furosemide 02466759 MNT ACDEFGVW

Teva-Furosemide 00337730 TEV ACDEFGVW

Tab Orl 40 mg

Apo-Furosemide 00362166 APX ACDEFGVW

Furosemide 02351439 SAS ACDEFGVW

Mint-Furosemide 02466767 MNT ACDEFGVW

Teva-Furosemide 00337749 TEV ACDEFGVW

Tab Orl 80 mg

Apo-Furosemide 00707570 APX ACDEFGVW

Furosemide 02351447 SAS ACDEFGVW

Mint-Furosemide 02466775 MNT ACDEFGVW

Teva-Furosemide 00765953 TEV ACDEFGVW

Tab Orl 500 mg

Lasix Special 02224755 SAV ACDEFGVW

C03CA02 BUMETANIDE

Tab Orl 1 mg

Burinex 00728284 KNI ACDEFV

Tab Orl 5 mg

Burinex 00728276 KNI ACDEFV

**C03CC ARYLOXYACETIC ACID DERIVATIVES**

C03CC01 ETHACRYNIC ACID

Tab Orl 25 mg

Edecrin 02258528 BSL ACDEFGV

**C03D POTASSIUM-SPARING DRUGS**

**C03DA ALDOSTERONE ANTAGONISTS**

C03DA01 SPIRONOLACTONE

**C03DA01 SPIRONOLACTONE**

Tab Orl 25 mg

Aldactone	00028606	PFI	ACDEFGV
Jamp Spironolactone	02518821	JPC	ACDEFGV
Mint-Spironolactone	02488140	MNT	ACDEFGV
Teva-Spironolactone	00613215	TEV	ACDEFGV

Tab Orl 100 mg

Aldactone	00285455	PFI	ACDEFGV
Jamp Spironolactone	02518848	JPC	ACDEFGV
Mint-Spironolactone	02488159	MNT	ACDEFGV
Teva-Spiroton	00613223	TEV	ACDEFGV

**C03DA04 EPLERENONE**

Tab Orl 25 mg

Inspra	02323052	BGP	(SA)
Mint-Eplerenone	02471442	MNT	(SA)

Tab Orl 50 mg

Inspra	02323060	BGP	(SA)
Mint-Eplerenone	02471450	MNT	(SA)

**C03DB OTHER POTASSIUM-SPARING AGENTS****C03DB01 AMILORIDE**

Tab Orl 5 mg

Midamor 02249510 AAP ACDEFGV

**C03E DIURETICS AND POTASSIUM-SPARING AGENTS IN COMBINATION****C03EA LOW-CEILING DIURETICS AND POTASSIUM-SPARING AGENTS****C03EA01 HYDROCHLOROTHIAZIDE AND POTASSIUM-SPARING DRUGS**

## HYDROCHLOROTHIAZIDE / AMILORIDE

Tab Orl 50 mg / 5 mg

AA-Amilzide 00784400 AAP ACDEFGV

## HYDROCHLOROTHIAZIDE / SPIRONOLACTONE

Tab Orl 25 mg / 25 mg

Teva-Spironolactone HCTZ 00613231 TEV ACDEFGV

Tab Orl 50 mg / 50 mg

Teva-Spironolactone HCTZ 00657182 TEV ACDEFGV

## HYDROCHLOROTHIAZIDE / TRIAMTERENE

Tab Orl 25 mg / 50 mg

Apo-Triazide	00441775	APX	ACDEFGV
Teva-Triamterene/HCTZ	00532657	TEV	ACDEFGV

**C04 PERIPHERAL VASODILATORS****C04A PERIPHERAL VASODILATORS****C04AA 2-AMINO-1-PHENYLETHANOL DERIVATIVES****C04AA02 BUPHENINE (NYLIDRIN)**

Tab Orl 6 mg

Arlidin 01926713 SLP ACDEFGV

**C04AD PURINE DERIVATIVES**

C04AD03 PENTOXIFYLLINE

SRT Orl 400 mg

Pentoxifylline SR 02230090 AAP ACDEFGV

**C05 VASOPROTECTIVES****C05A AGENTS FOR TREATMENT OF HEMORRHOIDS & ANAL FISSURES FOR TOPICAL USE****C05AA CORTICOSTEROIDS**

C05AA01 HYDROCORTISONE

HYDROCORTISONE / CINCHOCAINE / FRAMYCETIN / ESCULIN

Ont Rt 5 mg / 5 mg / 10 mg /  
10 mg

Proctosedyl 02223252 AXC ACDEFGV

Proctol Ointment 02247322 ODN ACDEFGV

Sup Rt 5 mg / 5 mg / 10 mg /  
10 mg

Proctol Suppositories 02247882 ODN ACDEFGV

HYDROCORTISONE / PRAMOXINE

Aer Rt 1% / 1%

Proctofoam HC 00363014 DUI ACDEFGV

HYDROCORTISONE / ZINC

Ont Rt 0.5% / 0.5%

Anodan HC 02128446 ODN ACDEFGV

Jamp-Zinc-HC 02387239 JPC ACDEFGV

Sup Rt 0.5% / 0.5%

Anodan HC 02236399 ODN ACDEFGV

**C05B ANTIVARICOSE THERAPY****C05BA HEPARINS OR HEPARINOIDS FOR TOPICAL USE**

C05BA04 PENTOSAN POLYSULFATE SODIUM

Cap Orl 100 mg

Elmiron 02029448 JAN ACDEFGV

**C07 BETA BLOCKING AGENTS****C07A BETA BLOCKING AGENTS, PLAIN****C07AA BETA BLOCKING AGENTS, NON-SELECTIVE**

C07AA03 PINDOLOL

Tab Orl 5 mg

Visken 00417270 XPI ACDEFGV

Apo-Pindol 00755877 APX ACDEFGV

Teva-Pindolol 00869007 TEV ACDEFGV

Tab Orl 10 mg

Visken 00443174 XPI ACDEFGV

Apo-Pindol 00755885 APX ACDEFGV

Teva-Pindolol 00869015 TEV ACDEFGV



C07AA03	PINDOLOL							
Tab	Orl	15 mg				Apo-Pindol	00755893	APX ACDEFGV
						Teva-Pindolol	00869023	TEV ACDEFGV
C07AA05	PROPRANOLOL							
Liq	Orl	3.75 mg/mL				Hemangiol	02457857	PFB (SA)
Tab	Orl	10 mg				Teva-Propranolol	00496480	TEV ACDEFGV
Tab	Orl	20 mg				Teva-Propranolol	00740675	TEV ACDEFGV
Tab	Orl	40 mg				Teva-Propranolol	00496499	TEV ACDEFGV
Tab	Orl	80 mg				Teva-Propranolol	00496502	TEV ACDEFGV
C07AA06	TIMOLOL							
Tab	Orl	5 mg				Timolol	00755842	AAP ACDEFGV
Tab	Orl	10 mg				Timolol	00755850	AAP ACDEFGV
Tab	Orl	20 mg				Timolol	00755869	AAP ACDEFGV
C07AA07	SOTALOL							
Tab	Orl	80 mg				Apo-Sotalol	02210428	APX ACDEFGV
						Jamp-Sotalol	02368617	JPC ACDEFGV
						pms-Sotalol	02238326	PMS ACDEFGV
						Sotalol	02385988	SIV ACDEFGV
Tab	Orl	160 mg				Apo-Sotalol	02167794	APX ACDEFGV
						Jamp-Sotalol	02368625	JPC ACDEFGV
						pms-Sotalol	02238327	PMS ACDEFGV
						Sotalol	02385996	SIV ACDEFGV
C07AA12	NADOLOL							
Tab	Orl	40 mg				Apo-Nadolol	00782505	APX ACDEFGV
						Mint-Nadolol	02496380	MNT ACDEFGV
Tab	Orl	80 mg				Apo-Nadolol	00782467	APX ACDEFGV
						Mint-Nadolol	02496399	MNT ACDEFGV
Tab	Orl	160 mg				Apo-Nadolol	00782475	APX ACDEFGV

**C07AB BETA BLOCKING AGENTS, SELECTIVE**

C07AB02 METOPROLOL

SRT	Orl	100 mg	AA-Metoprolol SR	02285169	AAP	ACDEFGV
Tab	Orl	25 mg	Apo-Metoprolol	02246010	APX	ACDEFGV
			Jamp-Metoprolol-L	02356813	JPC	ACDEFGV
			pms-Metoprolol-L	02248855	PMS	ACDEFGV
Tab	Orl	50 mg	Apo-Metoprolol (uncoated)	00618632	APX	ACDEFGV
			Apo-Metoprolol type "L"	00749354	APX	ACDEFGV
			Jamp-Metoprolol-L	02356821	JPC	ACDEFGV
			Metoprolol	02350394	SAS	ACDEFGV
			Metoprolol-L	02442124	SIV	ACDEFGV
			pms-Metoprolol-L	02230803	PMS	ACDEFGV
			Teva-Metoprolol (coated)	00648035	TEV	ACDEFGV
			Teva-Metoprolol (uncoated)	00842648	TEV	ACDEFGV
Tab	Orl	100 mg	Apo-Metoprolol (uncoated)	00618640	APX	ACDEFGV
			Apo-Metoprolol type "L"	00751170	APX	ACDEFGV
			Jamp-Metoprolol-L	02356848	JPC	ACDEFGV
			Metoprolol	02350408	SAS	ACDEFGV
			Metoprolol-L	02442132	SIV	ACDEFGV
			pms-Metoprolol-L	02230804	PMS	ACDEFGV
			Teva-Metoprolol (coated)	00648043	TEV	ACDEFGV
			Teva-Metoprolol (uncoated)	00842656	TEV	ACDEFGV

C07AB03 ATENOLOL

Tab	Orl	25 mg	Jamp-Atenolol	02367556	JPC	ACDEFGV
			Mar-Atenolol	02371979	MAR	ACDEFGV
			Mint-Atenolol	02368013	MNT	ACDEFGV
			pms-Atenolol	02246581	PMS	ACDEFGV
			Taro-Atenolol	02373963	SUN	ACDEFGV
			Teva-Atenolol	02266660	TEV	ACDEFGV

## C07AB03 ATENOLOL

Tab Orl 50 mg

Tenormin	02039532	SLP	ACDEFGV
Apo-Atenol	00773689	APX	ACDEFGV
Atenolol	02466465	SAS	ACDEFGV
Atenolol	02238316	SIV	ACDEFGV
Jamp-Atenolol	02367564	JPC	ACDEFGV
Mar-Atenolol	02371987	MAR	ACDEFGV
Mint-Atenolol	02368021	MNT	ACDEFGV
pms-Atenolol	02237600	PMS	ACDEFGV
Taro-Atenolol	02267985	SUN	ACDEFGV
Teva-Atenolol	02171791	TEV	ACDEFGV

Tab Orl 100 mg

Tenormin	02039540	SLP	ACDEFGV
Apo-Atenol	00773697	APX	ACDEFGV
Atenolol	02466473	SAS	ACDEFGV
Atenolol	02238318	SIV	ACDEFGV
Jamp-Atenolol	02367572	JPC	ACDEFGV
Mar-Atenolol	02371995	MAR	ACDEFGV
Mint-Atenolol	02368048	MNT	ACDEFGV
pms-Atenolol	02237601	PMS	ACDEFGV
Taro-Atenolol	02267993	SUN	ACDEFGV
Teva-Atenolol	02171805	TEV	ACDEFGV

## C07AB04 ACEBUTOLOL

Tab Orl 100 mg

Apo-Acebutolol	02147602	APX	ACDEFGV
Teva-Acebutolol	02204517	TEV	ACDEFGV

Tab Orl 200 mg

Apo-Acebutolol	02147610	APX	ACDEFGV
Teva-Acebutolol	02204525	TEV	ACDEFGV

Tab Orl 400 mg

Apo-Acebutolol	02147629	APX	ACDEFGV
Teva-Acebutolol	02204533	TEV	ACDEFGV

## C07AB07 BISOPROLOL

Tab Orl 5 mg

Apo-Bisoprolol	02256134	APX	ACDEFGV
Bisoprolol	02391589	SAS	ACDEFGV
Bisoprolol	02495562	SIV	ACDEFGV
Jamp Bisoprolol	02518805	JPC	ACDEFGV
Mint-Bisoprolol	02465612	MNT	ACDEFGV
Sandoz Bisoprolol	02494035	SDZ	ACDEFGV
Teva-Bisoprolol	02267470	TEV	ACDEFGV

C07AB07 BISOPROLOL

Tab Orl 10 mg

Apo-Bisoprolol 02256177 APX ACDEFGV  
Bisoprolol 02391597 SAS ACDEFGV  
Bisoprolol 02495570 SIV ACDEFGV  
Jamp Bisoprolol 02518791 JPC ACDEFGV  
Mint-Bisoprolol 02465620 MNT ACDEFGV  
Sandoz Bisoprolol 02494043 SDZ ACDEFGV  
Teva-Bisoprolol 02267489 TEV ACDEFGV

**C07AG ALPHA AND BETA BLOCKING AGENTS**

C07AG01 LABETALOL

Tab Orl 100 mg

Trandate 02106272 PAL ACDEFGV  
Apo-Labetalol 02243538 APX ACDEFGV  
Riva-Labetalol 02489406 RIV ACDEFGV

Tab Orl 200 mg

Trandate 02106280 PAL ACDEFGV  
Apo-Labetalol 02243539 APX ACDEFGV  
Riva-Labetalol 02489414 RIV ACDEFGV

C07AG02 CARVEDILOL

Tab Orl 3.125 mg

Apo-Carvedilol 02247933 APX ACDEFGV  
Auro-Carvedilol 02418495 ARO ACDEFGV  
Carvedilol 02364913 SAS ACDEFGV  
Carvedilol 02248752 SIV ACDEFGV  
Jamp-Carvedilol 02368897 JPC ACDEFGV  
pms-Carvedilol 02245914 PMS ACDEFGV  
ratio-Carvedilol 02252309 TEV ACDEFGV

Tab Orl 6.25 mg

Apo-Carvedilol 02247934 APX ACDEFGV  
Auro-Carvedilol 02418509 ARO ACDEFGV  
Carvedilol 02364921 SAS ACDEFGV  
Carvedilol 02248753 SIV ACDEFGV  
Jamp-Carvedilol 02368900 JPC ACDEFGV  
pms-Carvedilol 02245915 PMS ACDEFGV  
ratio-Carvedilol 02252317 TEV ACDEFGV

C07AG02 CARVEDILOL

Tab Orl 12.5 mg

Apo-Carvedilol 02247935 APX ACDEFGV  
Auro-Carvedilol 02418517 ARO ACDEFGV  
Carvedilol 02364948 SAS ACDEFGV  
Carvedilol 02248754 SIV ACDEFGV  
Jamp-Carvedilol 02368919 JPC ACDEFGV  
pms-Carvedilol 02245916 PMS ACDEFGV  
ratio-Carvedilol 02252325 TEV ACDEFGV

Tab Orl 25 mg

Apo-Carvedilol 02247936 APX ACDEFGV  
Auro-Carvedilol 02418525 ARO ACDEFGV  
Carvedilol 02364956 SAS ACDEFGV  
Carvedilol 02248755 SIV ACDEFGV  
Jamp-Carvedilol 02368927 JPC ACDEFGV  
pms-Carvedilol 02245917 PMS ACDEFGV  
ratio-Carvedilol 02252333 TEV ACDEFGV

**C07C BETA BLOCKING AGENTS AND OTHER DIURETICS**

**C07CA BETA BLOCKING AGENTS, NON-SELECTIVE, OTHER DIURETICS**

C07CA03 PINDOLOL AND OTHER DIURETICS  
PINDOLOL / HYDROCHLOROTHIAZIDE

Tab Orl 10 mg / 25 mg

Viskazine 00568627 XPI ACDEFGV

Tab Orl 10 mg / 50 mg

Viskazine 00568635 XPI ACDEFGV

**C07CB BETA BLOCKING AGENTS, SELECTIVE, AND OTHER DIURETICS**

C07CB03 ATENOLOL AND OTHER DIURETICS  
ATENOLOL / CHLORTHALIDONE

Tab Orl 50 mg / 25 mg

AA-Atenidone 02248763 AAP ACDEFGV

Tab Orl 100 mg / 25 mg

AA-Atenidone 02248764 AAP ACDEFGV

**C08 CALCIUM CHANNEL BLOCKERS**

**C08C SELECTIVE CALCIUM CHANNEL BLOCKERS WITH MAINLY VASCULAR EFFECTS**

**C08CA DIHYDROPYRIDINE DERIVATIVES**

C08CA01 AMLODIPINE

Liq Orl 1 mg/mL

pdp-Amlodipine 02484706 PDP (SA)

## C08CA01 AMLODIPINE

Tab Orl 2.5 mg

Amlodipine	02492199	JPC	ACDEFGV
Amlodipine	02326795	PDL	ACDEFGV
Amlodipine	02478587	SAS	ACDEFGV
Amlodipine	02385783	SIV	ACDEFGV
Amlodipine Besylate	02419556	AHI	ACDEFGV
Jamp-Amlodipine	02357186	JPC	ACDEFGV
M-Amlodipine	02468018	MRA	ACDEFGV
Mar-Amlodipine	02371707	MAR	ACDEFGV
NRA-Amlodipine	02476452	NRA	ACDEFGV
pharma-Amlodipine	02469022	PMS	ACDEFGV
pms-Amlodipine	02295148	PMS	ACDEFGV
PRZ-Amlodipine	02522500	PRZ	ACDEFGV
Sandoz Amlodipine	02330474	SDZ	ACDEFGV

Tab Orl 5 mg

Norvasc	00878928	BGP	ACDEFGV
Act Amlodipine	02297485	ATV	ACDEFGV
Amlodipine	02429217	JPC	ACDEFGV
Amlodipine	02326809	PDL	ACDEFGV
Amlodipine	02331284	SAS	ACDEFGV
Amlodipine	02385791	SIV	ACDEFGV
Amlodipine Besylate	02419564	AHI	ACDEFGV
Apo-Amlodipine	02273373	APX	ACDEFGV
Auro-Amlodipine	02397072	ARO	ACDEFGV
Jamp-Amlodipine	02357194	JPC	ACDEFGV
M-Amlodipine	02468026	MRA	ACDEFGV
Mar-Amlodipine	02371715	MAR	ACDEFGV
Mint-Amlodipine	02362651	MNT	ACDEFGV
Mylan-Amlodipine	02272113	MYL	ACDEFGV
NRA-Amlodipine	02476460	NRA	ACDEFGV
pharma-Amlodipine	02469030	PMS	ACDEFGV
pms-Amlodipine	02284065	PMS	ACDEFGV
PRZ-Amlodipine	02522519	PRZ	ACDEFGV
Ran-Amlodipine	02321858	RAN	ACDEFGV
Sandoz Amlodipine	02284383	SDZ	ACDEFGV
Septa-Amlodipine	02357712	SPT	ACDEFGV

C08CA01 AMLODIPINE

Tab Orl 10 mg

Norvasc	00878936	BGP	ACDEFGV
Act Amlodipine	02297493	ATV	ACDEFGV
Amlodipine	02429225	JPC	ACDEFGV
Amlodipine	02326817	PDL	ACDEFGV
Amlodipine	02331292	SAS	ACDEFGV
Amlodipine	02385805	SIV	ACDEFGV
Amlodipine Besylate	02419572	AHI	ACDEFGV
Apo-Amlodipine	02273381	APX	ACDEFGV
Auro-Amlodipine	02397080	ARO	ACDEFGV
Jamp-Amlodipine	02357208	JPC	ACDEFGV
M-Amlodipine	02468034	MRA	ACDEFGV
Mar-Amlodipine	02371723	MAR	ACDEFGV
Mint-Amlodipine	02362678	MNT	ACDEFGV
Mylan-Amlodipine	02272121	MYL	ACDEFGV
NRA-Amlodipine	02476479	NRA	ACDEFGV
pharma-Amlodipine	02469049	PMS	ACDEFGV
pms-Amlodipine	02284073	PMS	ACDEFGV
PRZ-Amlodipine	02522527	PRZ	ACDEFGV
Ran-Amlodipine	02321866	RAN	ACDEFGV
Sandoz Amlodipine	02284391	SDZ	ACDEFGV
Septa-Amlodipine	02357720	SPT	ACDEFGV

C08CA02 FELODIPINE

ERT Orl 2.5 mg

Plendil	02057778	AZE	ACDEFGV
Apo-Felodipine	02452367	APX	ACDEFGV

ERT Orl 5 mg

Plendil	00851779	AZE	ACDEFGV
Apo-Felodipine	02452375	APX	ACDEFGV
Sandoz Felodipine	02280264	SDZ	ACDEFGV

ERT Orl 10 mg

Plendil	00851787	AZE	ACDEFGV
Apo-Felodipine	02452383	APX	ACDEFGV
Sandoz Felodipine	02280272	SDZ	ACDEFGV

C08CA05 NIFEDIPINE

Cap Orl 5 mg

Nifedipine	00725110	AAP	ACDEFGV
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Cap Orl 10 mg

Nifedipine	00755907	AAP	ACDEFGV
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C08CA05	NIFEDIPINE							
ERT	Orl	30 mg		Adalat XL	02155907	TEV	ACDEFGV	
				Mylan-Nifedipine Extended Release	02349167	MYL	ACDEFGV	
ERT	Orl	60 mg		Mylan-Nifedipine Extended Release	02321149	MYL	ACDEFGV	

C08CA06	NIMODIPINE							
Tab	Orl	30 mg		Nimotop	02325926	BAY	ACDEFGV	

**C08D SELECTIVE CALCIUM CHANNEL BLOCKERS WITH DIRECT CARDIAC EFFECTS**

**C08DA PHENYLALKYLAMINE DERIVATIVES**

C08DA01	VERAPAMIL							
SRT	Orl	120 mg		Isoptin SR	01907123	BGP	ACDEFGVW	
				Apo-Verapamil SR	02246893	APX	ACDEFGVW	
				Mylan-Verapamil SR	02210347	MYL	ACDEFGVW	
SRT	Orl	180 mg		Isoptin SR	01934317	BGP	ACDEFGVW	
				Apo-Verap SR	02246894	APX	ACDEFGVW	
				Mylan-Verapamil SR	02450488	MYL	ACDEFGVW	
SRT	Orl	240 mg		Isoptin SR	00742554	BGP	ACDEFGVW	
				Mylan-Verapamil SR	02450496	MYL	ACDEFGVW	
Tab	Orl	80 mg		Apo-Verap	00782483	APX	ACDEFGVW	
				Mylan-Verapamil	02237921	MYL	ACDEFGVW	
Tab	Orl	120 mg		Apo-Verap	00782491	APX	ACDEFGVW	
				Mylan-Verapamil	02237922	MYL	ACDEFGVW	

**C08DB BENZOTHIAZEPINE DERIVATIVES**

C08DB01	DILTIAZEM							
CDC	Orl	120 mg		Cardizem CD	02097249	BSL	ACDEFGV	
				Act Diltiazem CD	02370611	TEV	ACDEFGV	
				Apo-Diltiaz CD	02230997	APX	ACDEFGV	
				Diltiazem CD	02400421	SAS	ACDEFGV	
				Diltiazem CD	02445999	SIV	ACDEFGV	
				Jamp Diltiazem CD	02528037	JPC	ACDEFGV	
				Mar-Diltiazem CD	02484064	MAR	ACDEFGV	
				Sandoz Diltiazem CD (Disc/non disp Jul 28/24)	02243338	SDZ	ACDEFGV	
				Teva-Diltiazem CD	02242538	TEV	ACDEFGV	



## C08DB01 DILTIAZEM

CDC Orl 180 mg

Cardizem CD	02097257	BSL	ACDEFGV
Apo-Diltiaz CD	02230998	APX	ACDEFGV
Diltiazem CD	02400448	SAS	ACDEFGV
Diltiazem CD	02446006	SIV	ACDEFGV
Jamp Diltiazem CD	02528045	JPC	ACDEFGV
Mar-Diltiazem CD	02484072	MAR	ACDEFGV
Sandoz Diltiazem CD (Disc/non disp Jul 28/24)	02243339	SDZ	ACDEFGV
Teva-Diltiazem CD	02242539	TEV	ACDEFGV

CDC Orl 240 mg

Cardizem CD	02097265	BSL	ACDEFGV
Apo-Diltiaz CD	02230999	APX	ACDEFGV
Diltiazem CD	02400456	SAS	ACDEFGV
Diltiazem CD	02446014	SIV	ACDEFGV
Jamp Diltiazem CD	02528053	JPC	ACDEFGV
Mar-Diltiazem CD	02484080	MAR	ACDEFGV
Sandoz Diltiazem CD (Disc/non disp Jul 28/24)	02243340	SDZ	ACDEFGV
Teva-Diltiazem CD	02242540	TEV	ACDEFGV

CDC Orl 300 mg

Cardizem CD	02097273	BSL	ACDEFGV
Act Diltiazem CD	02370654	TEV	ACDEFGV
Apo-Diltiaz CD	02229526	APX	ACDEFGV
Diltiazem CD	02400464	SAS	ACDEFGV
Diltiazem CD	02446022	SIV	ACDEFGV
Jamp Diltiazem CD	02528061	JPC	ACDEFGV
Mar-Diltiazem CD	02484099	MAR	ACDEFGV
Sandoz Diltiazem CD (Disc/non disp Jul 28/24)	02243341	SDZ	ACDEFGV
Teva-Diltiazem CD	02242541	TEV	ACDEFGV

ERC Orl 120 mg

Tiazac	02231150	BSL	ACDEFGV
Act Diltiazem T	02370441	TEV	ACDEFGV
Diltiazem T	02516101	SAS	ACDEFGV
Jamp-Diltiazem T	02495376	JPC	ACDEFGV
Mar-Diltiazem T	02465353	MAR	ACDEFGV
Sandoz Diltiazem T (Disc/non disp Jul 28/24)	02245918	SDZ	ACDEFGV
Teva-Diltiazem ER	02271605	BSL	ACDEFGV

C08DB01 DILTIAZEM

ERC Orl 180 mg

Tiazac 02231151 BSL ACDEFGV  
 Act Diltiazem T 02370492 TEV ACDEFGV  
 Diltiazem T 02516128 SAS ACDEFGV  
 Jamp-Diltiazem T 02495384 JPC ACDEFGV  
 Mar-Diltiazem T 02465361 MAR ACDEFGV  
 Sandoz Diltiazem T (Disc/non disp Jul 28/24) 02245919 SDZ ACDEFGV  
 Teva-Diltiazem ER 02271613 BSL ACDEFGV

ERC Orl 240 mg

Tiazac 02231152 BSL ACDEFGV  
 Act Diltiazem T 02370506 TEV ACDEFGV  
 Diltiazem T 02516136 SAS ACDEFGV  
 Jamp-Diltiazem T 02495392 JPC ACDEFGV  
 Mar-Diltiazem T 02465388 MAR ACDEFGV  
 Teva-Diltiazem ER 02271621 BSL ACDEFGV

ERC Orl 300 mg

Tiazac 02231154 BSL ACDEFGV  
 Diltiazem T 02516144 SAS ACDEFGV  
 Jamp-Diltiazem T 02495406 JPC ACDEFGV  
 Mar-Diltiazem T 02465396 MAR ACDEFGV  
 Sandoz Diltiazem T (Disc/non disp Jul 28/24) 02245921 SDZ ACDEFGV  
 Teva-Diltiazem ER 02271648 BSL ACDEFGV

ERC Orl 360 mg

Tiazac 02231155 BSL ACDEFGV  
 Act Diltiazem T 02370522 TEV ACDEFGV  
 Diltiazem T 02516152 SAS ACDEFGV  
 Jamp-Diltiazem T 02495414 JPC ACDEFGV  
 Mar-Diltiazem T 02465418 MAR ACDEFGV  
 Teva-Diltiazem ER 02271656 BSL ACDEFGV

ERT Orl 120 mg

Tiazac XC 02256738 BSL ACDEFGV

ERT Orl 180 mg

Tiazac XC 02256746 BSL ACDEFGV  
 Teva-Diltiazem XC 02429322 TEV ACDEFGV

ERT Orl 240 mg

Tiazac XC 02256754 BSL ACDEFGV  
 Teva-Diltiazem XC 02429330 TEV ACDEFGV

ERT Orl 300 mg

Tiazac XC 02256762 BSL ACDEFGV  
 Teva-Diltiazem XC 02429349 TEV ACDEFGV

C08DB01	DILTIAZEM						
ERT	Orl	360 mg		Tiazac XC	02256770	BSL	ACDEFGV
				Teva-Diltiazem XC	02429357	TEV	ACDEFGV
Tab	Orl	30 mg		AA-Diltiaz	00771376	AAP	ACDEFGV
				Teva-Diltiazem	00862924	TEV	ACDEFGV
Tab	Orl	60 mg		AA-Diltiaz	00771384	AAP	ACDEFGV
				Teva-Diltiazem	00862932	TEV	ACDEFGV

**C09 AGENTS ACTING ON THE RENIN-ANGIOTENSIN SYSTEM**

**C09A ACE INHIBITORS, PLAIN**

**C09AA ACE INHIBITORS, PLAIN**

C09AA01	CAPTOPRIL						
Tab	Orl	12.5 mg		Teva-Captopril	01942964	TEV	ACDEFGV
Tab	Orl	25 mg		Teva-Captopril	01942972	TEV	ACDEFGV
Tab	Orl	50 mg		Teva-Captopril	01942980	TEV	ACDEFGV
Tab	Orl	100 mg		Teva-Captopril	01942999	TEV	ACDEFGV
C09AA02	ENALAPRIL						
Tab	Orl	2.5 mg		Act Enalapril	02291878	TEV	ACDEFGV
				Apo-Enalapril	02020025	APX	ACDEFGV
				Enalapril	02400650	SAS	ACDEFGV
				Enalapril	02442957	SIV	ACDEFGV
				Jamp-Enalapril	02474786	JPC	ACDEFGV
				Mar-Enalapril	02459450	MAR	ACDEFGV
				Sandoz Enalapril	02299933	SDZ	ACDEFGV
				Taro-Enalapril	02352230	SUN	ACDEFGV
Tab	Orl	5 mg		Vasotec	00708879	ORG	ACDEFGV
				Act Enalapril	02291886	TEV	ACDEFGV
				Apo-Enalapril	02019884	APX	ACDEFGV
				Enalapril	02400669	SAS	ACDEFGV
				Enalapril	02442965	SIV	ACDEFGV
				Jamp-Enalapril	02474794	JPC	ACDEFGV
				Mar-Enalapril	02459469	MAR	ACDEFGV
				Sandoz Enalapril	02299941	SDZ	ACDEFGV
				Taro-Enalapril	02352249	SUN	ACDEFGV

C09AA02 ENALAPRIL

Tab Orl 10 mg

Vasotec 00670901 ORG ACDEFGV  
 Act Enalapril 02291894 TEV ACDEFGV  
 Apo-Enalapril 02019892 APX ACDEFGV  
 Enalapril 02400677 SAS ACDEFGV  
 Enalapril 02442973 SIV ACDEFGV  
 Jamp-Enalapril 02474808 JPC ACDEFGV  
 Mar-Enalapril 02444771 MAR ACDEFGV  
 Sandoz Enalapril 02299968 SDZ ACDEFGV  
 Taro-Enalapril 02352257 SUN ACDEFGV

Tab Orl 20 mg

Vasotec 00670928 ORG ACDEFGV  
 Act Enalapril 02291908 TEV ACDEFGV  
 Apo-Enalapril 02019906 APX ACDEFGV  
 Enalapril 02400685 SAS ACDEFGV  
 Enalapril 02442981 SIV ACDEFGV  
 Jamp-Enalapril 02474816 JPC ACDEFGV  
 Mar-Enalapril 02444798 MAR ACDEFGV  
 Sandoz Enalapril 02299976 SDZ ACDEFGV  
 Taro-Enalapril 02352265 SUN ACDEFGV

C09AA03 LISINOPRIL

Tab Orl 5 mg

Zestril 02049333 SLP ACDEFGV  
 Apo-Lisinopril 02217481 APX ACDEFGV  
 Auro-Lisinopril 02394472 ARO ACDEFGV  
 Lisinopril 02525186 SAS ACDEFGV  
 Lisinopril 02386232 SIV ACDEFGV  
 Teva-Lisinopril Z 02285118 TEV ACDEFGV

Tab Orl 10 mg

Zestril 02049376 SLP ACDEFGV  
 Apo-Lisinopril 02217503 APX ACDEFGV  
 Auro-Lisinopril 02394480 ARO ACDEFGV  
 Lisinopril 02525194 SAS ACDEFGV  
 Lisinopril 02386240 SIV ACDEFGV  
 Teva-Lisinopril Z 02285126 TEV ACDEFGV

## C09AA03 LISINOPRIL

Tab Orl 20 mg

Zestril	02049384	SLP	ACDEFGV
Apo-Lisinopril	02217511	APX	ACDEFGV
Auro-Lisinopril	02394499	ARO	ACDEFGV
Lisinopril	02525208	SAS	ACDEFGV
Lisinopril	02386259	SIV	ACDEFGV
Teva-Lisinopril Z	02285134	TEV	ACDEFGV

## C09AA04 PERINDOPRIL

Tab Orl 2 mg

Coversyl	02123274	SEV	ACDEFGV
Apo-Perindopril	02289261	APX	ACDEFGV
Auro-Perindopril	02459817	ARO	ACDEFGV
Jamp Perindopril Erbumine	02527200	JPC	ACDEFGV
Jamp-Perindopril	02477009	JPC	ACDEFGV
M-Perindopril Erbumine	02482924	MRA	ACDEFGV
Mar-Perindopril	02474824	MAR	ACDEFGV
Mint-Perindopril	02476762	MNT	ACDEFGV
NRA-Perindopril	02489015	NRA	ACDEFGV
Perindopril Erbumine	02481634	SAS	ACDEFGV
Perindopril Erbumine	02479877	SIV	ACDEFGV
pms-Perindopril	02470675	PMS	ACDEFGV
Sandoz Perindopril	02470225	SDZ	ACDEFGV
Teva-Perindopril	02464985	TEV	ACDEFGV

Tab Orl 4 mg

Coversyl	02123282	SEV	ACDEFGV
Apo-Perindopril	02289288	APX	ACDEFGV
Auro-Perindopril	02459825	ARO	ACDEFGV
Jamp Perindopril Erbumine	02527219	JPC	ACDEFGV
Jamp-Perindopril	02477017	JPC	ACDEFGV
M-Perindopril Erbumine	02482932	MRA	ACDEFGV
Mar-Perindopril	02474832	MAR	ACDEFGV
Mint-Perindopril	02476770	MNT	ACDEFGV
NRA-Perindopril	02489023	NRA	ACDEFGV
Perindopril Erbumine	02481642	SAS	ACDEFGV
Perindopril Erbumine	02479885	SIV	ACDEFGV
pms-Perindopril	02470683	PMS	ACDEFGV
Sandoz Perindopril	02470233	SDZ	ACDEFGV
Teva-Perindopril	02464993	TEV	ACDEFGV

C09AA04 PERINDOPRIL

Tab Orl 8 mg

Coversyl	02246624	SEV	ACDEFGV
Apo-Perindopril	02289296	APX	ACDEFGV
Auro-Perindopril	02459833	ARO	ACDEFGV
Jamp Perindopril Erbumine	02527227	JPC	ACDEFGV
Jamp-Perindopril	02477025	JPC	ACDEFGV
M-Perindopril Erbumine	02482940	MRA	ACDEFGV
Mar-Perindopril	02474840	MAR	ACDEFGV
Mint-Perindopril	02476789	MNT	ACDEFGV
NRA-Perindopril	02489031	NRA	ACDEFGV
Perindopril Erbumine	02481650	SAS	ACDEFGV
Perindopril Erbumine	02479893	SIV	ACDEFGV
pms-Perindopril	02470691	PMS	ACDEFGV
Sandoz Perindopril	02470241	SDZ	ACDEFGV
Teva-Perindopril	02465000	TEV	ACDEFGV

C09AA05 RAMIPRIL

Cap Orl 1.25 mg

Altace	02221829	BSL	ACDEFGV
Apo-Ramipril	02251515	APX	ACDEFGV
Auro-Ramipril	02387387	ARO	ACDEFGV
Mar-Ramipril	02420457	MAR	ACDEFGV
pharma-Ramipril	02469057	PMS	ACDEFGV
Pro-Ramipril	02310023	PDL	ACDEFGV
Ramipril	02308363	SIV	ACDEFGV
Taro-Ramipril	02310503	SUN	ACDEFGV

Cap Orl 2.5 mg

Altace	02221837	BSL	ACDEFGV
Apo-Ramipril	02251531	APX	ACDEFGV
Auro-Ramipril	02387395	ARO	ACDEFGV
Jamp-Ramipril	02331128	JPC	ACDEFGV
Mar-Ramipril	02420465	MAR	ACDEFGV
Mint-Ramipril	02421305	MNT	ACDEFGV
NRA-Ramipril	02486172	NRA	ACDEFGV
pharma-Ramipril	02469065	PMS	ACDEFGV
Pro-Ramipril	02310066	PDL	ACDEFGV
Ramipril	02374846	SAS	ACDEFGV
Ramipril	02287927	SIV	ACDEFGV
Taro-Ramipril	02310511	SUN	ACDEFGV
Teva-Ramipril	02247945	TEV	ACDEFGV

C09AA05 RAMIPRIL

Cap Orl 5 mg

Altace 02221845 BSL ACDEFGV  
 Apo-Ramipril 02251574 APX ACDEFGV  
 Auro-Ramipril 02387409 ARO ACDEFGV  
 Jamp-Ramipril 02331136 JPC ACDEFGV  
 Mar-Ramipril 02420473 MAR ACDEFGV  
 Mint-Ramipril 02421313 MNT ACDEFGV  
 NRA-Ramipril 02486180 NRA ACDEFGV  
 pharma-Ramipril 02469073 PMS ACDEFGV  
 Pro-Ramipril 02310074 PDL ACDEFGV  
 Ramipril 02374854 SAS ACDEFGV  
 Ramipril 02287935 SIV ACDEFGV  
 Taro-Ramipril 02310538 SUN ACDEFGV  
 Teva-Ramipril 02247946 TEV ACDEFGV

Cap Orl 10 mg

Altace 02221853 BSL ACDEFGV  
 Apo-Ramipril 02251582 APX ACDEFGV  
 Auro-Ramipril 02387417 ARO ACDEFGV  
 Jamp-Ramipril 02331144 JPC ACDEFGV  
 Mar-Ramipril 02420481 MAR ACDEFGV  
 Mint-Ramipril 02421321 MNT ACDEFGV  
 NRA-Ramipril 02486199 NRA ACDEFGV  
 pharma-Ramipril 02469081 PMS ACDEFGV  
 Pro-Ramipril 02310104 PDL ACDEFGV  
 Ramipril 02374862 SAS ACDEFGV  
 Ramipril 02287943 SIV ACDEFGV  
 Taro-Ramipril 02310546 SUN ACDEFGV  
 Teva-Ramipril 02247947 TEV ACDEFGV

Cap Orl 15 mg

Altace 02281112 BSL ACDEFGV  
 Apo-Ramipril 02325381 APX ACDEFGV

C09AA06 QUINAPRIL

Tab Orl 5 mg

Accupril 01947664 PFI ACDEFGV  
 Apo-Quinapril 02248499 APX ACDEFGV  
 pms-Quinapril 02340550 PMS ACDEFGV

Tab Orl 10 mg

Accupril 01947672 PFI ACDEFGV  
 Apo-Quinapril 02248500 APX ACDEFGV  
 pms-Quinapril 02340569 PMS ACDEFGV

C09AA06	QUINAPRIL							
Tab	Orl	20 mg		Accupril	01947680	PFI	ACDEFGV	
				Apo-Quinapril	02248501	APX	ACDEFGV	
				pms-Quinapril	02340577	PMS	ACDEFGV	
Tab	Orl	40 mg		Accupril	01947699	PFI	ACDEFGV	
				Apo-Quinapril	02248502	APX	ACDEFGV	
				pms-Quinapril	02340585	PMS	ACDEFGV	
C09AA07	BENAZEPRIL							
Tab	Orl	5 mg		Benazepril	02290332	AAP	ACDEFGV	
Tab	Orl	10 mg		Benazepril	02290340	AAP	ACDEFGV	
Tab	Orl	20 mg		Benazepril	02273918	AAP	ACDEFGV	
C09AA08	CILAZAPRIL							
Tab	Orl	1 mg		Mylan-Cilazapril	02283778	MYL	ACDEFGV	
Tab	Orl	2.5 mg		Inhibace	01911473	MSD	ACDEFGV	
				Apo-Cilazapril	02291142	APX	ACDEFGV	
				Mylan-Cilazapril	02283786	MYL	ACDEFGV	
Tab	Orl	5 mg		Inhibace	01911481	MSD	ACDEFGV	
				Apo-Cilazapril	02291150	APX	ACDEFGV	
				Mylan-Cilazapril	02283794	MYL	ACDEFGV	
C09AA09	FOSINOPRIL							
Tab	Orl	10 mg		Apo-Fosinopril	02266008	APX	ACDEFGV	
				Fosinopril	02459388	SAS	ACDEFGV	
				Jamp-Fosinopril	02331004	JPC	ACDEFGV	
				Ran-Fosinopril	02294524	RAN	ACDEFGV	
				Teva-Fosinopril	02247802	TEV	ACDEFGV	
Tab	Orl	20 mg		Apo-Fosinopril	02266016	APX	ACDEFGV	
				Fosinopril	02459396	SAS	ACDEFGV	
				Jamp-Fosinopril	02331012	JPC	ACDEFGV	
				Ran-Fosinopril	02294532	RAN	ACDEFGV	
				Teva-Fosinopril	02247803	TEV	ACDEFGV	
C09AA10	TRANDOLAPRIL							



C09AA10 TRANDOLAPRIL

Cap Orl 0.5 mg

Mavik 02231457 BGP ACDEFGV  
 Auro-Trandolapril 02471868 ARO ACDEFGV  
 pms-Trandolapril 02357755 PMS ACDEFGV  
 Sandoz Trandolapril 02325721 SDZ ACDEFGV

Cap Orl 1 mg

Mavik 02231459 BGP ACDEFGV  
 Auro-Trandolapril 02471876 ARO ACDEFGV  
 pms-Trandolapril 02357763 PMS ACDEFGV  
 Sandoz Trandolapril 02325748 SDZ ACDEFGV  
 Trandolapril 02525046 SAS ACDEFGV  
 Trandolapril 02526565 SIV ACDEFGV

Cap Orl 2 mg

Mavik 02231460 BGP ACDEFGV  
 Auro-Trandolapril 02471884 ARO ACDEFGV  
 pms-Trandolapril 02357771 PMS ACDEFGV  
 Sandoz Trandolapril 02325756 SDZ ACDEFGV  
 Trandolapril 02525054 SAS ACDEFGV  
 Trandolapril 02526573 SIV ACDEFGV

Cap Orl 4 mg

Mavik 02239267 BGP ACDEFGV  
 Auro-Trandolapril 02471892 ARO ACDEFGV  
 pms-Trandolapril 02357798 PMS ACDEFGV  
 Sandoz Trandolapril 02325764 SDZ ACDEFGV  
 Trandolapril 02525070 SAS ACDEFGV  
 Trandolapril 02526581 SIV ACDEFGV

**C09B ACE-INHIBITORS, COMBINATIONS**

**C09BA ACE-INHIBITORS AND DIURETICS**

C09BA02 ENALAPRIL AND DIURETICS  
 ENALAPRIL / HYDROCHLOROTHIAZIDE

Tab Orl 5 mg / 12.5 mg

Enalapril/HCTZ 02352923 AAP ACDEFGV

Tab Orl 10 mg / 25 mg

Vaseretic 00657298 ORG ACDEFGV  
 Enalapril/HCTZ 02352931 AAP ACDEFGV

C09BA03 LISINOPRIL AND DIURETICS  
 LISINOPRIL / HYDROCHLOROTHIAZIDE

C09BA03	LISINOPRIL AND DIURETICS							
	LISINOPRIL / HYDROCHLOROTHIAZIDE							
Tab	Orl	10 mg / 12.5 mg	Zestoretic	02103729	SLP	ACDEFGV		
			Lisinopril HCTZ (Type Z)	02362945	SAS	ACDEFGV		
			Sandoz Lisinopril HCT	02302365	SDZ	ACDEFGV		
			Teva-Lisinopril HCTZ (Type Z)	02301768	TEV	ACDEFGV		
Tab	Orl	20 mg / 12.5 mg	Zestoretic	02045737	SLP	ACDEFGV		
			Lisinopril HCTZ (Type Z)	02362953	SAS	ACDEFGV		
			Sandoz Lisinopril HCT	02302373	SDZ	ACDEFGV		
			Teva-Lisinopril HCTZ (Type Z)	02301776	TEV	ACDEFGV		
Tab	Orl	20 mg / 25 mg	Zestoretic	02045729	SLP	ACDEFGV		
			Lisinopril HCTZ (Type Z)	02362961	SAS	ACDEFGV		
			Sandoz Lisinopril HCT	02302381	SDZ	ACDEFGV		
			Teva-Lisinopril HCTZ (Type P)	02302152	TEV	ACDEFGV		
			Teva-Lisinopril HCTZ (Type Z)	02301784	TEV	ACDEFGV		
C09BA04	PERINDOPRIL AND DIURETICS							
	PERINDOPRIL / INDAPAMIDE							
Tab	Orl	4 mg / 1.25 mg	Coversyl Plus	02246569	SEV	ACDEFGV		
			Apo-Perindopril-Indapamide	02297574	APX	ACDEFGV		
			Perindopril Erbumine/Indapamide	02479834	SIV	ACDEFGV		
			Perindopril/Indapamide	02519720	SAS	ACDEFGV		
			Sandoz Perindopril/Indapamide	02470438	SDZ	ACDEFGV		
			Teva-Perindopril/Indapamide	02464020	TEV	ACDEFGV		
Tab	Orl	8 mg / 2.5 mg	Coversyl Plus HD	02321653	SEV	ACDEFGV		
			Apo-Perindopril-Indapamide	02453061	APX	ACDEFGV		
			Perindopril Erbumine/Indapamide	02479842	SIV	ACDEFGV		
			Perindopril/Indapamide	02519739	SAS	ACDEFGV		
			Sandoz Perindopril/Indapamide	02470446	SDZ	ACDEFGV		
			Teva-Perindopril/Indapamide	02464039	TEV	ACDEFGV		
C09BA05	RAMIPRIL AND DIURETICS							
	RAMIPRIL / HYDROCHLOROTHIAZIDE							
Tab	Orl	2.5 mg / 12.5 mg	Altace HCT	02283131	BSL	ACDEFGV		
			Taro-Ramipril HCTZ	02449439	SUN	ACDEFGV		
Tab	Orl	5 mg / 12.5 mg	Altace HCT	02283158	BSL	ACDEFGV		
			Taro-Ramipril HCTZ	02449447	SUN	ACDEFGV		

C09BA05	RAMIPRIL AND DIURETICS						
	RAMIPRIL / HYDROCHLOROTHIAZIDE						
Tab	Orl	5 mg / 25 mg		Altace HCT	02283174	BSL	ACDEFGV
				Taro-Ramipril HCTZ	02449463	SUN	ACDEFGV
Tab	Orl	10 mg / 12.5 mg		Altace HCT	02283166	BSL	ACDEFGV
				pms-Ramipril-HCTZ	02342154	PMS	ACDEFGV
				Taro-Ramipril HCTZ	02449455	SUN	ACDEFGV
Tab	Orl	10 mg / 25 mg		Altace HCT	02283182	BSL	ACDEFGV
				pms-Ramipril-HCTZ	02342170	PMS	ACDEFGV
				Taro-Ramipril HCTZ	02449471	SUN	ACDEFGV

C09BA06	QUINAPRIL AND DIURETICS						
	QUINAPRIL / HYDROCHLOROTHIAZIDE						
Tab	Orl	10 mg / 12.5 mg		Accuretic	02237367	PFI	ACDEFGV
				Apo-Quinapril/HCTZ	02408767	APX	ACDEFGV
				Auro-Quinapril HCTZ	02473291	ARO	ACDEFGV
Tab	Orl	20 mg / 12.5 mg		Accuretic	02237368	PFI	ACDEFGV
				Apo-Quinapril/HCTZ	02408775	APX	ACDEFGV
				Auro-Quinapril HCTZ	02473305	ARO	ACDEFGV
Tab	Orl	20 mg / 25 mg		Accuretic	02237369	PFI	ACDEFGV
				Apo-Quinapril/HCTZ	02408783	APX	ACDEFGV
				Auro-Quinapril HCTZ	02473321	ARO	ACDEFGV

C09BA08	CILAZAPRIL AND DIURETICS						
	CILAZAPRIL / HYDROCHLOROTHIAZIDE						
Tab	Orl	5 mg / 12.5 mg		Inhibace Plus	02181479	HLR	ACDEFGV
				Apo-Cilazapril/HCTZ	02284987	APX	ACDEFGV
				Teva-Cilazapril/HCTZ	02313731	TEV	ACDEFGV

**C09C ANGIOTENSIN II ANTAGONISTS, PLAIN**

**C09CA ANGIOTENSIN II ANTAGONISTS, PLAIN**

C09CA01 LOSARTAN

C09CA01 LOSARTAN

Tab Orl 25 mg

Cozaar 02182815 ORG ACDEFGV  
 Apo-Losartan 02379058 APX ACDEFGV  
 Auro-Losartan 02403323 ARO ACDEFGV  
 Jamp-Losartan 02398834 JPC ACDEFGV  
 Losartan 02388863 SAS ACDEFGV  
 Losartan 02388790 SIV ACDEFGV  
 Mint-Losartan 02405733 MNT ACDEFGV  
 pms-Losartan 02309750 PMS ACDEFGV  
 Sandoz Losartan 02313332 SDZ ACDEFGV  
 Septa-Losartan 02424967 SPT ACDEFGV  
 Teva-Losartan 02380838 TEV ACDEFGV

Tab Orl 50 mg

Cozaar 02182874 ORG ACDEFGV  
 Apo-Losartan 02353504 APX ACDEFGV  
 Auro-Losartan 02403331 ARO ACDEFGV  
 Jamp-Losartan 02398842 JPC ACDEFGV  
 Losartan 02388871 SAS ACDEFGV  
 Losartan 02388804 SIV ACDEFGV  
 Mint-Losartan 02405741 MNT ACDEFGV  
 pms-Losartan 02309769 PMS ACDEFGV  
 Sandoz Losartan 02313340 SDZ ACDEFGV  
 Septa-Losartan 02424975 SPT ACDEFGV  
 Teva-Losartan 02357968 TEV ACDEFGV

Tab Orl 100 mg

Cozaar 02182882 ORG ACDEFGV  
 Apo-Losartan 02353512 APX ACDEFGV  
 Auro-Losartan 02403358 ARO ACDEFGV  
 Jamp-Losartan 02398850 JPC ACDEFGV  
 Losartan 02388898 SAS ACDEFGV  
 Losartan 02388812 SIV ACDEFGV  
 Mint-Losartan 02405768 MNT ACDEFGV  
 pms-Losartan 02309777 PMS ACDEFGV  
 Sandoz Losartan 02313359 SDZ ACDEFGV  
 Septa-Losartan 02424983 SPT ACDEFGV  
 Teva-Losartan 02357976 TEV ACDEFGV

C09CA02 EPROSARTAN

Tab Orl 400 mg

Teveten 02240432 BGP ACDEFGV

Tab Orl 600 mg

Teveten 02243942 BGP ACDEFGV

C09CA03 VALSARTAN

Tab Orl 40 mg

Diovan 02270528 NVR ACDEFGV  
 Auro-Valsartan 02414201 ARO ACDEFGV  
 M-Valsartan 02524511 MRA ACDEFGV  
 Sandoz Valsartan 02356740 SDZ ACDEFGV  
 Taro-Valsartan 02363062 SUN ACDEFGV  
 Teva-Valsartan 02356643 TEV ACDEFGV  
 Valsartan 02366940 SAS ACDEFGV  
 Valsartan 02384523 SIV ACDEFGV

Tab Orl 80 mg

Diovan 02244781 NVR ACDEFGV  
 Auro-Valsartan 02414228 ARO ACDEFGV  
 M-Valsartan 02524538 MRA ACDEFGV  
 Sandoz Valsartan 02356759 SDZ ACDEFGV  
 Taro-Valsartan 02363100 SUN ACDEFGV  
 Teva-Valsartan 02356651 TEV ACDEFGV  
 Valsartan 02366959 SAS ACDEFGV  
 Valsartan 02384531 SIV ACDEFGV

Tab Orl 160 mg

Diovan 02244782 NVR ACDEFGV  
 Auro-Valsartan 02414236 ARO ACDEFGV  
 M-Valsartan 02524546 MRA ACDEFGV  
 Sandoz Valsartan 02356767 SDZ ACDEFGV  
 Taro-Valsartan 02363119 SUN ACDEFGV  
 Teva-Valsartan 02356678 TEV ACDEFGV  
 Valsartan 02366967 SAS ACDEFGV  
 Valsartan 02384558 SIV ACDEFGV

Tab Orl 320 mg

Diovan 02289504 NVR ACDEFGV  
 Auro-Valsartan 02414244 ARO ACDEFGV  
 Sandoz Valsartan 02356775 SDZ ACDEFGV  
 Teva-Valsartan 02356686 TEV ACDEFGV  
 Valsartan 02366975 SAS ACDEFGV  
 Valsartan 02384566 SIV ACDEFGV

C09CA04 IRBESARTAN

C09CA04 IRBESARTAN

Tab Orl 75 mg

Avapro	02237923	SAV	ACDEFGV
Auro-Irbesartan	02406098	ARO	ACDEFGV
Irbesartan	02365197	PDL	ACDEFGV
Irbesartan	02372347	SAS	ACDEFGV
Irbesartan	02385287	SIV	ACDEFGV
M-Irbesartan	02524813	MRA	ACDEFGV
Mint-Irbesartan	02422980	MNT	ACDEFGV
pms-Irbesartan	02317060	PMS	ACDEFGV
Sandoz Irbesartan	02328461	SDZ	ACDEFGV
Taro-Irbesartan	02406810	SUN	ACDEFGV
Teva-Irbesartan	02316390	TEV	ACDEFGV

Tab Orl 150 mg

Avapro	02237924	SAV	ACDEFGV
Auro-Irbesartan	02406101	ARO	ACDEFGV
Irbesartan	02365200	PDL	ACDEFGV
Irbesartan	02372371	SAS	ACDEFGV
Irbesartan	02385295	SIV	ACDEFGV
M-Irbesartan	02524821	MRA	ACDEFGV
Mint-Irbesartan	02422999	MNT	ACDEFGV
pms-Irbesartan	02317079	PMS	ACDEFGV
Sandoz Irbesartan	02328488	SDZ	ACDEFGV
Taro-Irbesartan	02406829	SUN	ACDEFGV
Teva-Irbesartan	02316404	TEV	ACDEFGV

Tab Orl 300 mg

Avapro	02237925	SAV	ACDEFGV
Auro-Irbesartan	02406128	ARO	ACDEFGV
Irbesartan	02365219	PDL	ACDEFGV
Irbesartan	02372398	SAS	ACDEFGV
Irbesartan	02385309	SIV	ACDEFGV
M-Irbesartan	02524848	MRA	ACDEFGV
Mint-Irbesartan	02423006	MNT	ACDEFGV
pms-Irbesartan	02317087	PMS	ACDEFGV
Sandoz Irbesartan	02328496	SDZ	ACDEFGV
Taro-Irbesartan	02406837	SUN	ACDEFGV
Teva-Irbesartan	02316412	TEV	ACDEFGV

C09CA06 CANDESARTAN

C09CA06 CANDESARTAN

Tab Orl 4 mg

Atacand	02239090	AZE	ACDEFGV
Apo-Candesartan	02365340	APX	ACDEFGV
Auro-Candesartan	02445786	ARO	ACDEFGV
Candesartan	02388901	SAS	ACDEFGV
Candesartan	02528258	SIV	ACDEFGV
Candesartan Cilexetil	02379260	AHI	ACDEFGV
Mint-Candesartan	02476908	MNT	ACDEFGV
pms-Candesartan	02391171	PMS	ACDEFGV
Sandoz Candesartan	02326957	SDZ	ACDEFGV
Taro-Candesartan	02380684	SUN	ACDEFGV

Tab Orl 8 mg

Atacand	02239091	AZE	ACDEFGV
Apo-Candesartan	02365359	APX	ACDEFGV
Auro-Candesartan	02445794	ARO	ACDEFGV
Candesartan	02388928	SAS	ACDEFGV
Candesartan	02388707	SIV	ACDEFGV
Candesartan Cilexetil	02379279	AHI	ACDEFGV
Jamp-Candesartan	02386518	JPC	ACDEFGV
Mint-Candesartan	02476916	MNT	ACDEFGV
pms-Candesartan	02391198	PMS	ACDEFGV
Sandoz Candesartan	02326965	SDZ	ACDEFGV
Taro-Candesartan	02380692	SUN	ACDEFGV
Teva-Candesartan	02366312	TEV	ACDEFGV

Tab Orl 16 mg

Atacand	02239092	AZE	ACDEFGV
Apo-Candesartan	02365367	APX	ACDEFGV
Auro-Candesartan	02445808	ARO	ACDEFGV
Candesartan	02388936	SAS	ACDEFGV
Candesartan	02388715	SIV	ACDEFGV
Candesartan Cilexetil	02379287	AHI	ACDEFGV
Jamp-Candesartan	02386526	JPC	ACDEFGV
Mint-Candesartan	02476924	MNT	ACDEFGV
pms-Candesartan	02391201	PMS	ACDEFGV
Sandoz Candesartan	02326973	SDZ	ACDEFGV
Taro-Candesartan	02380706	SUN	ACDEFGV
Teva-Candesartan	02366320	TEV	ACDEFGV

## C09CA06 CANDESARTAN

Tab Orl 32 mg

Atacand	02311658	AZE	ACDEFGV
Apo-Candesartan	02399105	APX	ACDEFGV
Auro-Candesartan	02445816	ARO	ACDEFGV
Candesartan	02435845	SAS	ACDEFGV
Candesartan	02528266	SIV	ACDEFGV
Candesartan Cilexetil	02379295	AHI	ACDEFGV
Jamp-Candesartan	02386534	JPC	ACDEFGV
pms-Candesartan	02391228	PMS	ACDEFGV
Sandoz Candesartan	02417340	SDZ	ACDEFGV
Taro-Candesartan	02380714	SUN	ACDEFGV
Teva-Candesartan	02366339	TEV	ACDEFGV

## C09CA07 TELMISARTAN

Tab Orl 40 mg

Micardis	02240769	BOE	ACDEFGV
Auro-Telmisartan	02453568	ARO	ACDEFGV
Jamp Telmisartan	02386755	JPC	ACDEFGV
Mint-Telmisartan	02486369	MNT	ACDEFGV
NRA-Telmisartan	02503794	NRA	ACDEFGV
pms-Telmisartan	02499622	PMS	ACDEFGV
Sandoz Telmisartan	02375958	SDZ	ACDEFGV
Telmisartan	02407485	AHI	ACDEFGV
Telmisartan	02388944	SAS	ACDEFGV
Telmisartan	02390345	SIV	ACDEFGV
Teva-Telmisartan	02320177	TEV	ACDEFGV

Tab Orl 80 mg

Micardis	02240770	BOE	ACDEFGV
Auro-Telmisartan	02453576	ARO	ACDEFGV
Jamp Telmisartan	02386763	JPC	ACDEFGV
Mint-Telmisartan	02486377	MNT	ACDEFGV
NRA-Telmisartan	02503808	NRA	ACDEFGV
pms-Telmisartan	02499630	PMS	ACDEFGV
Sandoz Telmisartan	02375966	SDZ	ACDEFGV
Telmisartan	02407493	AHI	ACDEFGV
Telmisartan	02388952	SAS	ACDEFGV
Telmisartan	02390353	SIV	ACDEFGV
Teva-Telmisartan	02320185	TEV	ACDEFGV

## C09CA08 OLMESARTAN MEDOXOMIL



C09CA08 OLMESARTAN MEDOXOMIL

Tab Orl 20 mg

Olmetec	02318660	ORG	ACDEFGV
Ach-Olmesartan	02456311	AHI	ACDEFGV
Act Olmesartan	02442191	TEV	ACDEFGV
Apo-Olmesartan	02453452	APX	ACDEFGV
Auro-Olmesartan	02443864	ARO	ACDEFGV
GLN-Olmesartan	02469812	GLM	ACDEFGV
Jamp-Olmesartan	02461641	JPC	ACDEFGV
NRA-Olmesartan	02499258	NRA	ACDEFGV
Olmesartan	02481057	SAS	ACDEFGV
pms-Olmesartan	02461307	PMS	ACDEFGV
Sandoz Olmesartan	02443414	SDZ	ACDEFGV

Tab Orl 40 mg

Olmetec	02318679	ORG	ACDEFGV
Ach-Olmesartan	02456338	AHI	ACDEFGV
Act Olmesartan	02442205	TEV	ACDEFGV
Apo-Olmesartan	02453460	APX	ACDEFGV
Auro-Olmesartan	02443872	ARO	ACDEFGV
GLN-Olmesartan	02469820	GLM	ACDEFGV
Jamp-Olmesartan	02461668	JPC	ACDEFGV
NRA-Olmesartan	02499266	NRA	ACDEFGV
Olmesartan	02481065	SAS	ACDEFGV
pms-Olmesartan	02461315	PMS	ACDEFGV
Sandoz Olmesartan	02443422	SDZ	ACDEFGV

**C09D ANGIOTENSIN II ANTAGONISTS, COMBINATIONS**

**C09DA ANGIOTENSIN II ANTAGONISTS AND DIURETICS**

C09DA01 LOSARTAN AND DIURETICS

LOSARTAN / HYDROCHLOROTHIAZIDE

Tab Orl 50 mg / 12.5 mg

Hyzaar	02230047	ORG	ACDEFGV
Auro-Losartan HCT	02423642	ARO	ACDEFGV
Losartan HCT	02388960	SIV	ACDEFGV
Losartan/HCTZ	02427648	SAS	ACDEFGV
Mint-Losartan/HCTZ	02389657	MNT	ACDEFGV
pms-Losartan-HCTZ	02392224	PMS	ACDEFGV
Sandoz Losartan HCT	02313375	SDZ	ACDEFGV
Teva-Losartan HCTZ	02358263	TEV	ACDEFGV

C09DA01		LOSARTAN AND DIURETICS							
		LOSARTAN / HYDROCHLOROTHIAZIDE							
Tab	Orl	100 mg / 12.5 mg		Hyzaar	02297841	ORG	ACDEFGV		
				Auro-Losartan HCT	02423650	ARO	ACDEFGV		
				Losartan HCT	02388979	SIV	ACDEFGV		
				Losartan/HCTZ	02427656	SAS	ACDEFGV		
				Mint-Losartan/HCTZ	02389665	MNT	ACDEFGV		
				pms-Losartan-HCTZ	02392232	PMS	ACDEFGV		
				Sandoz Losartan HCT	02362449	SDZ	ACDEFGV		
				Teva-Losartan HCTZ	02377144	TEV	ACDEFGV		
Tab	Orl	100 mg / 25 mg		Hyzaar DS	02241007	ORG	ACDEFGV		
				Auro-Losartan HCT	02423669	ARO	ACDEFGV		
				Losartan HCT	02388987	SIV	ACDEFGV		
				Losartan/HCTZ	02427664	SAS	ACDEFGV		
				Mint-Losartan/HCTZ DS	02389673	MNT	ACDEFGV		
				pms-Losartan-HCTZ	02392240	PMS	ACDEFGV		
				Sandoz Losartan HCT	02313383	SDZ	ACDEFGV		
				Teva-Losartan HCTZ	02377152	TEV	ACDEFGV		
C09DA02		EPROSARTAN AND DIURETICS							
		EPROSARTAN / HYDROCHLOROTHIAZIDE							
Tab	Orl	600 mg / 12.5 mg		Teveten Plus	02253631	BGP	ACDEFGV		
C09DA03		VALSARTAN AND DIURETICS							
		VALSARTAN / HYDROCHLOROTHIAZIDE							
Tab	Orl	80 mg / 12.5 mg		Diovan HCT	02241900	NVR	ACDEFGV		
				Auro-Valsartan HCT	02408112	ARO	ACDEFGV		
				Sandoz Valsartan HCT	02356694	SDZ	ACDEFGV		
				Teva-Valsartan/ HCTZ	02356996	TEV	ACDEFGV		
				Valsartan HCT	02384736	SIV	ACDEFGV		
				Valsartan/HCTZ	02367009	SAS	ACDEFGV		
Tab	Orl	160 mg / 12.5 mg		Diovan HCT	02241901	NVR	ACDEFGV		
				Auro-Valsartan HCT	02408120	ARO	ACDEFGV		
				Sandoz Valsartan HCT	02356708	SDZ	ACDEFGV		
				Teva-Valsartan/ HCTZ	02357003	TEV	ACDEFGV		
				Valsartan HCT	02384744	SIV	ACDEFGV		
				Valsartan/HCTZ	02367017	SAS	ACDEFGV		

C09DA03 VALSARTAN AND DIURETICS  
 VALSARTAN / HYDROCHLOROTHIAZIDE

Tab	Orl	160 mg / 25 mg	Diovan HCT	02246955	NVR	ACDEFGV
			Auro-Valsartan HCT	02408139	ARO	ACDEFGV
			Sandoz Valsartan HCT	02356716	SDZ	ACDEFGV
			Teva-Valsartan/ HCTZ	02357011	TEV	ACDEFGV
			Valsartan HCT	02384752	SIV	ACDEFGV
			Valsartan/HCTZ	02367025	SAS	ACDEFGV

Tab	Orl	320 mg / 12.5 mg	Diovan HCT	02308908	NVR	ACDEFGV
			Auro-Valsartan HCT	02408147	ARO	ACDEFGV
			Sandoz Valsartan HCT	02356724	SDZ	ACDEFGV
			Teva-Valsartan/ HCTZ	02357038	TEV	ACDEFGV
			Valsartan HCT	02384760	SIV	ACDEFGV
			Valsartan/HCTZ	02367033	SAS	ACDEFGV

Tab	Orl	320 mg / 25 mg	Diovan HCT	02308916	NVR	ACDEFGV
			Auro-Valsartan HCT	02408155	ARO	ACDEFGV
			Sandoz Valsartan HCT	02356732	SDZ	ACDEFGV
			Teva-Valsartan/ HCTZ	02357046	TEV	ACDEFGV
			Valsartan/HCTZ	02367041	SAS	ACDEFGV

C09DA04 IRBESARTAN AND DIURETICS  
 IRBESARTAN / HYDROCHLOROTHIAZIDE

Tab	Orl	150 mg / 12.5 mg	Avalide	02241818	SAV	ACDEFGV
			Auro-Irbesartan HCT	02447878	ARO	ACDEFGV
			Irbesartan HCT	02385317	SIV	ACDEFGV
			Irbesartan/HCTZ	02372886	SAS	ACDEFGV
			pms-Irbesartan HCTZ	02328518	PMS	ACDEFGV
			Sandoz Irbesartan HCT	02337428	SDZ	ACDEFGV
			Teva-Irbesartan HCTZ	02330512	TEV	ACDEFGV

Tab	Orl	300 mg / 12.5 mg	Avalide	02241819	SAV	ACDEFGV
			Auro-Irbesartan HCT	02447886	ARO	ACDEFGV
			Irbesartan HCT	02385325	SIV	ACDEFGV
			Irbesartan/HCTZ	02372894	SAS	ACDEFGV
			pms-Irbesartan HCTZ	02328526	PMS	ACDEFGV
			Sandoz Irbesartan HCT	02337436	SDZ	ACDEFGV
			Teva-Irbesartan HCTZ	02330520	TEV	ACDEFGV

C09DA04 IRBESARTAN AND DIURETICS  
 IRBESARTAN / HYDROCHLOROTHIAZIDE

Tab	Orl	300 mg / 25 mg	Auro-Irbesartan HCT	02447894	ARO	ACDEFGV
			Irbesartan HCT	02385333	SIV	ACDEFGV
			Irbesartan/HCTZ	02372908	SAS	ACDEFGV
			pms-Irbesartan HCTZ	02328534	PMS	ACDEFGV
			Sandoz Irbesartan HCT	02337444	SDZ	ACDEFGV
			Teva-Irbesartan HCTZ	02330539	TEV	ACDEFGV

C09DA06 CANDESARTAN AND DIURETICS  
 CANDESARTAN / HYDROCHLOROTHIAZIDE

Tab	Orl	16 mg / 12.5 mg	Atacand Plus	02244021	AZE	ACDEFGV
			Auro-Candesartan HCT	02421038	ARO	ACDEFGV
			Candesartan HCT	02394812	SIV	ACDEFGV
			Candesartan/HCTZ	02394804	SAS	ACDEFGV
			Jamp-Candesartan HCT	02473240	JPC	ACDEFGV
			pms-Candesartan-HCTZ	02391295	PMS	ACDEFGV
			Sandoz Candesartan Plus	02327902	SDZ	ACDEFGV
			Teva-Candesartan/HCTZ	02395541	TEV	ACDEFGV

Tab	Orl	32 mg / 12.5 mg	Atacand Plus	02332922	AZE	ACDEFGV
			Auro-Candesartan HCT	02421046	ARO	ACDEFGV
			Jamp-Candesartan HCT	02473259	JPC	ACDEFGV
			Sandoz Candesartan Plus	02420732	SDZ	ACDEFGV
			Teva-Candesartan/HCTZ	02395568	TEV	ACDEFGV

Tab	Orl	32 mg / 25 mg	Atacand Plus	02332957	AZE	ACDEFGV
			Auro-Candesartan HCT	02421054	ARO	ACDEFGV
			Jamp-Candesartan HCT	02473267	JPC	ACDEFGV
			Sandoz Candesartan Plus	02420740	SDZ	ACDEFGV

C09DA07 TELMISARTAN AND DIURETICS  
 TELMISARTAN / HYDROCHLOROTHIAZIDE

C09DA07 TELMISARTAN AND DIURETICS  
 TELMISARTAN / HYDROCHLOROTHIAZIDE

Tab	Orl	80 mg / 12.5 mg	Micardis Plus	02244344	BOE	ACDEFGV
			ACH-Telmisartan HCTZ	02419114	AHI	ACDEFGV
			Auro-Telmisartan HCTZ	02456389	ARO	ACDEFGV
			Jamp Telmisartan-HCT	02389940	JPC	ACDEFGV
			NRA-Telmisartan HCTZ	02504146	NRA	ACDEFGV
			Sandoz Telmisartan HCT	02393557	SDZ	ACDEFGV
			Telmisartan HCTZ	02390302	SIV	ACDEFGV
			Telmisartan/HCTZ	02395355	SAS	ACDEFGV
			Teva-Telmisartan HCTZ	02330288	TEV	ACDEFGV

Tab	Orl	80 mg / 25 mg	Micardis Plus	02318709	BOE	ACDEFGV
			ACH-Telmisartan HCTZ	02419122	AHI	ACDEFGV
			Auro-Telmisartan HCTZ	02456397	ARO	ACDEFGV
			Jamp Telmisartan-HCT	02389959	JPC	ACDEFGV
			NRA-Telmisartan HCTZ	02504138	NRA	ACDEFGV
			Sandoz Telmisartan HCT	02393565	SDZ	ACDEFGV
			Telmisartan HCTZ	02390310	SIV	ACDEFGV
			Telmisartan/HCTZ	02395363	SAS	ACDEFGV
			Teva-Telmisartan HCTZ	02379252	TEV	ACDEFGV

C09DA08 OLMESARTAN AND DIURETICS  
 OLMESARTAN / HYDROCHLOROTHIAZIDE

Tab	Orl	20 mg / 12.5 mg	Olmetec Plus	02319616	ORG	ACDEFGV
			ACH-Olmesartan HCTZ	02468948	AHI	ACDEFGV
			Act Olmesartan HCT	02443112	TEV	ACDEFGV
			Apo-Olmesartan/HCTZ	02453606	APX	ACDEFGV
			Auro-Olmesartan HCTZ	02476487	ARO	ACDEFGV
			GLN-Olmesartan HCTZ	02475707	GLM	ACDEFGV
			NRA-Olmesartan HCTZ	02508273	NRA	ACDEFGV
			Olmesartan/HCTZ	02509601	SAS	ACDEFGV
			PRZ-Olmesartan/HCTZ	02526468	PRZ	ACDEFGV

C09DA08 OLMESARTAN AND DIURETICS  
 OLMESARTAN / HYDROCHLOROTHIAZIDE

Tab	Orl	40 mg / 12.5 mg	Olmetec Plus	02319624	ORG	ACDEFGV
			ACH-Olmesartan HCTZ	02468956	AHI	ACDEFGV
			Act Olmesartan HCT	02443120	TEV	ACDEFGV
			Apo-Olmesartan/HCTZ	02453614	APX	ACDEFGV
			Auro-Olmesartan HCTZ	02476495	ARO	ACDEFGV
			GLN-Olmesartan HCTZ	02475715	GLM	ACDEFGV
			NRA-Olmesartan HCTZ	02508281	NRA	ACDEFGV
			Olmesartan/HCTZ	02509636	SAS	ACDEFGV
			PRZ-Olmesartan/HCTZ	02526476	PRZ	ACDEFGV

Tab	Orl	40 mg / 25 mg	Olmetec Plus	02319632	ORG	ACDEFGV
			ACH-Olmesartan HCTZ	02468964	AHI	ACDEFGV
			Act Olmesartan HCT	02443139	TEV	ACDEFGV
			Apo-Olmesartan/HCTZ	02453622	APX	ACDEFGV
			Auro-Olmesartan HCTZ	02476509	ARO	ACDEFGV
			GLN-Olmesartan HCTZ	02475723	GLM	ACDEFGV
			NRA-Olmesartan HCTZ	02508303	NRA	ACDEFGV
			Olmesartan/HCTZ	02509628	SAS	ACDEFGV
			PRZ-Olmesartan/HCTZ	02526484	PRZ	ACDEFGV

**C09DB ANGIOTENSIN II ANTAGONISTS AND CALCIUM CHANNEL BLOCKERS**

C09DB04 TELMISARTAN AND AMLODIPINE

Tab	Orl	40 mg / 5 mg	Twynsta	02371022	BOE	ACDEFGV
Tab	Orl	40 mg / 10 mg	Twynsta	02371030	BOE	ACDEFGV
Tab	Orl	80 mg / 5 mg	Twynsta	02371049	BOE	ACDEFGV
Tab	Orl	80 mg / 10 mg	Twynsta	02371057	BOE	ACDEFGV

**C09DX ANGIOTENSIN II ANTAGONISTS, OTHER COMBINATIONS**

C09DX04 VALSARTAN AND SACUBITRIL

Tab	Orl	26 mg / 24 mg	Entresto	02446928	NVR	(SA)
Tab	Orl	51 mg / 49 mg	Entresto	02446936	NVR	(SA)
Tab	Orl	103 mg / 97 mg	Entresto	02446944	NVR	(SA)

**C10 LIPID MODIFYING AGENTS**

**C10A LIPID MODIFYING AGENTS, PLAIN**

**C10AA HMG COA REDUCTASE INHIBITORS**

C10AA01 SIMVASTATIN

Tab Orl 5 mg

Apo-Simvastatin 02247011 APX ACDEFGV  
 Auro-Simvastatin 02405148 ARO ACDEFGV  
 Jamp-Simvastatin 02375591 JPC ACDEFGV  
 Mint-Simvastatin 02372932 MNT ACDEFGV  
 pharma-Simvastatin 02469979 PMS ACDEFGV  
 Simvastatin 02284723 SAS ACDEFGV  
 Simvastatin 02386291 SIV ACDEFGV  
 Taro-Simvastatin 02329131 SUN ACDEFGV  
 Teva-Simvastatin 02250144 TEV ACDEFGV

Tab Orl 10 mg

Zocor 00884332 ORG ACDEFGV  
 Apo-Simvastatin 02247012 APX ACDEFGV  
 Auro-Simvastatin 02405156 ARO ACDEFGV  
 Jamp-Simvastatin 02375605 JPC ACDEFGV  
 Mar-Simvastatin 02375044 MAR ACDEFGV  
 Mint-Simvastatin 02372940 MNT ACDEFGV  
 pharma-Simvastatin 02469987 PMS ACDEFGV  
 Simvastatin 02284731 SAS ACDEFGV  
 Simvastatin 02386305 SIV ACDEFGV  
 Simvastatin-10 02247221 PDL ACDEFGV  
 Taro-Simvastatin 02329158 SUN ACDEFGV  
 Teva-Simvastatin 02250152 TEV ACDEFGV

Tab Orl 20 mg

Zocor 00884340 ORG ACDEFGV  
 Apo-Simvastatin 02247013 APX ACDEFGV  
 Auro-Simvastatin 02405164 ARO ACDEFGV  
 Jamp-Simvastatin 02375613 JPC ACDEFGV  
 Mar-Simvastatin 02375052 MAR ACDEFGV  
 Mint-Simvastatin 02372959 MNT ACDEFGV  
 pharma-Simvastatin 02469995 PMS ACDEFGV  
 Simvastatin 02284758 SAS ACDEFGV  
 Simvastatin 02386313 SIV ACDEFGV  
 Simvastatin-20 02247222 PDL ACDEFGV  
 Taro-Simvastatin 02329166 SUN ACDEFGV  
 Teva-Simvastatin 02250160 TEV ACDEFGV

C10AA01 SIMVASTATIN

Tab Orl 40 mg

Zocor 00884359 ORG ACDEFGV  
 Apo-Simvastatin 02247014 APX ACDEFGV  
 Auro-Simvastatin 02405172 ARO ACDEFGV  
 Jamp-Simvastatin 02375621 JPC ACDEFGV  
 Mar-Simvastatin 02375060 MAR ACDEFGV  
 Mint-Simvastatin 02372967 MNT ACDEFGV  
 pharma-Simvastatin 02470004 PMS ACDEFGV  
 Simvastatin 02284766 SAS ACDEFGV  
 Simvastatin 02386321 SIV ACDEFGV  
 Simvastatin-40 02247223 PDL ACDEFGV  
 Taro-Simvastatin 02329174 SUN ACDEFGV  
 Teva-Simvastatin 02250179 TEV ACDEFGV

Tab Orl 80 mg

Apo-Simvastatin 02247015 APX ACDEFGV  
 Auro-Simvastatin 02405180 ARO ACDEFGV  
 Jamp-Simvastatin 02375648 JPC ACDEFGV  
 Mint-Simvastatin 02372975 MNT ACDEFGV  
 pharma-Simvastatin 02470012 PMS ACDEFGV  
 Simvastatin 02284774 SAS ACDEFGV  
 Simvastatin 02386348 SIV ACDEFGV  
 Simvastatin-80 02247224 PDL ACDEFGV  
 Taro-Simvastatin 02329182 SUN ACDEFGV  
 Teva-Simvastatin 02250187 TEV ACDEFGV

C10AA02 LOVASTATIN

Tab Orl 20 mg

Act Lovastatin 02248572 TEV ACDEFGV  
 Lovastatin 02220172 AAP ACDEFGV

Tab Orl 40 mg

Act Lovastatin 02248573 TEV ACDEFGV  
 Lovastatin 02220180 AAP ACDEFGV

C10AA03 PRAVASTATIN



C10AA03 PRAVASTATIN

Tab Orl 10 mg

Ach-Pravastatin	02440644	AHI	ACDEFGV
Apo-Pravastatin	02243506	APX	ACDEFGV
Auro-Pravastatin	02458977	ARO	ACDEFGV
Jamp-Pravastatin	02330954	JPC	ACDEFGV
M-Pravastatin	02476274	MRA	ACDEFGV
Mar-Pravastatin	02432048	MAR	ACDEFGV
Mint-Pravastatin	02317451	MNT	ACDEFGV
pms-Pravastatin	02247655	PMS	ACDEFGV
Pravastatin	02356546	SAS	ACDEFGV
Pravastatin	02389703	SIV	ACDEFGV
Pravastatin-10	02243824	PDL	ACDEFGV
Sandoz Pravastatin	02468700	SDZ	ACDEFGV
Taro-Pravastatin	02284421	SUN	ACDEFGV
Teva-Pravastatin	02247008	TEV	ACDEFGV

Tab Orl 20 mg

Ach-Pravastatin	02440652	AHI	ACDEFGV
Apo-Pravastatin	02243507	APX	ACDEFGV
Auro-Pravastatin	02458985	ARO	ACDEFGV
Jamp-Pravastatin	02330962	JPC	ACDEFGV
M-Pravastatin	02476282	MRA	ACDEFGV
Mar-Pravastatin	02432056	MAR	ACDEFGV
Mint-Pravastatin	02317478	MNT	ACDEFGV
pms-Pravastatin	02247656	PMS	ACDEFGV
Pravastatin	02356554	SAS	ACDEFGV
Pravastatin	02389738	SIV	ACDEFGV
Pravastatin-20	02243825	PDL	ACDEFGV
Sandoz Pravastatin	02468719	SDZ	ACDEFGV
Taro-Pravastatin	02284448	SUN	ACDEFGV
Teva-Pravastatin	02247009	TEV	ACDEFGV

C10AA03 PRAVASTATIN

Tab Orl 40 mg

Ach-Pravastatin 02440660 AHI ACDEFGV  
Apo-Pravastatin 02243508 APX ACDEFGV  
Auro-Pravastatin 02458993 ARO ACDEFGV  
Jamp-Pravastatin 02330970 JPC ACDEFGV  
M-Pravastatin 02476290 MRA ACDEFGV  
Mar-Pravastatin 02432064 MAR ACDEFGV  
Mint-Pravastatin 02317486 MNT ACDEFGV  
pms-Pravastatin 02247657 PMS ACDEFGV  
Pravastatin 02356562 SAS ACDEFGV  
Pravastatin 02389746 SIV ACDEFGV  
Pravastatin-40 02243826 PDL ACDEFGV  
Sandoz Pravastatin 02468727 SDZ ACDEFGV  
Taro-Pravastatin 02284456 SUN ACDEFGV  
Teva-Pravastatin 02247010 TEV ACDEFGV

C10AA04 FLUVASTATIN

Cap Orl 20 mg

Teva-Fluvastatin 02299224 TEV ACDEFGV

Cap Orl 40 mg

Teva-Fluvastatin 02299232 TEV ACDEFGV

C10AA05 ATORVASTATIN

## C10AA05 ATORVASTATIN

Tab Orl 10 mg

Lipitor	02230711	BGP	ACDEFGV
ACH-Atorvastatin Calcium	02457741	AHI	ACDEFGV
Apo-Atorvastatin	02295261	APX	ACDEFGV
Atorvastatin	02346486	PDL	ACDEFGV
Atorvastatin	02475022	RIV	ACDEFGV
Atorvastatin	02348705	SAS	ACDEFGV
Atorvastatin	02411350	SIV	ACDEFGV
Auro-Atorvastatin	02407256	ARO	ACDEFGV
Jamp Atorvastatin Calcium	02504197	JPC	ACDEFGV
Jamp-Atorvastatin	02391058	JPC	ACDEFGV
M-Atorvastatin	02471167	MRA	ACDEFGV
Mar-Atorvastatin	02454017	MAR	ACDEFGV
Mint-Atorvastatin	02479508	MNT	ACDEFGV
Mylan-Atorvastatin	02392933	MYL	ACDEFGV
NRA-Atorvastatin	02476517	NRA	ACDEFGV
pms-Atorvastatin	02477149	PMS	ACDEFGV
pmsc-Atorvastatin	02507234	PMS	ACDEFGV
PRZ-Atorvastatin	02521555	PRZ	ACDEFGV
Reddy-Atorvastatin	02417936	RCH	ACDEFGV
Sandoz Atorvastatin	02324946	SDZ	ACDEFGV
Taro-Atorvastatin	02313707	SUN	ACDEFGV
Teva-Atorvastatin	02310899	TEV	ACDEFGV

## C10AA05 ATORVASTATIN

Tab Orl 20 mg

Lipitor	02230713	BGP	ACDEFGV
ACH-Atorvastatin Calcium	02457768	AHI	ACDEFGV
Apo-Atorvastatin	02295288	APX	ACDEFGV
Atorvastatin	02346494	PDL	ACDEFGV
Atorvastatin	02475030	RIV	ACDEFGV
Atorvastatin	02348713	SAS	ACDEFGV
Atorvastatin	02411369	SIV	ACDEFGV
Auro-Atorvastatin	02407264	ARO	ACDEFGV
Jamp Atorvastatin Calcium	02504200	JPC	ACDEFGV
Jamp-Atorvastatin	02391066	JPC	ACDEFGV
M-Atorvastatin	02471175	MRA	ACDEFGV
Mar-Atorvastatin	02454025	MAR	ACDEFGV
Mint-Atorvastatin	02479516	MNT	ACDEFGV
Mylan-Atorvastatin	02392941	MYL	ACDEFGV
NRA-Atorvastatin	02476525	NRA	ACDEFGV
pms-Atorvastatin	02477157	PMS	ACDEFGV
pmsc-Atorvastatin	02507242	PMS	ACDEFGV
PRZ-Atorvastatin	02521563	PRZ	ACDEFGV
Reddy-Atorvastatin	02417944	RCH	ACDEFGV
Sandoz Atorvastatin	02324954	SDZ	ACDEFGV
Taro-Atorvastatin	02313715	SUN	ACDEFGV
Teva-Atorvastatin	02310902	TEV	ACDEFGV

## C10AA05 ATORVASTATIN

Tab Orl 40 mg

Lipitor	02230714	BGP	ACDEFGV
ACH-Atorvastatin Calcium	02457776	AHI	ACDEFGV
Apo-Atorvastatin	02295296	APX	ACDEFGV
Atorvastatin	02346508	PDL	ACDEFGV
Atorvastatin	02475049	RIV	ACDEFGV
Atorvastatin	02348721	SAS	ACDEFGV
Atorvastatin	02411377	SIV	ACDEFGV
Auro-Atorvastatin	02407272	ARO	ACDEFGV
Jamp Atorvastatin Calcium	02504219	JPC	ACDEFGV
Jamp-Atorvastatin	02391074	JPC	ACDEFGV
M-Atorvastatin	02471183	MRA	ACDEFGV
Mar-Atorvastatin	02454033	MAR	ACDEFGV
Mint-Atorvastatin	02479524	MNT	ACDEFGV
Mylan-Atorvastatin	02392968	MYL	ACDEFGV
NRA-Atorvastatin	02476533	NRA	ACDEFGV
pms-Atorvastatin	02477165	PMS	ACDEFGV
pmsc-Atorvastatin	02507250	PMS	ACDEFGV
PRZ-Atorvastatin	02521571	PRZ	ACDEFGV
Reddy-Atorvastatin	02417952	RCH	ACDEFGV
Sandoz Atorvastatin	02324962	SDZ	ACDEFGV
Taro-Atorvastatin	02313723	SUN	ACDEFGV
Teva-Atorvastatin	02310910	TEV	ACDEFGV

## C10AA05 ATORVASTATIN

Tab Orl 80 mg

Lipitor	02243097	BGP	ACDEFGV
ACH-Atorvastatin Calcium	02457784	AHI	ACDEFGV
Apo-Atorvastatin	02295318	APX	ACDEFGV
Atorvastatin	02346516	PDL	ACDEFGV
Atorvastatin	02475057	RIV	ACDEFGV
Atorvastatin	02348748	SAS	ACDEFGV
Atorvastatin	02411385	SIV	ACDEFGV
Auro-Atorvastatin	02407280	ARO	ACDEFGV
Jamp Atorvastatin Calcium	02504235	JPC	ACDEFGV
Jamp-Atorvastatin	02391082	JPC	ACDEFGV
M-Atorvastatin	02471191	MRA	ACDEFGV
Mar-Atorvastatin	02454041	MAR	ACDEFGV
Mylan-Atorvastatin	02392976	MYL	ACDEFGV
NRA-Atorvastatin	02476541	NRA	ACDEFGV
pms-Atorvastatin	02477173	PMS	ACDEFGV
pmsc-Atorvastatin	02507269	PMS	ACDEFGV
PRZ-Atorvastatin	02521598	PRZ	ACDEFGV
Reddy-Atorvastatin	02417960	RCH	ACDEFGV
Sandoz Atorvastatin	02324970	SDZ	ACDEFGV
Taro-Atorvastatin	02313758	SUN	ACDEFGV
Teva-Atorvastatin	02310929	TEV	ACDEFGV

## C10AA07 ROSUVASTATIN

Tab Orl 5 mg

Crestor	02265540	AZE	ACDEFGV
ACH-Rosuvastatin	02438917	AHI	ACDEFGV
Apo-Rosuvastatin	02337975	APX	ACDEFGV
Auro-Rosuvastatin	02442574	ARO	ACDEFGV
Jamp Rosuvastatin Calcium	02498332	JPC	ACDEFGV
Jamp-Rosuvastatin	02391252	JPC	ACDEFGV
M-Rosuvastatin	02496534	MRA	ACDEFGV
NRA-Rosuvastatin	02477483	NRA	ACDEFGV
pms-Rosuvastatin	02378523	PMS	ACDEFGV
PRZ-Rosuvastatin	02505576	PRZ	ACDEFGV
Rosuvastatin	02381176	PDL	ACDEFGV
Rosuvastatin	02405628	SAS	ACDEFGV
Rosuvastatin	02411628	SIV	ACDEFGV
Sandoz Rosuvastatin	02338726	SDZ	ACDEFGV
Taro-Rosuvastatin	02382644	SUN	ACDEFGV
Teva-Rosuvastatin	02354608	TEV	ACDEFGV

C10AA07 ROSUVASTATIN

Tab Orl 10 mg

	Crestor	02247162	AZE	ACDEFGV
	ACH-Rosuvastatin	02438925	AHI	ACDEFGV
	Apo-Rosuvastatin	02337983	APX	ACDEFGV
	Auro-Rosuvastatin	02442582	ARO	ACDEFGV
Jamp	Rosuvastatin Calcium	02498340	JPC	ACDEFGV
	Jamp-Rosuvastatin	02391260	JPC	ACDEFGV
	M-Rosuvastatin	02496542	MRA	ACDEFGV
	NRA-Rosuvastatin	02477491	NRA	ACDEFGV
	pms-Rosuvastatin	02378531	PMS	ACDEFGV
	PRZ-Rosuvastatin	02505584	PRZ	ACDEFGV
	Rosuvastatin	02381184	PDL	ACDEFGV
	Rosuvastatin	02405636	SAS	ACDEFGV
	Rosuvastatin	02411636	SIV	ACDEFGV
Sandoz	Rosuvastatin	02338734	SDZ	ACDEFGV
	Taro-Rosuvastatin	02382652	SUN	ACDEFGV
	Teva-Rosuvastatin	02354616	TEV	ACDEFGV

Tab Orl 20 mg

	Crestor	02247163	AZE	ACDEFGV
	ACH-Rosuvastatin	02438933	AHI	ACDEFGV
	Apo-Rosuvastatin	02337991	APX	ACDEFGV
	Auro-Rosuvastatin	02442590	ARO	ACDEFGV
Jamp	Rosuvastatin Calcium	02498359	JPC	ACDEFGV
	Jamp-Rosuvastatin	02391279	JPC	ACDEFGV
	M-Rosuvastatin	02496550	MRA	ACDEFGV
	NRA-Rosuvastatin	02477505	NRA	ACDEFGV
	pms-Rosuvastatin	02378558	PMS	ACDEFGV
	PRZ-Rosuvastatin	02505592	PRZ	ACDEFGV
	Rosuvastatin	02381192	PDL	ACDEFGV
	Rosuvastatin	02405644	SAS	ACDEFGV
	Rosuvastatin	02411644	SIV	ACDEFGV
Sandoz	Rosuvastatin	02338742	SDZ	ACDEFGV
	Taro-Rosuvastatin	02382660	SUN	ACDEFGV
	Teva-Rosuvastatin	02354624	TEV	ACDEFGV

C10AA07 ROSUVASTATIN

Tab Orl 40 mg

Crestor	02247164	AZE	ACDEFGV
ACH-Rosuvastatin	02438941	AHI	ACDEFGV
Apo-Rosuvastatin	02338009	APX	ACDEFGV
Auro-Rosuvastatin	02442604	ARO	ACDEFGV
Jamp Rosuvastatin Calcium	02498367	JPC	ACDEFGV
Jamp-Rosuvastatin	02391287	JPC	ACDEFGV
M-Rosuvastatin	02496569	MRA	ACDEFGV
NRA-Rosuvastatin	02477513	NRA	ACDEFGV
pms-Rosuvastatin	02378566	PMS	ACDEFGV
PRZ-Rosuvastatin	02505606	PRZ	ACDEFGV
Rosuvastatin	02381206	PDL	ACDEFGV
Rosuvastatin	02405652	SAS	ACDEFGV
Rosuvastatin	02411652	SIV	ACDEFGV
Sandoz Rosuvastatin	02338750	SDZ	ACDEFGV
Taro-Rosuvastatin	02382679	SUN	ACDEFGV
Teva-Rosuvastatin	02354632	TEV	ACDEFGV

**C10AB FIBRATES**

C10AB04 GEMFIBROZIL

Cap Orl 300 mg

pms-Gemfibrozil 02239951 PMS ACDEFGV

Tab Orl 600 mg

Teva-Gemfibrozil 02142074 TEV ACDEFGV

C10AB05 FENOFIBRATE

Cap Orl 67 mg

AA-Feno Micro 02243180 AAP ACDEFGV

Cap Orl 200 mg

AA-Feno-Micro 02239864 AAP ACDEFGV

Tab Orl 48 mg

Lipidil EZ 02269074 BGP ACDEFGV  
Sandoz Fenofibrate E 02390698 SDZ ACDEFGV

Tab Orl 100 mg

AA-Feno-Super 02246859 AAP ACDEFGV

Tab Orl 145 mg

Lipidil EZ 02269082 BGP ACDEFGV  
Sandoz Fenofibrate E 02390701 SDZ ACDEFGV  
Taro-Fenofibrate E 02454696 SUN ACDEFGV

Tab Orl 160 mg

Lipidil Supra 02241602 BGP ACDEFGV  
AA-Feno-Super 02246860 AAP ACDEFGV

**C10AC BILE ACID SEQUESTRANTS**



C10AC01	CHOLESTYRAMINE						
Pws	Orl	4 g					
			Cholestyramine-Odan	02455609	ODN	ACDEFGV	
			Jamp-Cholestyramine	02478595	JPC	ACDEFGV	

C10AC02	COLESTIPOL						
Pws	Orl	5 g					
			Colestid	00642975	PFI	ACDEFGV	
Tab	Orl	1 mg					
			Colestid	02132680	PFI	ACDEFGV	

C10AC04	COLESEVELAM						
Pws	Orl	3.75 g					
			Lodalis	02432463	BSL	ACDEFGV	
Tab	Orl	625 mg					
			Lodalis	02373955	BSL	ACDEFGV	
			Apo-Colesevelam	02494051	APX	ACDEFGV	

**C10AX OTHER LIPID MODIFYING AGENTS**

C10AX06	OMEGA-3-TRIGLYCERIDES INCL. OTHER ESTERS AND ACIDS						
	ICOSAPENT ETHYL						
Cap	Orl	1 g					
			Vascepa	02495244	HLS	(SA)	

C10AX09	EZETIMIBE						
Tab	Orl	10 mg					
			Ezetrol	02247521	ORG	ACDEFGV	
			ACH-Ezetimibe	02425610	AHI	ACDEFGV	
			Apo-Ezetimibe	02427826	APX	ACDEFGV	
			Auro-Ezetimibe	02469286	ARO	ACDEFGV	
			Ezetimibe	02422549	PDL	ACDEFGV	
			Ezetimibe	02431300	SAS	ACDEFGV	
			Ezetimibe	02429659	SIV	ACDEFGV	
			GLN-Ezetimibe	02460750	GLM	ACDEFGV	
			Jamp-Ezetimibe	02423235	JPC	ACDEFGV	
			M-Ezetimibe	02467437	MRA	ACDEFGV	
			Mar-Ezetimibe	02422662	MAR	ACDEFGV	
			Mint-Ezetimibe	02423243	MNT	ACDEFGV	
			NRA-Ezetimibe	02481669	NRA	ACDEFGV	
			pms-Ezetimibe	02416409	PMS	ACDEFGV	
			Ran-Ezetimibe	02419548	RAN	ACDEFGV	
			Sandoz Ezetimibe	02416778	SDZ	ACDEFGV	
			Teva-Ezetimibe	02354101	TEV	ACDEFGV	

C10AX13	EVOLOCUMAB						
Liq	SC	120 mg/mL					
			Repatha (prefilled mini-doser)	02459779	AGA	(SA)	

C10AX13	EVOLOCUMAB								
	Liq	SC	140 mg/mL					Repatha (autoinjector)	02446057 AGA (SA)
C10AX14	ALIROCUMAB								
	Liq	SC	75 mg/mL					Praluent (prefilled pen)	02453819 SAV (SA)
	Liq	SC	150 mg/mL					Praluent (prefilled pen)	02453835 SAV (SA)
<b>D</b>	<b>DERMATOLOGICALS</b>								
<b>D01</b>	<b>ANTIFUNGALS FOR DERMATOLOGICAL USE</b>								
<b>D01A</b>	<b>ANTIFUNGALS FOR TOPICAL USE</b>								
<b>D01AA</b>	<b>ANTIBIOTICS</b>								
D01AA01	NYSTATIN								
	Crm	Top	100 000 IU					Nyaderm	00716871 TAR ACDEFGV
<b>D01AC</b>	<b>IMIDAZOLE AND TRIAZOLE DERIVATIVES</b>								
D01AC01	CLOTRIMAZOLE								
	Crm	Top	1%					Canesten	02150867 BAY ACDEFGV
								Clotrimaderm	00812382 TAR ACDEFGV
D01AC02	MICONAZOLE								
	Crm	Top	2%					Micatin	02085852 WLS ACDEFGV
								Monistat Derm	02126567 INP ACDEFGV
D01AC08	KETOCONAZOLE								
	Crm	Top	2%					Ketoderm	02245662 TPH ACDEFGV
D01AC20	IMIDAZOLES AND TRIAZOLES IN COMBINATION WITH CORTICOSTEROIDS								
	CLOTRIMAZOLE / BETAMETHASONE								
	Crm	Top	1% / 0.05%					Lotriderm	00611174 ORG ACDEFGV
								Taro-Clotrimazole/Betamethasone Dipropionate	02496410 TAR ACDEFGV
<b>D01AE</b>	<b>OTHER ANTIFUNGALS FOR TOPICAL USE</b>								
D01AE14	CICLOPIROX								
	Crm	Top	1%					Loprox	02221802 BSL ACDEFGV
	Lot	Top	1%					Loprox	02221810 BSL ACDEFGV
D01AE15	TERBINAFINE								
	Crm	Top	1%					Lamisil	02031094 NVR ACDEFGV
	Spr	Top	1%					Lamisil	02238703 NVR ACDEFGV

**D01B ANTIFUNGALS, SYSTEMIC PREPARATIONS****D01BA ANTIFUNGALS FOR SYSTEMIC USE**

## D01BA02 TERBINAFINE

Tab Orl 250 mg

Lamisil 02031116 NVR ACDEFGV

Act Terbinafine 02254727 TEV ACDEFGV

Apo-Terbinafine 02239893 APX ACDEFGV

Auro-Terbinafine 02320134 ARO ACDEFGV

pms-Terbinafine 02294273 PMS ACDEFGV

Terbinafine 02353121 SAS ACDEFGV

Terbinafine 02385279 SIV ACDEFGV

**D04 ANTIPRURITICS, INCL. ANTIHISTAMINES, ANESTHETICS, ETC.****D04A ANTIPRURITICS, INCL. ANTIHISTAMINES, ANESTHETICS, ETC.****D04AB ANESTHETICS FOR TOPICAL USE**

## D04AB01 LIDOCAINE

Gel Top 2%

Lidodan Jelly 02143879 ODN ACDEFGV

Xylocaine Jelly 00001694 APN ACDEFGV

Xylocaine Jelly 00385484 APN ACDEFGV

Ont Top 5%

Xylocaine Ointment 5% 00001961 APN ACDEFGV

**D05 ANTIPSORIATICS****D05A ANTIPSORIATICS FOR TOPICAL USE****D05AA TARS**

## D05AA99 TARS

Liq Top 20%

Odans LCD 00358495 ODN ACDEFGV

**D05AX OTHER ANTIPSORIATICS FOR TOPICAL USE**

## D05AX02 CALCIPOTRIOL

Ont Top 50 mcg

Dovonex 01976133 LEO ACDEFV

## D05AX05 TAZAROTENE

HALOBÉTASOL PROPIONATE / TAZAROTÈNE

Lot Top 0.01% / 0.045%

Duobrii 02499967 BSL ACDEFGV

Crm Top 0.05%

Tazorac Cream (Disc/non disp Mar 6/24) 02243894 ALL (SA)

Gel Top 0.1%

Tazorac Gel (Disc/non disp Mar 6/24) 02230785 ALL (SA)

Lot Top 0.045%

Arazlo 02517868 BSL ACDEFGV

## D05AX52 CALCIPOTRIOL, COMBINATIONS

D05AX52 CALCIPOTRIOL, COMBINATIONS  
CALCIPOTRIOL / BETAMETHASONE

Aer	Top	50 mcg 0.5 mg	Enstilar	02457393	LEO	ACDEFGV
Gel	Top	50 mcg / 0.5 mg	Dovobet	02319012	LEO	ACDEFGV
			Taro-Calcipotriol/Betamethasone Gel	02525178	TAR	ACDEFGV
Ont	Top	50 mcg / 0.5 mg	Dovobet	02244126	LEO	ACDEFGV
			Teva-Betamethasone/Calcipotriol	02427419	TEV	ACDEFGV

**D05B ANTIPSORIATICS FOR SYSTEMIC USE**

**D05BB RETINOIDS FOR TREATMENT OF PSORIASIS**

D05BB02 ACITRETIN

Cap	Orl	10 mg	Soriatane	02070847	ALL	ACDEFGV
			Mint-Acitreten	02468840	MNT	ACDEFGV
			Taro-Acitreten	02466074	TAR	ACDEFGV
Cap	Orl	25 mg	Soriatane	02070863	ALL	ACDEFGV
			Mint-Acitreten	02468859	MNT	ACDEFGV
			Taro-Acitreten	02466082	TAR	ACDEFGV

**D06 ANTIBIOTICS AND CHEMOTHERAPEUTICS FOR DERMATOLOGICAL USE**

**D06A ANTIBIOTICS FOR TOPICAL USE**

**D06AX OTHER ANTIBIOTICS FOR TOPICAL USE**

D06AX01 FUSIDIC ACID

Crm	Top	2%	Fucidin	00586668	LEO	ACDEFGV
Ont	Top	2%	Fucidin	00586676	LEO	ACDEFGV

D06AX09 MUPIROCIN

Ont	Top	2%	Taro-Mupirocin	02279983	TAR	ACDEFGV
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**D06B CHEMOTHERAPEUTICS FOR TOPICAL USE**

**D06BA SULFONAMIDES**

D06BA01 SILVER SULFADIAZINE

Crm	Top	1%	Flamazine	00323098	SNE	ACDEFGVW
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**D06BB ANTIVIRALS**

D06BB10 IMIQUIMOD

Crm	Top	5%	Aldara P	02239505	BSL	(SA)
			Taro-Imiquimod Pump	02482983	TAR	(SA)

**D06BX OTHER CHEMOTHERAPEUTICS****D06BX01 METRONIDAZOLE**

Crm	Top	1%	Noritate	02156091	BSL	ACDEFGV
Gel	Top	1%	Metrogel	02297809	GAC	ACDEFGV

**D07 CORTICOSTEROIDS, DERMATOLOGICAL PREPARATIONS****D07A CORTICOSTEROIDS, PLAIN****D07AA CORTICOSTEROIDS, WEAK (GROUP I)****D07AA02 HYDROCORTISONE**

## HYDROCORTISONE ACETATE

Crm	Top	0.5%	Cortate	80021088	BAY	AEFGV
			Hydrosone	00564281	TEV	AEFGV
Crm	Top	1%	Hyderm	00716839	TAR	ACDEFGV
			Jamp-Hydrocortisone	80057189	JPC	ACDEFGV
			Jamp-Hydrocortisone Acetate	80057178	JPC	ACDEFGV
			Sandoz Hydrocortisone	02412926	SDZ	ACDEFGV

Lot	Top	1%	Jamp-Hydrocortisone	80057191	JPC	ACDEFGV
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Ont	Top	1%	Cortoderm	00716693	TAR	ACDEFGV
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## HYDROCORTISONE VALERATE

Crm	Top	0.2%	Hydroval	02242984	TPH	ACDEFGV
Ont	Top	0.2%	Hydroval	02242985	TPH	ACDEFGV

## HYDROCORTISONE/UREA

Crm	Top	1%	Dermaflex HC	00681989	PAL	ACDEFGV
			Jamp-Hydrocortisone Acetate-Urea	80061501	JPC	ACDEFGV
			M-HC 1% Urea 10%	80073645	MRA	ACDEFGV
Lot	Top	1%	Dermaflex HC	00681997	PAL	ACDEFGV

**D07AB CORTICOSTEROIDS, MODERATELY POTENT (GROUP II)****D07AB01 CLOBETASONE**

Crm	Top	0.05%	Spectro Eczemacare	02214415	GCH	AEFGV
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**D07AB08 DESONIDE**

Crm	Top	0.05%	pdp-Desonide	02229315	PDP	ACDEFGV
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D07AB08	DESONIDE						
Ont	Top	0.05%		pdp-Desonide	02229323	PDP	ACDEFGV
D07AB09	TRIAMCINOLONE						
Crm	Top	0.1%		Aristocort R	02194058	BSL	ACDEFGV
Crm	Top	0.5%		Aristocort C	02194066	BSL	ACDEFGV
Ont	Top	0.1%		Aristocort R	02194031	BSL	ACDEFGV
<b>D07AC</b>	<b>CORTICOSTEROIDS, POTENT (GROUP III)</b>						
D07AC01	BETAMETHASONE						
	BETAMETHASONE DIPROPIONATE						
Crm	Top	0.05%		Diprosone	00323071	ORG	ACDEFGV
				Teva-Topilene	00849650	TEV	ACDEFGV
				Teva-Topisone	00804991	TEV	ACDEFGV
Lot	Top	0.05%		Diprosone	00417246	ORG	ACDEFGV
				Teva-Topilene	01927914	TEV	ACDEFGV
				Teva-Topisone	00809187	TEV	ACDEFGV
Ont	Top	0.05%		Diprolene Glycol	00629367	ORG	ACDEFGV
				Diprosone	00344923	ORG	ACDEFGV
				Teva-Topilene Glycol	00849669	TEV	ACDEFGV
				Teva-Topisone	00805009	TEV	ACDEFGV
	BETAMETHASONE VALERATE						
Crm	Top	0.05%		Celestoderm V/2	02357860	BSL	ACDEFGV
				Betaderm	00716618	TAR	ACDEFGV
				Teva-Ectosone Mild	00535427	TEV	ACDEFGV
Crm	Top	0.1%		Betaderm	00716626	TAR	ACDEFGV
				Celestoderm V	02357844	PMS	ACDEFGV
				Teva-Ectosone	00535435	TEV	ACDEFGV
Lot	Top	0.05%		Teva-Ectosone Mild	00653209	TEV	ACDEFGV
Lot	Top	0.1%		Betaderm	00716634	TAR	ACDEFGV
				Teva-Ectosone	00750050	TEV	ACDEFGV
				Teva-Ectosone Scalp	00653217	TEV	ACDEFGV

D07AC01	BETAMETHASONE							
	BETAMETHASONE VALERATE							
Ont	Top	0.05%	Celestoderm V/2	02357879	BSL	ACDEFGV		
			Betaderm	00716642	TAR	ACDEFGV		
Ont	Top	0.1%	Celestoderm V	02357852	BSL	ACDEFGV		
			Betaderm	00716650	TAR	ACDEFGV		
D07AC03	DESOXIMETASONE							
Crm	Top	0.05%	Topicort Mild	02221918	BSL	ACDEFGV		
Crm	Top	0.25%	Topicort	02221896	BSL	ACDEFGV		
Gel	Top	0.05%	Topicort	02221926	BSL	ACDEFGV		
Ont	Top	0.25%	Topicort	02221934	BSL	ACDEFGV		
D07AC04	FLUOCINOLONE							
Liq	Top	0.01%	Derma-Smoothe	00873292	HLZ	ACDEFGV		
D07AC08	FLUOCINONIDE							
Crm	Top	0.05%	Lidemol	02163152	BSL	ACDEFGV		
			Lidex	02161923	BSL	ACDEFGV		
			Lyderm	00716863	TPH	ACDEFGV		
Gel	Top	0.05%	Lidex Gel	02161974	BSL	ACDEFGV		
			Lyderm	02236997	TPH	ACDEFGV		
Ont	Top	0.05%	Lidex	02161966	BSL	ACDEFGV		
			Lyderm	02236996	TPH	ACDEFGV		
D07AC11	AMCINONIDE							
Crm	Top	0.1%	Taro-Amcinonide	02246714	TAR	ACDEFGV		
D07AC13	MOMETASONE							
Crm	Top	0.1%	Elocom	00851744	ORG	ACDEFGV		
			Taro-Mometasone	02367157	TAR	ACDEFGV		
Lot	Top	0.1%	Elocom	00871095	ORG	ACDEFGV		
			Taro-Mometasone	02266385	TAR	ACDEFGV		

D07AC13	MOMETASONE								
Ont	Top	0.1%				Elocom	00851736	ORG	ACDEFGV
						Teva-Mometasone	02248130	TEV	ACDEFGV
D07AC21	ULOBETASOL								
Lot	Top	0.01%				Bryhali	02506262	BSL	ACDEFGV
<b>D07AD</b>	<b>CORTICOSTEROIDS, VERY POTENT (GROUP IV)</b>								
D07AD01	CLOBETASOL								
Crm	Top	0.05%				Dermovate	02213265	TPH	ACDEFGV
						Mylan-Clobetasol	02024187	MYL	ACDEFGV
						ratio-Clobetasol	01910272	TEV	ACDEFGV
						Taro-Clobetasol Cream	02245523	TAR	ACDEFGV
Lot	Top	0.05%				Dermovate	02213281	TPH	ACDEFGV
						Mylan-Clobetasol Propionate	02216213	MYL	ACDEFGV
						ratio-Clobetasol	01910299	TEV	ACDEFGV
						Taro-Clobetasol Topical Sol'n	02245522	TAR	ACDEFGV
Ont	Top	0.05%				Dermovate	02213273	TPH	ACDEFGV
						Mylan-Clobetasol	02026767	MYL	ACDEFGV
						ratio-Clobetasol	01910280	TEV	ACDEFGV
						Taro-Clobetasol Ointment	02245524	TAR	ACDEFGV
<b>D07C</b>	<b>CORTICOSTEROIDS, COMBINATIONS WITH ANTIBIOTICS</b>								
<b>D07CA</b>	<b>CORTICOSTEROIDS, WEAK, COMBINATIONS WITH ANTIBIOTICS</b>								
D07CA01	HYDROCORTISONE AND ANTIBIOTICS								
	HYDROCORTISONE / CLIOQUINOL								
Crm	Top	1% / 3%				Vioform HC	00074500	PAL	ACDEFGV
	HYDROCORTISONE / FUSIDIC ACID								
Crm	Top	1% / 2%				Fucidin H	02238578	LEO	ACDEFGV
<b>D07CB</b>	<b>CORTICOSTEROIDS, MODERATELY POTENT, COMBINATIONS WITH ANTIBIOTICS</b>								
D07CB01	TRIAMCINOLONE AND ANTIBIOTICS								
	TRIAMCINOLONE / NEOMYCIN / NYSTATIN / GRAMICIDIN								
Crm	Top	1 mg / 2.5 mg / 100 000 IU / 0.25 mg				Viaderm K-C	00717002	TAR	ACDEFGV
Ont	Top	1 mg / 2.5 mg / 100 000 IU / 0.25 mg				Viaderm K-C	00717029	TAR	ACDEFGV



D07CB05	FLUMETASONE AND ANTIBIOTICS FLUMETASONE / CLIOQUINOL	Crm Top 0.02% / 3%	Locacorten-Vioform	00074462	PAL	ACDEFGV
<b>D07CC</b>	<b>CORTICOSTEROIDS, POTENT, COMBINATIONS WITH ANTIBIOTICS</b>					
D07CC01	BETAMETHASONE AND ANTIBIOTICS BETAMETHASONE / GENTAMICIN	Crm Top 0.1% / 0.1%	Valisone G	00177016	BSL	ACDEFGV
<b>D07X</b>	<b>CORTICOSTEROIDS, OTHER COMBINATIONS</b>					
<b>D07XA</b>	<b>CORTICOSTEROIDS, WEAK, OTHER COMBINATIONS</b>					
D07XA01	HYDROCORTISONE HYDROCORTISONE / PRAMOXINE	Crm Top 1% / 1%	Pramox HC	00770957	DPT	ACDEFGV
<b>D07XC</b>	<b>CORTICOSTEROIDS, POTENT, OTHER COMBINATIONS</b>					
D07XC01	BETAMETHASONE BETAMETHASONE / SALICYLIC ACID	Lot Top 0.5 mg / 20 mg	ratio-Topisalic	02245688	TEV	ACDEFGV
		Ont Top 0.5 mg / 30 mg	Diprosalic	00578436	ORG	ACDEFGV
<b>D09</b>	<b>MEDICATED DRESSINGS</b>					
<b>D09A</b>	<b>MEDICATED DRESSINGS</b>					
<b>D09AA</b>	<b>MEDICATED DRESSINGS WITH ANTIINFECTIVES</b>					
D09AA01	FRAMYCETIN	Dre Top 1%	Sofra-Tulle (10cm x 10cm)	01988840	ERF	ACDEFGVW
			Sofra-Tulle (10cm x 30cm)	01987682	ERF	ACDEFGVW
<b>D10</b>	<b>ANTI-ACNE PREPARATIONS</b>					
<b>D10A</b>	<b>ANTI-ACNE PREPARATIONS FOR TOPICAL USE</b>					
<b>D10AD</b>	<b>RETINOIDS FOR TOPICAL USE IN ACNE</b>					
D10AD01	TRETINOIN	Crm Top 0.01%	Stieva-A	00657204	GSK	CDEFG
		Crm Top 0.025%	Stieva-A	00578576	GSK	CDEFG
		Crm Top 0.05%	Retin-A	00443794	BSL	CDEFG
			Stieva-A	00518182	GSK	CDEFG
		Gel Top 0.01%	Vitamin A Acid	01926462	BSL	CDEFG

D10AD01	TRETINOIN							
Gel	Top	0.025%		Vitamin A Acid	01926470	BSL	CDEFG	
Gel	Top	0.05%		Vitamin A Acid	01926489	BSL	CDEFG	

**D10AF ANTIINFECTIVES FOR TREATMENT OF ACNE**

D10AF01	CLINDAMYCIN							
Liq	Top	1%		Clindamycin Phosphate Topical Solution	02483769	HIK	ACDEFGV	
				Taro-Clindamycin	02266938	TAR	ACDEFGV	

**D10AX OTHER ANTI ACNE PREPARATIONS FOR TOPICAL USE**

D10AX03	AZELAIC ACID							
Gel	Top	15%		Finacea	02270811	LEO	ACDEFGV	

**D10B ANTI ACNE PREPARATIONS FOR SYSTEMIC USE**

**D10BA RETINOIDS FOR TREATMENT OF ACNE**

D10BA01	ISOTRETINOIN							
Cap	Orl	10 mg		Accutane Roche	00582344	HLR	ACDEFGV	
				Epuris	02396971	CIP	ACDEFGV	
				Clarus	02257955	MYL	ACDEFGV	
Cap	Orl	20 mg		Epuris	02396998	CIP	ACDEFGV	
Cap	Orl	30 mg		Epuris	02397005	CIP	ACDEFGV	
Cap	Orl	40 mg		Accutane Roche	00582352	HLR	ACDEFGV	
				Epuris	02397013	CIP	ACDEFGV	
				Clarus	02257963	MYL	ACDEFGV	

**D11 OTHER DERMATOLOGICAL PREPARATIONS**

**D11A OTHER DERMATOLOGICAL PREPARATIONS**

**D11AH AGENTS FOR DERMATITIS, EXCLUDING CORTICOSTEROIDS**

D11AH01	TACROLIMUS							
Ont	Top	0.03%		Protopic	02244149	LEO	(SA)	
Ont	Top	0.1%		Protopic	02244148	LEO	(SA)	
D11AH05	DUPILUMAB							
Liq	SC	200 mg / 1.14 mL		Dupixent	02492504	SAV	(SA)	
				Dupixent (prefilled pen)	02524252	SAV	(SA)	

D11AH05 DUPILUMAB

Liq SC 300 mg / 2 mL

Dupixent (autoinjector) 02510049 SAV (SA)

Dupixent (prefilled syringe) 02470365 SAV (SA)

**G GENITO URINARY SYSTEM AND SEX HORMONES**

**G01 GYNECOLOGICAL ANTIINFECTIVES AND ANTISEPTICS**

**G01A ANTIINFECTIVES AND ANTISEPTICS, EXCLUDING COMBINATIONS WITH CORTICOSTEROIDS**

**G01AA ANTIBIOTICS**

G01AA01 NYSTATIN

Crm Vag 25 000 IU

Nyaderm 00716901 TAR ACDEFGV

G01AA10 CLINDAMYCIN

Crm Vag 20 mg/g

Dalacin Vaginal Cream 02060604 PAL ACDEFGV

**G01AF IMIDAZOLE DERIVATIVES**

G01AF01 METRONIDAZOLE

Crm Vag 10%

Flagyl 01926861 SAV ACDEFGV

Gel Vag 0.75%

Nidagel 02125226 BSL ACDEFGV

G01AF02 CLOTRIMAZOLE

Crm Vag 1%

Canesten 02150891 BAY ACDEFGV

Crm Vag 2%

Canesten 3 02150905 BAY ACDEFGV

Crm Vag 500 mg/1%

Canesten 1 Comfortab 02264102 BAY ACDEFGV

Canesten 3 Comfortab Combi-Pak 02264099 BAY ACDEFGV

G01AF04 MICONAZOLE

Crm Vag 2%

Monistat 7 02084309 INP ACDEFGV

Micozole Vaginal 2% 02231106 TAR ACDEFGV

Crm Vag 1 200 mg / 2%

Monistat 3 Dual Pak 02126249 INP ACDEFGV

Sup Vag 400 mg

Monistat-3 02126605 INP ACDEFGV

**G01AG TRIAZOLE DERIVATIVES**

G01AG02 TERCONAZOLE

Crm Vag 0.4%

Taro-Terconazole 02247651 TAR ACDEFGV

**G02 OTHER GYNECOLOGICALS**

**G02B CONTRACEPTIVES FOR TOPICAL USE**

**G02BA INTRAUTERINE CONTRACEPTIVES**

G02BA03 PLASTIC IUD WITH PROGESTERONE  
LEVONORGESTREL

Ins	Vag	19.5 mg	Kyleena	02459523	BAY	CDEFGV
Ins	Vag	52 mg	Mirena	02243005	BAY	ACDEFGV

**G02BB INTRAVAGINAL CONTRACEPTIVES**

G02BB01 VAGINAL RING WITH PROGESTOGEN AND ESTROGEN  
ETHINYL ESTRADIOL / ETONOGESTREL

Ins	Vag	2.6 mg / 11.4 mg	NuvaRing	02253186	ORG	CDEFG
			Haloette	02520028	SLP	CDEFG

**G02C OTHER GYNECOLOGICALS****G02CB PROLACTINE INHIBITORS**

G02CB01 BROMOCRIPTINE

Cap	Orl	5 mg	Bromocriptine	02230454	AAP	ACDEFGV
Tab	Orl	2.5 mg	Bromocriptine	02087324	AAP	ACDEFGV

G02CB03 CABERGOLINE

Tab	Orl	0.5 mg	Dostinex	02242471	PAL	ACDEFGV
			Apo-Cabergoline	02455897	APX	ACDEFGV

**G03 SEX HORMONES AND MODULATORS OF THE GENITAL SYSTEM****G03A HORMONAL CONTRACEPTIVES FOR SYSTEMIC USE****G03AA PROGESTOGENS AND ESTROGENS, FIXED COMBINATIONS**

G03AA05 NORETHISTERONE (NORETHINDRONE) AND ETHINYL ESTRADIOL

Tab	Orl	0.5 mg / 0.035 mg	Brevicon (21)	02187086	PFI	CDEFGV
			Brevicon (28)	02187094	PFI	CDEFGV
Tab	Orl	1 mg / 0.02 mg	Minestrin 1/20 (21)	00315966	ALL	CDEFGV
			Minestrin 1/20 (28)	00343838	ALL	CDEFGV
Tab	Orl	1 mg / 0.035 mg	Brevicon 1/35 (21)	02189054	PFI	CDEFGV
			Brevicon 1/35 (28)	02189062	PFI	CDEFGV
			Select 1/35 (21)	02197502	PFI	CDEFGV
			Select 1/35 (28)	02199297	PFI	CDEFGV
Tab	Orl	1.5 mg / 0.03 mg	Loestrin 1.5/30 (21)	00297143	WNC	CDEFGV
			Loestrin 1.5/30 (28)	00353027	WNC	CDEFGV

## G03AA07 LEVONORGESTREL AND ETHINYL ESTRADIOL

Tab	Orl	0.1 mg / 0.02 mg	Alesse (21)	02236974	PFI	CDEFGV
			Alesse (28)	02236975	PFI	CDEFGV
			Alysen (21)	02387875	APX	CDEFGV
			Alysen (28)	02387883	APX	CDEFGV
			Audrina (21)	02532174	JPC	CDEFGV
			Audrina (28)	02532182	JPC	CDEFGV
			Aviane (21)	02298538	TEV	CDEFGV
			Aviane (28)	02298546	TEV	CDEFGV

Tab	Orl	0.15 mg / 0.03 mg	Min-Ovral (21)	02042320	PFI	CDEFGV
			Min-Ovral (28)	02042339	PFI	CDEFGV
			Ovima (21)	02387085	APX	CDEFGV
			Ovima (28)	02387093	APX	CDEFGV
			Portia (21)	02295946	TEV	CDEFGV
			Portia (28)	02295954	TEV	CDEFGV

## G03AA09 DESOGESTREL AND ETHINYL ESTRADIOL

Tab	Orl	0.1 mg, 0.125 mg, 0.15 mg / 0.025 mg	Linessa (21)	02272903	APN	CDEFGV
			Linessa (28)	02257238	APN	CDEFGV

Tab	Orl	0.15 mg / 0.03 mg	Marvelon (21)	02042487	ORG	CDEFGV
			Marvelon (28)	02042479	ORG	CDEFGV
			Apri (21)	02317192	TEV	CDEFGV
			Apri (28)	02317206	TEV	CDEFGV
			Freya (21)	02396491	MYL	CDEFGV
			Freya (28)	02396610	MYL	CDEFGV
			Mirvala (21)	02410249	APX	CDEFGV
			Mirvala (28)	02410257	APX	CDEFGV

## G03AA12 DROSPIRENONE AND ETHINYLESTRADIOL

Tab	Orl	3 mg / 0.02 mg	Yaz	02321157	BAY	CDEFGV
			Drospirenone and Ethinyl Estradiol	02462060	GLM	CDEFGV
			Mya	02415380	APX	CDEFGV

Tab	Orl	3 mg / 0.03 mg	Yasmin (21)	02261723	BAY	CDEFGV
			Yasmin (28)	02261731	BAY	CDEFGV
			Drospirenone and Ethinyl Estradiol-21	02421437	GLM	CDEFGV
			Drospirenone and Ethinyl Estradiol-28	02421445	GLM	CDEFGV
			Zamine (21)	02410788	APX	CDEFGV
			Zamine (28)	02410796	APX	CDEFGV

**G03AB      PROGESTOGENS AND ESTROGENS, SEQUENTIAL PREPARATIONS****G03AB03      LEVONORGESTREL AND ETHINYL ESTRADIOL**

Tab	Orl	0.05 mg / 0.03 mg, 0.075 mg / 0.04 mg, 0.125 mg / 0.03 mg	Triquilar (21)	00707600	BAY	CDEFGV
			Triquilar (28)	00707503	BAY	CDEFGV

**G03AB04      NORETHISTERONE (NORETHINDRONE) AND ETHINYL ESTRADIOL**

Tab	Orl	0.5 mg / 0.035 mg, 1 mg / 0.035 mg	Synphasic (21)	02187108	PFI	CDEFGV
			Synphasic (28)	02187116	PFI	CDEFGV

**G03AB09      NORGESTIMATE AND ETHINYL ESTRADIOL**

Tab	Orl	0.18 mg,0.215 mg, 0.25 mg / 0.035 mg	Tri-Cira (21)	02508087	APX	CDEFGV
			Tri-Cira (28)	02508095	APX	CDEFGV
			Tri-Jordyna (21)	02486296	GLM	CDEFGV
			Tri-Jordyna (28)	02486318	GLM	CDEFGV
Tab	Orl	0.215 mg,0.18 mg, 0.025 mg / 0.025 mg	Tricira LO (21)	02401967	APX	CDEFGV
			Tricira LO (28)	02401975	APX	CDEFGV

**G03AC      PROGESTOGENS****G03AC01      NORETHISTERONE (NORETHINDRONE)**

Tab	Orl	0.35 mg	Jencycla (28)	02441306	LUP	CDEFGV
			Movisse (28)	02410303	MYL	CDEFGV

**G03AC06      MEDROXYPROGESTERONE**

Sus	Inj	150 mg/mL	Depo-Provera	02523493	PFI	CDEFGV
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**G03AC08      ETONOGESTREL**

Imp	SC	68 mg	Nexplanon	02499509	ORG	CDEFGV
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**G03AD      EMERGENCY CONTRACEPTIVES****G03AD01      LEVONORGESTREL**

Tab	Orl	1.5 mg	Plan B	02293854	PAL	CDEFGV
			Backup Plan Onestep	02433532	APX	CDEFGV
			Contingency One	02425009	MYL	CDEFGV

**G03B      ANDROGENS****G03BA      3-OXOANDROSTEN (4) DERIVATIVES****G03BA03      TESTOSTERONE  
TESTOSTERONE UNDECANOATE**

Cap	Orl	40 mg	pms-Testosterone	02322498	PMS	(SA)
			Taro-Testosterone	02421186	TAR	(SA)

G03BA03 TESTOSTERONE

Gel	Top	1%	Testim	02280248	PAL	(SA)
Gel	Top	25 mg	AndroGel Packets	02245345	BGP	(SA)
			Taro-Testosterone Gel	02463792	TAR	(SA)
Gel	Top	50 mg	AndroGel Packets	02245346	BGP	(SA)
			Taro-Testosterone Gel	02463806	TAR	(SA)
Liq	IM	100 mg/mL	Depo-Testosterone	00030783	PFI	ACDEFGV
			Taro-Testosterone	02496003	TAR	ACDEFGV
Liq	Inj	200 mg/mL	Delatestryl	00029246	BSL	ACDEFGV

**G03C ESTROGENS**

**G03CA NATURAL AND SEMISYNTHETIC ESTROGENS, PLAIN**

G03CA03 ESTRADIOL

Gel	Trd	0.06%	Estrogel	02238704	ORG	ACDEFGV
Gel	Trd	0.25 mg	Divigel	02424924	SLP	ACDEFGV
Gel	Trd	0.5 mg	Divigel	02424835	SLP	ACDEFGV
Gel	Trd	1 mg	Divigel	02424843	SLP	ACDEFGV
Ins	Vag	2 mg	Estring	02168898	PAL	ACDEFGV
Pth	Trd	25 mcg	Climara 25	02247499	BAY	ACDEFGV
			Estradot	02245676	SDZ	ACDEFGV
Pth	Trd	37.5 mcg	Estradot	02243999	SDZ	ACDEFGV
Pth	Trd	50 mcg	Climara 50	02231509	BAY	ACDEFGV
			Estradot	02244000	SDZ	ACDEFGV
			Sandoz Estradiol Derm Srd	02246967	SDZ	ACDEFGV
Pth	Trd	75 mcg	Climara 75	02247500	BAY	ACDEFGV
			Estradot	02244001	SDZ	ACDEFGV
			Sandoz Estradiol Derm Srd	02246968	SDZ	ACDEFGV
Pth	Trd	100 mcg	Estradot	02244002	SDZ	ACDEFGV
			Sandoz Estradiol Derm Srd	02246969	SDZ	ACDEFGV

G03CA03	ESTRADIOL							
Tab	Orl	0.5 mg	Estrace	02225190	PMS	ACDEFGV		
			Lupin-Estradiol	02449048	LUP	ACDEFGV		
Tab	Orl	1 mg	Estrace	02148587	PMS	ACDEFGV		
			Lupin-Estradiol	02449056	LUP	ACDEFGV		
Tab	Orl	2 mg	Estrace	02148595	PMS	ACDEFGV		
			Lupin-Estradiol	02449064	LUP	ACDEFGV		
Tab	Vag	10 mcg	Vagifem 10	02325462	NNO	ACDEFGV		

G03CA07	ESTRONE							
Crm	Vag	1 mg	Estragyn	00727369	SLP	ACDEFGV		

G03CA57	CONJUGATED ESTROGENS							
Crm	Vag	0.625 mg	Premarin	02043440	PFI	ACDEFGV		
Tab	Orl	0.3 mg	Premarin	02414678	PFI	ACDEFGV		
Tab	Orl	0.625 mg	Premarin	02414686	PFI	ACDEFGV		
Tab	Orl	1.25 mg	Premarin	02414694	PFI	ACDEFGV		

**G03D      PROGESTOGENS**

**G03DA     PREGNEN (4) DERIVATIVES**

G03DA02	MEDROXYPROGESTERONE							
Tab	Orl	2.5 mg	Provera	00708917	PFI	ACDEFGV		
			Apo-Medroxy	02244726	APX	ACDEFGV		
			Teva-Medroxyprogesterone	02221284	TEV	ACDEFGV		
Tab	Orl	5 mg	Provera	00030937	PFI	ACDEFGV		
			Apo-Medroxy	02244727	APX	ACDEFGV		
			Teva-Medroxyprogesterone	02221292	TEV	ACDEFGV		
Tab	Orl	10 mg	Provera	00729973	PFI	ACDEFGV		
			Apo-Medroxy	02277298	APX	ACDEFGV		
			Teva-Medroxyprogesterone	02221306	TEV	ACDEFGV		
Tab	Orl	100 mg	Apo-Medroxy	02267640	APX	ACDEFGV		

G03DA04	PROGESTERONE							
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**G03DA04**    **PROGESTERONE**

Cap    Orl    100 mg

Prometrium	02166704	ORG	ACDEFGV
Auro-Progesterone	02493578	ARO	ACDEFGV
pms-Progesterone	02476576	PMS	ACDEFGV
Progesterone	02516187	SAS	ACDEFGV
Reddy-Progesterone	02463113	RCH	ACDEFGV
Teva-Progesterone	02439913	TEV	ACDEFGV

**G03DB**    **PREGNADIEN DERIVATIVES****G03DB08**    **DIENOGEST**

Tab    Orl    2 mg

Visanne	02374900	BAY	(SA)
Aspen-Dienogest	02493055	APN	(SA)
Jamp Dienogest	02498189	JPC	(SA)

**G03DC**    **ESTREN DERIVATIVES****G03DC02**    **NORETHISTERONE (NORETHINDRONE)**

Tab    Orl    5 mg

Norlutate    00023760    SLP    (SA)

**G03F**    **PROGESTOGENS AND ESTROGENS IN COMBINATION****G03FA**    **PROGESTOGENS AND ESTROGENS, FIXED COMBINATIONS****G03FA01**    **NORETHISTERONE (NORETHINDRONE) AND ESTROGEN**

Pad    Trd    140 mcg / 50 mcg

Estalis    02241835    SDZ    ACDEFGV

Pad    Trd    250 mcg / 50 mcg

Estalis    02241837    SDZ    ACDEFGV

**G03H**    **ANTIANDROGENS****G03HA**    **ANTIANDROGENS, PLAIN****G03HA01**    **CYPROTERONE**

Tab    Orl    50 mg

Androcur	00704431	PMS	ACDEFV
Med-Cyproterone	02390760	GMP	ACDEFV

**G03HB**    **ANTIANDROGENS AND ESTROGENS****G03HB01**    **CYPROTERONE AND ESTROGENS**

Tab    Orl    2 mg / 0.035 mg

Diane-35	02233542	BAY	CDEFGV
Cléo-35	02436736	ATS	CDEFGV
Cyestra-35	02290308	PAL	CDEFGV
Teva-Cyproterone/Ethinyl Estradiol	02309556	TEV	CDEFGV

**G03X**    **OTHER SEX HORMONES AND MODULATORS OF THE GENITAL SYSTEM****G03XA**    **ANTIGONADOTROPHINS AND SIMILAR AGENTS****G03XA01**    **DANAZOL**

G03XA01	DANAZOL								
Cap	Orl	50 mg				Cyclomen	02018144	SAV	ACDEFV
Cap	Orl	100 mg				Cyclomen	02018152	SAV	ACDEFV
Cap	Orl	200 mg				Cyclomen	02018160	SAV	ACDEFV
<b>G03XB</b>	<b>PROGESTERONE RECEPTOR MODULATORS</b>								
G03XB51	MIFEPRISTONE, COMBINATIONS								
	MIFEPRISTONE / MISOPROSTOL								
Tab	Orl	200 mg / 200 mcg				Mifegymiso	02444038	LIN	CJ
<b>G03XC</b>	<b>OTHER SEX HORMONES</b>								
G03XC01	RALOXIFENE								
Tab	Orl	60 mg				Evista	02239028	LIL	ACDEFV
						Act Raloxifene	02358840	TEV	ACDEFV
						Apo-Raloxifene	02279215	APX	ACDEFV
<b>G04</b>	<b>UROLOGICALS</b>								
<b>G04B</b>	<b>UROLOGICALS</b>								
<b>G04BD</b>	<b>DRUGS FOR URINARY FREQUENCY AND INCONTINENCE</b>								
G04BD04	OXYBUTYNIN								
Syr	Orl	1 mg				pms-Oxybutynin	02223376	PMS	ACDEFGV
Tab	Orl	5 mg				Apo-Oxybutynin	02163543	APX	ACDEFGV
						Novo-Oxybutynin	02230394	TEV	ACDEFGV
						Oxybutynin	02350238	SAS	ACDEFGV
						pms-Oxybutynin	02240550	PMS	ACDEFGV
G04BD06	PROPIVERINE								
Tab	Orl	5 mg				Mictoryl Pediatric	02460289	DUI	(SA)
G04BD07	TOLTERODINE								
ERC	Orl	2 mg				Detrol LA	02244612	BGP	ACDEFGV
						Sandoz Tolterodine LA	02413140	SDZ	ACDEFGV
						Teva-Tolterodine LA	02412195	TEV	ACDEFGV
ERC	Orl	4 mg				Detrol LA	02244613	BGP	ACDEFGV
						Sandoz Tolterodine LA	02413159	SDZ	ACDEFGV
						Teva-Tolterodine LA	02412209	TEV	ACDEFGV

## G04BD07 TOLTERODINE

Tab Orl 1 mg

Detrol	02239064	UJC	ACDEFGV
Jamp Tolterodine	02496836	JPC	ACDEFGV
Mint-Tolterodine	02423308	MNT	ACDEFGV
Teva-Tolterodine	02299593	TEV	ACDEFGV

Tab Orl 2 mg

Detrol	02239065	UJC	ACDEFGV
Jamp Tolterodine	02496844	JPC	ACDEFGV
Mint-Tolterodine	02423316	MNT	ACDEFGV
Teva-Tolterodine	02299607	TEV	ACDEFGV

## G04BD08 SOLIFENACIN

Tab Orl 5 mg

Vesicare	02277263	ASL	ACDEFGV
ACH-Solifenacin Succinate	02439344	AHI	ACDEFGV
Auro-Solifenacin	02446375	ARO	ACDEFGV
Jamp-Solifenacin	02424339	JPC	ACDEFGV
pms-Solifenacin	02417723	PMS	ACDEFGV
PRZ-Solifenacin	02493039	PRZ	ACDEFGV
Sandoz Solifenacin	02399032	SDZ	ACDEFGV
Solifenacin	02458241	SAS	ACDEFGV
Taro-Solifenacin	02437988	SUN	ACDEFGV
Teva-Solifenacin	02397900	TEV	ACDEFGV

Tab Orl 10 mg

Vesicare	02277271	ASL	ACDEFGV
ACH-Solifenacin Succinate	02439352	AHI	ACDEFGV
Auro-Solifenacin	02446383	ARO	ACDEFGV
Jamp-Solifenacin	02424347	JPC	ACDEFGV
pms-Solifenacin	02417731	PMS	ACDEFGV
PRZ-Solifenacin	02493047	PRZ	ACDEFGV
Sandoz Solifenacin	02399040	SDZ	ACDEFGV
Solifenacin	02458268	SAS	ACDEFGV
Taro-Solifenacin	02437996	SUN	ACDEFGV
Teva-Solifenacin	02397919	TEV	ACDEFGV

## G04BD09 TROSPIUM

Tab Orl 20 mg

Trosec	02275066	SNV	(SA)
Mar-Trospium	02488353	MAR	(SA)

## G04BD10 DARIFENACIN

G04BD10	DARIFENACIN						
ERT	Orl	7.5 mg	Enablex	02273217	SLP	(SA)	
			Apo-Darifenacin	02452510	APX	(SA)	
			Jamp Darifenacin	02491869	JPC	(SA)	
ERT	Orl	15 mg	Enablex	02273225	SLP	(SA)	
			Apo-Darifenacin	02452529	APX	(SA)	
			Jamp Darifenacin	02491877	JPC	(SA)	
G04BD11	FESOTERODINE						
ERT	Orl	4 mg	Toviaz	02380021	PFI	(SA)	
			Sandoz Fesoterodine Fumarate	02521768	SDZ	(SA)	
ERT	Orl	8 mg	Toviaz	02380048	PFI	(SA)	
			Sandoz Fesoterodine Fumarate	02521776	SDZ	(SA)	
G04BD12	MIRABEGRON						
ERT	Orl	25 mg	Myrbetriq	02402874	ASL	(SA)	
ERT	Orl	50 mg	Myrbetriq	02402882	ASL	(SA)	
<b>G04BX</b>	<b>OTHER UROLOGICAL</b>						
G04BX13	DIMETHYL SULFOXIDE						
	Liq	ITV 500 mg/g	Rimso-50	00493392	MYL	ACDEFGV	
<b>G04C</b>	<b>DRUGS USED IN BENIGN PROSTATIC HYPERTROPHY</b>						
<b>G04CA</b>	<b>ALPHA-ADRENORECEPTOR ANTAGONISTS</b>						
G04CA01	ALFUZOSIN						
ERT	Orl	10 mg	Xatral	02245565	SAV	ACDEFGV	
			Alfuzosin	02519844	SAS	ACDEFGV	
			Alfuzosin	02447576	SIV	ACDEFGV	
			Apo-Alfuzosin	02315866	APX	ACDEFGV	
			Auro-Alfuzosin	02443201	ARO	ACDEFGV	
			Sandoz Alfuzosin	02304678	SDZ	ACDEFGV	
G04CA02	TAMSULOSIN						

G04CA02 TAMSULOSIN

ERT Orl 0.4 mg

Flomax CR 02270102 BOE ACDEFV  
 Apo-Tamsulosin CR 02362406 APX ACDEFV  
 Sandoz Tamsulosin CR 02340208 SDZ ACDEFV  
 Tamsulosin CR 02427117 SAS ACDEFV  
 Tamsulosin CR 02429667 SIV ACDEFV  
 Teva-Tamsulosin CR 02368242 TEV ACDEFV

SRC Orl 0.4 mg

Sandoz Tamsulosin 02319217 SDZ ACDEFV

G04CA03 TERAZOSIN

Tab Orl 1 mg

Apo-Terazosin 02234502 APX ACDEFV  
 pms-Terazosin 02243518 PMS ACDEFV

Tab Orl 2 mg

Apo-Terazosin 02234503 APX ACDEFV  
 pms-Terazosin 02243519 PMS ACDEFV

Tab Orl 5 mg

Apo-Terazosin 02234504 APX ACDEFV  
 pms-Terazosin 02243520 PMS ACDEFV  
 Teva-Terazosin 02230807 TEV ACDEFV

Tab Orl 10 mg

Apo-Terazosin 02234505 APX ACDEFV  
 pms-Terazosin 02243521 PMS ACDEFV

**G04CB TESTOSTERONE-5-ALPHA REDUCTASE INHIBITORS**

G04CB01 FINASTERIDE

Tab Orl 5 mg

Proscar 02010909 ORG ACDEFGV  
 Apo-Finasteride 02365383 APX ACDEFGV  
 Auro-Finasteride 02405814 ARO ACDEFGV  
 Finasteride 02355043 AHI ACDEFGV  
 Finasteride 02445077 SAS ACDEFGV  
 Finasteride 02447541 SIV ACDEFGV  
 Jamp-Finasteride 02357224 JPC ACDEFGV  
 M-Finasteride 02522489 MRA ACDEFGV  
 Mint-Finasteride 02389878 MNT ACDEFGV  
 pms-Finasteride 02310112 PMS ACDEFGV  
 Riva-Finasteride 02455013 RIV ACDEFGV  
 Sandoz Finasteride 02322579 SDZ ACDEFGV  
 Teva-Finasteride 02348500 TEV ACDEFGV

G04CB02 DUTASTERIDE

## G04CB02 DUTASTERIDE

Cap Orl 0.5 mg

Avodart	02247813	GSK	ACDEFGV
Apo-Dutasteride	02404206	APX	ACDEFGV
Auro-Dutasteride	02469308	ARO	ACDEFGV
Dutasteride	02443058	SAS	ACDEFGV
Dutasteride	02429012	SIV	ACDEFGV
Jamp-Dutasteride	02484870	JPC	ACDEFGV
Med-Dutasteride	02416298	GMP	ACDEFGV
Mint-Dutasteride	02428873	MNT	ACDEFGV
pms-Dutasteride	02393220	PMS	ACDEFGV
Priva-Dutasteride	02490587	NRA	ACDEFGV
Sandoz Dutasteride	02424444	SDZ	ACDEFGV
Teva-Dutasteride	02408287	TEV	ACDEFGV

**H SYSTEMIC HORMONAL PREPARATIONS EXCLUDING SEX HORMONES****H01 PITUITARY AND HYPOTHALAMIC HORMONES AND ANALOGUES****H01A ANTERIOR PITUITARY LOBE HORMONES AND ANALOGUES****H01AC SOMATROPIN AND SOMATROPIN AGONISTS**

## H01AC01 SOMATROPIN

Ctg Inj 6 mg

Humatrope 02243077 LIL T (SA)

Ctg Inj 12 mg

Humatrope 02243078 LIL T (SA)

Ctg Inj 24 mg

Humatrope 02243079 LIL T (SA)

Liq Inj 5 mg / 1.5 mL

Norditropin Nordiflex 02334852 NNO T (SA)

Omnitrope 02325063 SDZ T (SA)

Liq Inj 5 mg / 2 mL

Nutropin AQ NuSpin 02399091 HLR T (SA)

Liq Inj 6 mg

Saizen 02350122 EMD T (SA)

Liq Inj 10 mg / 1.5 mL

Norditropin Nordiflex 02334860 NNO T (SA)

Omnitrope 02325071 SDZ T (SA)

Liq Inj 10 mg / 2 mL

Nutropin AQ NuSpin 02376393 HLR T (SA)

Liq Inj 12 mg

Saizen 02350130 EMD T (SA)

Liq Inj 15 mg / 1.5 mL

Norditropin Nordiflex 02334879 NNO T (SA)

Omnitrope 02459647 SDZ T (SA)

H01AC01 SOMATROPIN

Liq	Inj	20 mg	Saizen	02350149	EMD	T (SA)
Liq	Inj	20 mg / 2 mL	Nutropin AQ NuSpin	02399083	HLR	T (SA)
Pwd	Inj	5 mg	Saizen	02237971	EMD	T (SA)
Pws	SC	0.6 mg	Genotropin MiniQuick	02401762	PFI	T (SA)
Pws	SC	0.8 mg	Genotropin MiniQuick	02401770	PFI	T (SA)
Pws	SC	1 mg	Genotropin MiniQuick	02401789	PFI	T (SA)
Pws	SC	1.2 mg	Genotropin MiniQuick	02401797	PFI	T (SA)
Pws	SC	1.4 mg	Genotropin MiniQuick	02401800	PFI	T (SA)
Pws	SC	1.6 mg	Genotropin MiniQuick	02401819	PFI	T (SA)
Pws	SC	1.8 mg	Genotropin MiniQuick	02401827	PFI	T (SA)
Pws	SC	2 mg	Genotropin MiniQuick	02401835	PFI	T (SA)
Pws	SC	5.3 mg	Genotropin GoQuick	02401703	PFI	T (SA)
Pws	SC	12 mg	Genotropin GoQuick	02401711	PFI	T (SA)

H01AC03 MECASERMIN

Liq	SC	10 mg/mL	Increlex	02509733	IPS	(SA)
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**H01B POSTERIOR PITUITARY LOBE HORMONES**

**H01BA VASOPRESSIN AND ANALOGUES**

H01BA02 DESMOPRESSIN

Aem	Nas	10 mcg	Desmopressin Spray	02242465	AAP	(SA)
Liq	Inj	4 mcg/mL	DDAVP	00873993	FEI	ACDEFGV
			Bipazen	02513579	KVR	ACDEFGV
ODT	Slg	60 mcg	DDAVP Melt	02284995	FEI	CDEF-18G (SA)
ODT	Slg	120 mcg	DDAVP Melt	02285002	FEI	CDEF-18G (SA)

**H01BA02 DESMOPRESSIN**

Tab	Orl	0.1 mg	Apo-Desmopressin	02284030	APX	CDEF-18G (SA)
			pms-Desmopressin	02304368	PMS	CDEF-18G (SA)
Tab	Orl	0.2 mg	Apo-Desmopressin	02284049	APX	CDEF-18G (SA)
			pms-Desmopressin	02304376	PMS	CDEF-18G (SA)

**H01C HYPOTHALAMIC HORMONES****H01CA GONADOTROPIN-RELEASING HORMONES****H01CA02 NAFARELIN**

Liq	Nas	2 mg/mL	Synarel	02188783	PFI	ACDEFGV
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**H01CB SOMATOSTATIN AND ANALOGUES****H01CB02 OCTREOTIDE**

Liq	Inj	0.05 mg/mL	Sandostatin	00839191	NVR	ACDEFGVW
			Octreotide Acetate Omega	02248639	OMG	ACDEFGVW
Liq	Inj	0.1 mg/mL	Sandostatin	00839205	NVR	ACDEFGVW
			Octreotide Acetate Omega	02248640	OMG	ACDEFGVW
Liq	Inj	0.2 mg/mL	Octreotide Acetate Omega	02248642	OMG	ACDEFGVW
Liq	Inj	0.5 mg/mL	Octreotide Acetate Omega	02248641	OMG	ACDEFGVW
Pws	Inj	10 mg	Sandostatin LAR	02239323	NVR	ACDEFGVW
			Octreotide for Injectable Suspension	02503751	TEV	ACDEFGVW
Pws	Inj	20 mg	Sandostatin LAR	02239324	NVR	ACDEFGVW
			Octreotide for Injectable Suspension	02503778	TEV	ACDEFGVW
Pws	Inj	30 mg	Sandostatin LAR	02239325	NVR	ACDEFGVW
			Octreotide for Injectable Suspension	02503786	TEV	ACDEFGVW

**H01CB03 LANREOTIDE**

Liq	SC	60 mg / 0.5 mL	Somatuline Autogel (prefilled syringe)	02283395	IPS	ACDEFGV
Liq	SC	90 mg / 0.5 mL	Somatuline Autogel (prefilled syringe)	02283409	IPS	ACDEFGV
Liq	SC	120 mg / 0.5 mL	Somatuline Autogel (prefilled syringe)	02283417	IPS	ACDEFGV

**H02 CORTICOSTEROIDS FOR SYSTEMIC USE****H02A CORTICOSTEROIDS FOR SYSTEMIC USE, PLAIN**



**H02AA MINERALOCORTICIDS**

H02AA02 FLUDROCORTISONE

Tab Orl 0.1 mg Florinef 02086026 PAL ACDEFGV

**H02AB GLUCOCORTICIDS**

H02AB01 BETAMETHASONE

Sus IM 3 mg / 3 mg Celestone Soluspan 00028096 ORG ACDEFGV

H02AB02 DEXAMETHASONE

Liq Inj 4 mg/mL Dexamethasone sodium phosphate 00664227 SDZ ACDEFGVW

Dexamethasone sodium phosphate 01977547 STR ACDEFGVW

Dexamethasone-Omega 02204266 OMG ACDEFGVW

Tab Orl 0.5 mg Apo-Dexamethasone 02261081 APX ACDEFGVW

pms-Dexamethasone 01964976 PMS ACDEFGVW

Tab Orl 2 mg pms-Dexamethasone 02279363 PMS ACDEFGVW

Tab Orl 4 mg Apo-Dexamethasone 02250055 APX ACDEFGVW

pms-Dexamethasone 01964070 PMS ACDEFGVW

H02AB04 METHYLPREDNISOLONE

Pws Inj 40 mg Solu-Medrol (Act-O-Vial) 02367947 PFI ACDEFGVW

Pws Inj 125 mg Solu-Medrol (Act-O-Vial) 02367955 PFI ACDEFGVW

Pws Inj 500 mg Solu-Medrol 00030678 PFI ACDEFGVW

Solu-Medrol (Act-O-Vial) 02367963 PFI ACDEFGVW

Pws Inj 1 g Solu-Medrol 00036137 PFI ACDEFGVW

Solu-Medrol (Act-O-Vial) 02367971 PFI ACDEFGVW

Sus Inj 20 mg/mL Depo-Medrol 01934325 PFI ACDEFGVW

Sus Inj 40 mg/mL Depo-Medrol 00030759 PFI ACDEFGVW

Depo-Medrol 01934333 PFI ACDEFGVW

Sus Inj 80 mg/mL Depo-Medrol 00030767 PFI ACDEFGVW

Depo-Medrol 01934341 PFI ACDEFGVW

Tab Orl 4 mg Medrol 00030988 PFI ACDEFGVW

Tab Orl 16 mg Medrol 00036129 PFI ACDEFGVW

H02AB06	PREDNISOLONE							
	Liq	Orl	5 mg / 5 mL		Pediapred	02230619	SAV	ACDEFGVW
					pms-Prednisolone	02245532	PMS	ACDEFGVW
H02AB07	PREDNISONONE							
	Tab	Orl	1 mg		Winpred	00271373	AAP	ACDEFGRVW
	Tab	Orl	5 mg		Apo-Prednisone	00312770	APX	ABCDEFGRVW
					Teva-Prednisone	00021695	TEV	ABCDEFGRVW
	Tab	Orl	50 mg		Apo-Prednisone	00550957	APX	ACDEFGRVW
					Teva-Prednisone	00232378	TEV	ACDEFGRVW
H02AB08	TRIAMCINOLONE							
	Sus	IA	10 mg/mL		Kenalog-10	01999761	BRI	ACDEFGV
	Sus	IA	20 mg/mL		Trispan	02470632	MDX	(SA)
	Sus	IA	40 mg/mL		Kenalog-40	01999869	BRI	ACDEFGV
					Triamcinolone Acetonide	01977563	STR	ACDEFGV
H02AB09	HYDROCORTISONE							
	Pws	Inj	100 mg		Solu-Cortef (Act-O-Vial)	00030600	PFI	ACDEFGVW
	Pws	Inj	250 mg		Solu-Cortef (Act-O-Vial)	00030619	PFI	ACDEFGVW
	Pws	Inj	500 mg		Solu-Cortef (Act-O-Vial)	00030627	PFI	ACDEFGVW
	Pws	Inj	1 g		Solu-Cortef (Act-O-Vial)	00030635	PFI	ACDEFGVW
	Tab	Orl	10 mg		Cortef	00030910	PFI	ACDEFGVW
					Auro-Hydrocortisone	02524465	ARO	ACDEFGVW
	Tab	Orl	20 mg		Cortef	00030929	PFI	ACDEFGVW
					Auro-Hydrocortisone	02524473	ARO	ACDEFGVW
H02AB10	CORTISONE							
	Tab	Orl	25 mg		Cortisone	00280437	BSL	ACDEFGVW
<b>H02B</b>	<b>CORTICOSTEROIDS FOR SYSTEMIC USE, COMBINATIONS</b>							
<b>H02BX</b>	<b>CORTICOSTEROIDS FOR SYSTEMIC USE, COMBINATIONS</b>							
H02BX01	METHYLPREDNISOLONE, COMBINATIONS							
	METHYLPREDNISOLONE / LIDOCAINE							

H02BX01 METHYLPREDNISOLONE, COMBINATIONS  
METHYLPREDNISOLONE / LIDOCAINE

Sus IA 40 mg / 10 mg

Depo-Medrol with Lidocaine 00260428 PFI ACDEFGVW

**H03 THYROID THERAPY**

**H03A THYROID PREPARATIONS**

**H03AA THYROID HORMONES**

H03AA01 LEVOTHYROXINE SODIUM

Tab Orl 0.025 mg

Synthroid 02172062 BGP ACDEFGV

Tab Orl 0.05 mg

Synthroid 02172070 BGP ACDEFGV

Eltroxin 02213192 APN ACDEFGV

Tab Orl 0.075 mg

Synthroid 02172089 BGP ACDEFGV

Tab Orl 0.088 mg

Synthroid 02172097 BGP ACDEFGV

Tab Orl 0.1 mg

Synthroid 02172100 BGP ACDEFGV

Eltroxin 02213206 APN ACDEFGV

Tab Orl 0.112 mg

Synthroid 02171228 BGP ACDEFGV

Tab Orl 0.125 mg

Synthroid 02172119 BGP ACDEFGV

Tab Orl 0.137 mg

Synthroid 02233852 BGP ACDEFGV

Tab Orl 0.15 mg

Synthroid 02172127 BGP ACDEFGV

Eltroxin 02213214 APN ACDEFGV

Tab Orl 0.175 mg

Synthroid 02172135 BGP ACDEFGV

Tab Orl 0.2 mg

Synthroid 02172143 BGP ACDEFGV

Eltroxin 02213222 APN ACDEFGV

Tab Orl 0.3 mg

Synthroid 02172151 BGP ACDEFGV

H03AA02 LIOTHYRONINE SODIUM

Tab Orl 5 mcg

Cytomel 01919458 PFI ACDEFGV

Teva-Liothyronine 02494337 TEV ACDEFGV

Tab Orl 25 mcg

Cytomel 01919466 PFI ACDEFGV

Teva-Liothyronine 02494345 TEV ACDEFGV

H03AA05 THYROID GLAND PREPARATIONS  
DESICCATED THYROID

Tab	Orl	30 mg	Thyroid	00023949	ERF	ACDEFGV
Tab	Orl	60 mg	Thyroid	00023957	ERF	ACDEFGV
Tab	Orl	125 mg	Thyroid	00023965	ERF	ACDEFGV

**H03B ANTITHYROID PREPARATIONS**

**H03BA THIOURACILS**

H03BA02 PROPYLTHIOURACIL

Tab	Orl	50 mg	Halycil	02521059	ARN	ACDEFGV
			Propylthiouracil	02523019	PCI	ACDEFGV

**H03BB SULPHUR-CONTAINING IMIDAZOLE DERIVATIVES**

H03BB02 THIAMAZOLE (METHIMAZOLE)

Tab	Orl	5 mg	Tapazole	00015741	PAL	ACDEFGV
			Jamp Methimazole	02490625	JPC	ACDEFGV
			Mar-Methimazole	02480107	MAR	ACDEFGV
Tab	Orl	10 mg	Tapazole	02296039	PAL	ACDEFGV
			Jamp Methimazole	02490633	JPC	ACDEFGV
			Mar-Methimazole	02480115	MAR	ACDEFGV

**H04 PANCREATIC HORMONES**

**H04A GLYCOGENOLYTIC HORMONES**

**H04AA GLYCOGENOLYTIC HORMONES**

H04AA01 GLUCAGON

Pws	Inj	1 mg	Glucagen	02333619	PAL	ACDEFGV
			Glucagen Hypokit	02333627	PAL	ACDEFGV
			Glucagon	02243297	LIL	ACDEFGV
Pws	Nas	3 mg	Baqsimi	02492415	LIL	(SA)

**H05 CALCIUM HOMEOSTASIS**

**H05B ANTI-PARATHYROID AGENTS**

**H05BA CALCITONIN PREPARATIONS**

H05BA01 CALCITONIN (SALMON SYNTHETIC)

Liq	Inj	200 U/mL	Calcimar	01926691	SAV	ACDEFGV
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**H05BX OTHER ANTI-PARATHYROID AGENTS**

H05BX01 CINACALCET

H05BX01 CINACALCET

Tab Orl 30 mg

Sensipar 02257130 AGA ACDEFGV  
Apo-Cinacalcet 02452693 APX ACDEFGV  
Auro-Cinacalcet 02478900 ARO ACDEFGV  
Cinacalcet 02524880 SAS ACDEFGV  
Jamp Cinacalcet 02500094 JPC ACDEFGV  
M-Cinacalcet 02481987 MRA ACDEFGV  
Mar-Cinacalcet 02480298 MAR ACDEFGV  
pms-Cinacalcet 02517604 PMS ACDEFGV  
Teva-Cinacalcet 02441624 TEV ACDEFGV

Tab Orl 60 mg

Sensipar 02257149 AGA ACDEFGV  
Apo-Cinacalcet 02452707 APX ACDEFGV  
Auro-Cinacalcet 02478919 ARO ACDEFGV  
Jamp Cinacalcet 02500108 JPC ACDEFGV  
M-Cinacalcet 02481995 MRA ACDEFGV  
Mar-Cinacalcet 02480301 MAR ACDEFGV  
pms-Cinacalcet 02517612 PMS ACDEFGV  
Teva-Cinacalcet 02441632 TEV ACDEFGV

Tab Orl 90 mg

Sensipar 02257157 AGA ACDEFGV  
Apo-Cinacalcet 02452715 APX ACDEFGV  
Auro-Cinacalcet 02478943 ARO ACDEFGV  
Jamp Cinacalcet 02500116 JPC ACDEFGV  
M-Cinacalcet 02482002 MRA ACDEFGV  
Mar-Cinacalcet 02480328 MAR ACDEFGV  
pms-Cinacalcet 02517620 PMS ACDEFGV  
Teva-Cinacalcet 02441640 TEV ACDEFGV

**J ANTIINFECTIVES FOR SYSTEMIC USE**

**J01 ANTIBACTERIALS FOR SYSTEMIC USE**

**J01A TETRACYCLINES**

**J01AA TETRACYCLINES**

J01AA02 DOXYCYCLINE

Cap Orl 100 mg

Apo-Doxy 00740713 APX ABCDEFGVW  
Doxycycline 02351234 SAS ABCDEFGVW  
Teva-Doxycycline 00725250 TEV ABCDEFGVW

J01AA02	DOXYCYCLINE							
Tab	Orl	100 mg		Doxycin	00860751	RIV	ABCDEFGVW	
				Apo-Doxy	00874256	APX	ABCDEFGVW	
				Doxycycline	02351242	SAS	ABCDEFGVW	
				Teva-Doxycycline	02158574	TEV	ABCDEFGVW	
J01AA07	TETRACYCLINE							
Cap	Orl	250 mg		Tetra	00580929	AAP	ACDEFGVW	
J01AA08	MINOCYCLINE							
Cap	Orl	50 mg		Minocycline	02084090	AAP	ACDEFGV	
Cap	Orl	100 mg		Minocycline	02084104	AAP	ACDEFGV	
J01AA12	TIGECYCLINE							
Pws	IV	50 mg		Tygacil	02285401	PFI	W (SA)	
<b>J01C</b>	<b>BETA LACTAM ANTIBACTERIALS, PENICILLINS</b>							
<b>J01CA</b>	<b>PENICILLIN WITH EXTENDED SPECTRUMS</b>							
J01CA01	AMPICILLIN							
Cap	Orl	250 mg		Teva-Ampicillin	00020877	TEV	ACDEFGVW	
Cap	Orl	500 mg		Teva-Ampicillin	00020885	TEV	ACDEFGVW	
Pws	Inj	500 mg		Ampicillin Sodium	00872652	TEV	ACDEFGVW	
Pws	Inj	1 g		Ampicillin Sodium	01933345	TEV	ACDEFGVW	
Pws	Inj	2 g		Ampicillin Sodium	01933353	TEV	ACDEFGVW	
J01CA04	AMOXICILLIN							
Cap	Orl	250 mg		Amoxicillin (Disc/non disp Nov 4/23)	02352710	SAS	ABCDEFGVW	
				Amoxicillin Capsules BP	02525348	SAS	ABCDEFGVW	
				Apo-Amoxi	00628115	APX	ABCDEFGVW	
				Auro-Amoxicillin	02388073	ARO	ABCDEFGVW	
				Jamp-Amoxicillin	02433060	JPC	ABCDEFGVW	
				Novamoxin	00406724	TEV	ABCDEFGVW	

J01CA04 AMOXICILLIN

Cap	Orl	500 mg	Amoxicillin	02401509	SIV	ABCDEFGVW
			Amoxicillin (Disc/non disp Nov 4/23)	02352729	SAS	ABCDEFGVW
			Amoxicillin Capsules BP	02525356	SAS	ABCDEFGVW
			Apo-Amoxi	00628123	APX	ABCDEFGVW
			Auro-Amoxicillin	02388081	ARO	ABCDEFGVW
			Jamp-Amoxicillin	02433079	JPC	ABCDEFGVW
			Novamoxin	00406716	TEV	ABCDEFGVW
Pws	Orl	25 mg	Apo-Amoxi	00628131	APX	ABCDEFGVW
			Jamp-Amoxicillin	02535793	JPC	ABCDEFGVW
Pws	Orl	50 mg	Amoxicillin	02352753	SAS	ABCDEFGVW
			Amoxicillin	02401541	SIV	ABCDEFGVW
			Amoxicillin (sugar-reduced)	02352788	SAS	ABCDEFGVW
			Apo-Amoxi	00628158	APX	ABCDEFGVW
			Jamp-Amoxicillin	02535815	JPC	ABCDEFGVW
			Moxilen (Temporary Benefit)	09858237	JNO	ABCDEFGVW
			Novamoxin	00452130	TEV	ABCDEFGVW
			Novamoxin 125 (sugar-reduced)	01934163	TEV	ABCDEFGVW
TabC	Orl	250 mg	Novamoxin chew	02036355	TEV	ABCDEFGVW

**J01CE BETA-LACTAMASE SENSITIVE PENICILLINS**

J01CE02 PHENOXYMETHYLPENICILLIN (PENICILLIN V)

Tab	Orl	300 mg	Pen VK	00642215	AAP	ACDEFGVW
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J01CE08 BENZATHINE BENZYL PENICILLIN (PENICILLIN G BENZATHINE)

Sus	Inj	1 200 000 unit / 2 mL	Bicillin L-A	02291924	PFI	ACDEFGV
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**J01CF BETA-LACTAMASE RESISTANT PENICILLINS**

J01CF02 CLOXACILLIN

Cap	Orl	250 mg	Jamp Cloxacillin	02510731	JPC	ACDEFGVW
			Teva-Cloxacillin	00337765	TEV	ACDEFGVW
Cap	Orl	500 mg	Jamp Cloxacillin	02510758	JPC	ACDEFGVW
			Teva-Cloxacillin	00337773	TEV	ACDEFGVW
Pws	Inj	2 g	Cloxacillin	02367424	STR	ACDEFGVW
Pws	Orl	25 mg	Teva-Cloxacillin	00337757	TEV	ACDEFGVW

**J01CR COMBINATIONS PENICILLINS INCLUDING BETA LACTAMASE INHIBITORS**

**J01CR02 AMOXICILLIN AND ENZYME INHIBITOR  
AMOXICILLIN / CLAVULANIC ACID**

Pws	Orl	125 mg / 31.25 mg / 5 mL	Clavulin	01916882	GSK	ABCDEFGVW
Pws	Orl	200 mg / 28.5 mg / 5 mL	Clavulin 200	02238831	GSK	ABCDEFGVW
Pws	Orl	250 mg / 62.5 mg / 5 mL	Clavulin-250 F	01916874	GSK	ABCDEFGVW
Pws	Orl	400 mg / 57 mg / 5 mL	Clavulin 400	02238830	GSK	ABCDEFGVW
			M-Amoxi Clav	02530694	MRA	ABCDEFGVW
Tab	Orl	250 mg / 125 mg	Apo-Amoxi Clav	02243350	APX	ABCDEFGVW
			Auro-Amoxi Clav	02471671	ARO	ABCDEFGVW
			Jamp Amoxi Clav	02508249	JPC	ABCDEFGVW
Tab	Orl	500 mg / 125 mg	Clavulin-500 F	01916858	GSK	ABCDEFGVW
			Apo-Amoxi Clav	02243351	APX	ABCDEFGVW
			Auro-Amoxi Clav	02471698	ARO	ABCDEFGVW
			Jamp Amoxi Clav	02508257	JPC	ABCDEFGVW
			Sandoz Amoxi-Clav	02482576	SDZ	ABCDEFGVW
Tab	Orl	875 mg / 125 mg	Clavulin	02238829	GSK	ABCDEFGVW
			Apo-Amoxi Clav	02245623	APX	ABCDEFGVW
			Auro-Amoxi Clav	02471701	ARO	ABCDEFGVW
			Jamp Amoxi Clav	02508265	JPC	ABCDEFGVW
			Sandoz Amoxi-Clav	02482584	SDZ	ABCDEFGVW

**J01CR05 PIPERACILLIN AND ENZYME INHIBITOR  
PIPERACILLIN / TAZOBACTAM**

Pws	Inj	2 g / 0.25 g	Piperacillin and Tazobactam	02308444	APX	ACDEFGVW
			Piperacillin and Tazobactam	02401312	HIK	ACDEFGVW
			Piperacillin and Tazobactam	02299623	SDZ	ACDEFGVW
			Piperacillin and Tazobactam	02362619	STR	ACDEFGVW
Pws	Inj	3 g / 0.375 g	Piperacillin and Tazobactam	02308452	APX	ACDEFGVW
			Piperacillin and Tazobactam	02401320	HIK	ACDEFGVW
			Piperacillin and Tazobactam	02299631	SDZ	ACDEFGVW
			Piperacillin and Tazobactam	02362627	STR	ACDEFGVW
			Piperacillin/Tazobactam	02370166	TEV	ACDEFGVW



J01CR05 PIPERACILLIN AND ENZYME INHIBITOR  
 PIPERACILLIN / TAZOBACTAM

Pws	Inj	4 g / 0.5 g	Piperacillin and Tazobactam	02308460	APX	ACDEFGVW
			Piperacillin and Tazobactam	02401339	HIK	ACDEFGVW
			Piperacillin and Tazobactam	02299658	SDZ	ACDEFGVW
			Piperacillin and Tazobactam	02362635	STR	ACDEFGVW
			Piperacillin/Tazobactam	02370174	TEV	ACDEFGVW
Pws	Inj	12 g / 1.5 g	Piperacillin and Tazobactam	02330547	SDZ	ACDEFGVW
			Piperacillin and Tazobactam	02377748	STR	ACDEFGVW

**J01D OTHER BETA LACTAM ANTIBACTERIALS**

**J01DB FIRST GENERATION CEPHALOSPORINS**

J01DB01 CEPHALEXIN

Cap	Orl	250 mg	Teva-Cephalexin	00342084	TEV	ABCDEFGHIW
Cap	Orl	500 mg	Teva-Cephalexin	00342114	TEV	ABCDEFGHIW
Pws	Orl	25 mg	Teva-Cephalexin	00342106	TEV	ABCDEFGHIW
Pws	Orl	50 mg	Teva-Cephalexin	00342092	TEV	ABCDEFGHIW
Tab	Orl	250 mg	Apo-Cephalex	00768723	APX	ABCDEFGHIW
			Auro-Cephalexin	02470578	ARO	ABCDEFGHIW
			Cephalexin	02521253	SAS	ABCDEFGHIW
			Jamp Cephalexin	02494698	JPC	ABCDEFGHIW
			Teva-Cephalexin	00583413	TEV	ABCDEFGHIW
Tab	Orl	500 mg	Apo-Cephalex	00768715	APX	ABCDEFGHIW
			Auro-Cephalexin	02470586	ARO	ABCDEFGHIW
			Cephalexin	02521261	SAS	ABCDEFGHIW
			Cephalexin	02495651	SIV	ABCDEFGHIW
			Jamp Cephalexin	02494701	JPC	ABCDEFGHIW
			Teva-Cephalexin	00583421	TEV	ABCDEFGHIW

J01DB04 CEFAZOLIN

Pws	Inj	500 mg	Cefazolin for Injection	02108119	TEV	ACDEFGVW
			Cefazolin Sodium	02308932	SDZ	ACDEFGVW
Pws	Inj	1 g	Cefazolin for Injection	02108127	TEV	ACDEFGVW
			Cefazolin Sodium	02308959	SDZ	ACDEFGVW

J01DB04	CEFAZOLIN								
Pws	Inj	10 g				Cefazolin for Injection	02437120	HIK	ACDEFGVW
						Cefazolin for Injection	02108135	TEV	ACDEFGVW
						Cefazolin for Injection USP	02465477	STR	ACDEFGVW

J01DB05	CEFADROXIL								
Cap	Orl	500 mg				Apo-Cefadroxil	02240774	APX	ACDEFGVW
						Teva-Cefadroxil	02235134	TEV	ACDEFGVW

**J01DC SECOND GENERATION CEPHALOSPORINS**

J01DC01	CEFOXITIN								
Pws	Inj	1 g				Cefoxitin Sodium	02128187	TEV	ACDEFGVW
Pws	Inj	2 g				Cefoxitin Sodium	02128195	TEV	ACDEFGVW

J01DC02	CEFUROXIME								
Liq	Orl	125 mg/mL				Ceftin	02212307	SDZ	ABCDEFGHIJ
Pws	Inj	750 mg				Cefuroxime	02241638	FKB	ACDEFGVW
Pws	Inj	1.5 g				Cefuroxime	02241639	FKB	ACDEFGVW
Tab	Orl	250 mg				Apo-Cefuroxime	02244393	APX	ABCDEFGHIJ
						Auro-Cefuroxime	02344823	ARO	ABCDEFGHIJ
Tab	Orl	500 mg				Apo-Cefuroxime	02244394	APX	ABCDEFGHIJ
						Auro-Cefuroxime	02344831	ARO	ABCDEFGHIJ

J01DC10	CEFPROZIL								
Pws	Orl	25 mg				Taro-Cefprozil	02329204	SUN	ACDEFGVW
Pws	Orl	50 mg				Taro-Cefprozil	02293579	SUN	ACDEFGVW
Tab	Orl	250 mg				Taro-Cefprozil	02293528	SUN	ACDEFGVW
Tab	Orl	500 mg				Auro-Cefprozil	02347253	ARO	ACDEFGVW
						Taro-Cefprozil	02293536	SUN	ACDEFGVW

**J01DD THIRD GENERATION CEPHALOSPORINS**

J01DD01	CEFOTAXIME								
Pws	Inj	1 g				Cefotaxime Sodium	02434091	STR	ACDEFGVW
Pws	Inj	2 g				Cefotaxime Sodium	02434105	STR	ACDEFGVW

J01DD02	CEFTAZIDIME								
Pws	Inj	1 g		Ceftazidime	00886971	FKB	ACDEFGVW		
Pws	Inj	2 g		Ceftazidime	00886955	FKB	ACDEFGVW		
Pws	Inj	6 g		Ceftazidime for Injection	02437864	STR	ACDEFGVW		

J01DD04	CEFTRIAZONE								
Pws	Inj	250 mg		Ceftriaxone Sodium	02325594	STR	ACDEFGVW		
Pws	Inj	1 g		Ceftriaxone Sodium	02325616	STR	ACDEFGVW		
				Ceftriaxone Sodium	02287633	TEV	ACDEFGVW		
				Ceftriaxone Sodium for Injection	02292270	SDZ	ACDEFGVW		
Pws	Inj	2 g		Ceftriaxone Sodium	02325624	STR	ACDEFGVW		
				Ceftriaxone Sodium for Injection	02292289	SDZ	ACDEFGVW		
Pws	Inj	10 g		Ceftriaxone Sodium for Injection	02292297	SDZ	ACDEFGVW		
				Ceftriaxone Sodium for Injection	02325632	STR	ACDEFGVW		

J01DD08	CEFIXIME								
Pws	Orl	20 mg		Suprax	00868965	ODN	ACDEFGVW		
				Auro-Cefixime	02468689	ARO	ACDEFGVW		
Tab	Orl	400 mg		Suprax	00868981	ODN	ACDEFGVW		
				Auro-Cefixime	02432773	ARO	ACDEFGVW		

**J01DE FOURTH GENERATION CEPHALOSPORINS**

J01DE01	CEFEPIME								
Pws	Inj	1 g		Apo-Cefepime	02467496	APX	ACDEFGVW		
Pws	Inj	2 g		Apo-Cefepime	02467518	APX	ACDEFGVW		

**J01DF MONOBACTAMS**

J01DF01	AZTREONAM								
Pwr	Inh	75 mg		Cayston	02329840	GIL	(SA)		

**J01DH CARBAPENEMS**

J01DH02	MEROPENEM								
Pws	Inj	500 mg		Meropenem	02378787	SDZ	ACDEFGVW		
				Meropenem for Injection	02493330	STR	ACDEFGVW		
				Taro-Meropenem	02421518	SUN	ACDEFGVW		

J01DH02	MEROPENEM								
Pws	Inj	1 g		Meropenem for Injection	02378795	SDZ	ACDEFGVW		
				Meropenem for Injection	02493349	STR	ACDEFGVW		
				Taro-Meropenem	02421526	SUN	ACDEFGVW		
J01DH03	ERTAPENEM								
Pws	Inj	1 g		Invanz	02247437	FRS	ACDEFGVW		
J01DH51	IMIPENEM AND ENZYME INHIBITOR								
				IMIPENEM / CILASTATIN					
Pws	Inj	250 mg / 250 mg		Taro-Imipenem-Cilastatin	02351692	SUN	ACDEFGVW		
Pws	Inj	500 mg / 500 mg		Taro-Imipenem-Cilastatin	02351706	SUN	ACDEFGVW		
<b>J01DI</b>	<b>OTHER CEPHALOSPORINS AND PENEMS</b>								
J01DI54	CEFTOLOZANE AND BETA-LACTAMASE INHIBITOR								
				CEFTOLOZANE / TAZOBACTAM					
Pws	IV	1 g / 0.5 g		Zerbaxa	02446901	FRS	W (SA)		
<b>J01E</b>	<b>SULFONAMIDES AND TRIMETHOPRIM</b>								
<b>J01EA</b>	<b>TRIMETHOPRIM AND DERIVATIVES</b>								
J01EA01	TRIMETHOPRIM								
Tab	Orl	100 mg		Trimethoprim	02243116	AAP	ACDEFGV		
Tab	Orl	200 mg		Trimethoprim	02243117	AAP	ACDEFGV		
<b>J01EE</b>	<b>COMBINATIONS OF SULFONAMIDES AND TRIMETHOPRIM, INCLUDING DERIVATIVES</b>								
J01EE01	SULFAMETHOXASOLE AND TRIMETHOPRIM								
Sus	Orl	40 mg / 8 mg		Teva-Trimel	00726540	TEV	ABCDEFGVW		
Tab	Orl	100 mg / 20 mg		Sulfatrim	00445266	AAP	ABCDEFGVW		
Tab	Orl	400 mg / 80 mg		Sulfatrim	00445274	AAP	ABCDEFGVW		
				Teva-Trimel	00510637	TEV	ABCDEFGVW		
Tab	Orl	800 mg / 160 mg		Sulfatrim DS	00445282	AAP	ABCDEFGVW		
<b>J01F</b>	<b>MACROLIDES, LINCOSAMIDES AND STREPTOGRAMINS</b>								
<b>J01FA</b>	<b>MACROLIDES</b>								
J01FA01	ERYTHROMYCIN								
ECC	Orl	333 mg		Eryc (Disc/non disp Apr 27/24)	00873454	PFI	ACDEFGVW		

J01FA01	ERYTHROMYCIN							
Tab	Orl	250 mg		Erythro	00682020	AAP	ACDEFGVW	
J01FA02	SPIRAMYCIN							
Cap	Orl	750 000 IU		Rovamycine 250	01927825	ODN	ACDEFGVW	
Cap	Orl	1 500 000 IU		Rovamycine 500	01927817	ODN	ACDEFGVW	
J01FA09	CLARITHROMYCIN							
ERT	Orl	500 mg		Act Clarithromycin XL	02403196	TEV	ACDEFGVW	
				Apo-Clarithromycin XL	02413345	APX	ACDEFGVW	
Pws	Orl	125 mg / 5 mL		Biaxin	02146908	ABB	ACDEFGVW	
				Clarithromycin	02408988	SAS	ACDEFGVW	
				Taro-Clarithromycin	02390442	TAR	ACDEFGVW	
Pws	Orl	250 mg / 5 mL		Biaxin	02244641	ABB	ACDEFGVW	
				Clarithromycin	02408996	SAS	ACDEFGVW	
				Taro-Clarithromycin	02390450	TAR	ACDEFGVW	
Tab	Orl	250 mg		Biaxin BID	01984853	ABB	ACDEFGVW	
				Apo-Clarithromycin	02274744	APX	ACDEFGVW	
				Clarithromycin	02466120	SAS	ACDEFGVW	
				Clarithromycin	02442469	SIV	ACDEFGVW	
				M-Clarithromycin	02471388	MRA	ACDEFGVW	
				pms-Clarithromycin	02247573	PMS	ACDEFGVW	
				Sandoz Clarithromycin	02266539	SDZ	ACDEFGVW	
				Taro-Clarithromycin	02361426	SUN	ACDEFGVW	
Tab	Orl	500 mg		Biaxin BID	02126710	ABB	ACDEFGVW	
				Apo-Clarithromycin	02274752	APX	ACDEFGVW	
				Clarithromycin	02466139	SAS	ACDEFGVW	
				Clarithromycin	02442485	SIV	ACDEFGVW	
				M-Clarithromycin	02471396	MRA	ACDEFGVW	
				pms-Clarithromycin	02247574	PMS	ACDEFGVW	
				Sandoz Clarithromycin	02266547	SDZ	ACDEFGVW	
				Taro-Clarithromycin	02361434	SUN	ACDEFGVW	
J01FA10	AZITHROMYCIN							
Pws	Inj	500 mg		Zithromax	02239952	PFI	ACDEFGVW	

J01FA10 AZITHROMYCIN

Pws Orl 100 mg / 5 mL

Zithromax 02223716 PFI ABCDEFGVW  
Auro-Azithromycin 02482363 ARO ABCDEFGVW  
Sandoz Azithromycin 02332388 SDZ ABCDEFGVW

Pws Orl 200 mg / 5 mL

Zithromax 02223724 PFI ABCDEFGVW  
Auro-Azithromycin 02482371 ARO ABCDEFGVW  
Sandoz Azithromycin 02332396 SDZ ABCDEFGVW

Tab Orl 250 mg

Zithromax 02212021 PFI ABCDEFGVW  
Apo-Azithromycin Z 02415542 APX ABCDEFGVW  
Azithromycin 02330881 SAS ABCDEFGVW  
Azithromycin 02442434 SIV ABCDEFGVW  
Jamp-Azithromycin 02452308 JPC ABCDEFGVW  
M-Azithromycin 02502038 MRA ABCDEFGVW  
Mar-Azithromycin 02452324 MAR ABCDEFGVW  
NRA-Azithromycin 02479680 NRA ABCDEFGVW  
pms-Azithromycin 02261634 PMS ABCDEFGVW  
Riva-Azithromycin 02275309 RIV ABCDEFGVW  
Sandoz Azithromycin 02265826 SDZ ABCDEFGVW  
Teva-Azithromycin 02267845 TEV ABCDEFGVW  
  
pms-Azithromycin 02261642 PMS (SA)

Tab Orl 600 mg

J01FF LINCOSAMIDES

J01FF01 CLINDAMYCIN

Cap Orl 150 mg

Dalacin C 00030570 PFI ACDEFGVW  
Auro-Clindamycin 02436906 ARO ACDEFGVW  
Clindamycin 02400529 SAS ACDEFGVW  
Jamp-Clindamycin 02483734 JPC ACDEFGVW  
M-Clindamycin 02479923 MRA ACDEFGVW  
Med-Clindamycin 02462656 GMP ACDEFGVW  
NRA-Clindamycin 02493748 NRA ACDEFGVW  
Riva-Clindamycin 02468476 RIV ACDEFGVW  
Teva-Clindamycin 02241709 TEV ACDEFGVW

**J01FF01 CLINDAMYCIN**

Cap Orl 300 mg

Dalacin C	02182866	PFI	ACDEFGVW
Auro-Clindamycin	02436914	ARO	ACDEFGVW
Clindamycin	02400537	SAS	ACDEFGVW
Jamp-Clindamycin	02483742	JPC	ACDEFGVW
M-Clindamycin	02479931	MRA	ACDEFGVW
Med-Clindamycin	02462664	GMP	ACDEFGVW
NRA-Clindamycin	02493756	NRA	ACDEFGVW
Riva-Clindamycin	02468484	RIV	ACDEFGVW
Teva-Clindamycin	02241710	TEV	ACDEFGVW

Liq Inj 150 mg/mL

Dalacin C Phosphate	00260436	PFI	ACDEFGVW
Clindamycin (2mL, 4mL, 6mL vials)	02230540	SDZ	ACDEFGVW
Clindamycin (bulk vials)	02230535	SDZ	ACDEFGVW

Pws Orl 75 mg / 5 mL

Dalacin C 00225851 PFI ACDEFGVW

**J01G AMINOGLYCOSIDE ANTIBACTERIALS****J01GB OTHER AMINOGLYCOSIDES****J01GB01 TOBRAMYCIN**

Liq Inh 300 mg / 5 mL

Tobi	02239630	BGP	ABCDEFVW
Teva-Tobramycin	02389622	TEV	ABCDEFVW

Liq Inj 40 mg/mL

Tobramycin (PF) 02241210 SDZ ABCDEFGVW

Pwr Inh 28 mg

Tobi Podhaler 02365154 BGP (SA)

**J01GB03 GENTAMICIN**

Liq Inj 40 mg/mL

Gentamicin 02242652 SDZ ACDEFGVW

**J01GB06 AMIKACIN**

Liq Inj 250 mg/mL

Amikacin 02242971 SDZ ACDEFGPVW

**J01M QUINOLONE ANTIBACTERIALS****J01MA FLUOROQUINOLONES****J01MA02 CIPROFLOXACIN**

Liq IV 2 mg/mL

Ciprofloxacin Intravenous Infusion BP 02304759 SDZ ACDEFGVW

Liq Orl 500 mg / 5 mL

Cipro Oral Suspension 02237514 BAY W (SA)

J01MA02 CIPROFLOXACIN

Tab Orl 250 mg

Act Ciprofloxacin 02247339 TEV BW (SA)  
 Auro-Ciprofloxacin 02381907 ARO BW (SA)  
 Ciprofloxacin 02353318 SAS BW (SA)  
 Ciprofloxacin 02386119 SIV BW (SA)  
 Jamp-Ciprofloxacin 02380358 JPC BW (SA)  
 Mar-Ciprofloxacin 02379686 MAR BW (SA)  
 pms-Ciprofloxacin 02248437 PMS BW (SA)  
 Sandoz Ciprofloxacin 02248756 SDZ BW (SA)  
 Taro-Ciproflox 02303728 SUN BW (SA)

Tab Orl 500 mg

Act Ciprofloxacin 02247340 TEV BW (SA)  
 Auro-Ciprofloxacin 02381923 ARO BW (SA)  
 Ciprofloxacin 02353326 SAS BW (SA)  
 Ciprofloxacin 02386127 SIV BW (SA)  
 Jamp-Ciprofloxacin 02380366 JPC BW (SA)  
 Mar-Ciprofloxacin 02379694 MAR BW (SA)  
 NRA-Ciprofloxacin 02492008 NRA BW (SA)  
 pms-Ciprofloxacin 02248438 PMS BW (SA)  
 Sandoz Ciprofloxacin 02248757 SDZ BW (SA)  
 Taro-Ciproflox 02303736 SUN BW (SA)

Tab Orl 750 mg

Act Ciprofloxacin 02247341 TEV BW (SA)  
 Jamp-Ciprofloxacin 02380374 JPC BW (SA)  
 Mar-Ciprofloxacin 02379708 MAR BW (SA)  
 pms-Ciprofloxacin 02248439 PMS BW (SA)  
 Sandoz Ciprofloxacin 02248758 SDZ BW (SA)  
 Taro-Ciproflox 02303744 SUN BW (SA)

J01MA06 NORFLOXACIN

Tab Orl 400 mg

Norfloxacin 02229524 AAP ACDEFGVW

J01MA12 LEVOFLOXACIN

Liq Inh 240 mg / 2.4 mL

Quinsair 02442302 HRZ (SA)

Liq Inj 5 mg/mL

Levofloxacin 02314932 PFI W

Tab Orl 250 mg

Act Levofloxacin 02315424 TEV BVW (SA)  
 Apo-Levofloxacin 02284707 APX BVW (SA)  
 Mint-Levofloxacin 02505797 MNT BVW (SA)  
 Sandoz Levofloxacin 02298635 SDZ BVW (SA)



J01MA12 LEVOFLOXACIN

Tab Orl 500 mg

Act Levofloxacin 02315432 TEV BVW (SA)  
Apo-Levofloxacin 02284715 APX BVW (SA)  
Mint-Levofloxacin 02505819 MNT BVW (SA)  
Sandoz Levofloxacin 02298643 SDZ BVW (SA)

Tab Orl 750 mg

Apo-Levofloxacin 02325942 APX BVW (SA)  
Sandoz Levofloxacin 02298651 SDZ BVW (SA)

J01MA14 MOXIFLOXACIN

Tab Orl 400 mg

Apo-Moxifloxacin 02404923 APX BVW (SA)  
Auro-Moxifloxacin 02432242 ARO BVW (SA)  
Jamp-Moxifloxacin 02443929 JPC BVW (SA)  
Jamp-Moxifloxacin 02447061 JPC BVW (SA)  
M-Moxifloxacin 02472791 MRA BVW (SA)  
Mar-Moxifloxacin 02447053 MAR BVW (SA)  
Med-Moxifloxacin 02457814 GMP BVW (SA)  
Moxifloxacin 02520710 SAS BVW (SA)  
Sandoz Moxifloxacin 02383381 SDZ BVW (SA)  
Teva-Moxifloxacin 02375702 TEV BVW (SA)

J01X OTHER ANTIBACTERIALS

J01XA GLYCOPEPTIDE ANTIBACTERIALS

J01XA01 VANCOMYCIN

Cap Orl 125 mg

Vancocin 00800430 SLP ACDEFGVW  
Jamp-Vancomycin 02407744 JPC ACDEFGVW

Pws Inj 500 mg

Sterile Vancomycin 02230191 PFI ABCDEFGVW  
Vancomycin Hydrochloride 02502593 JPC ABCDEFGVW  
Vancomycin Hydrochloride USP 02342855 STR ABCDEFGVW  
Vancomycin 02394626 SDZ ABCDEFGVW

Pws Inj 1 g

Jamp-Vancomycin (Disc/non disp Nov 16/23) 02420309 JPC ABCDEFGVW  
Vancomycin 02394634 SDZ ABCDEFGVW  
Vancomycin 02342863 STR ABCDEFGVW  
Vancomycin Hydrochloride 02502607 JPC ABCDEFGVW

Pws Inj 5g

Vancomycin Hydrochloride 02394642 SDZ ABCDEFGVW  
Vancomycin Hydrochloride 02405822 STR ABCDEFGVW

J01XB POLYMYXINS

J01XB01	COLISTIN							
	Pws	IM	150 mg		Coly-Mycin M Parenteral	00476420	ERF	ACDEFGV
<b>J01XD</b>	<b>IMIDAZOLE DERIVATIVES</b>							
J01XD01	METRONIDAZOLE							
	Liq	Inj	5 mg/mL		Metronidazole	00870420	BAX	ACDEFGVW
					Metronidazole	00649074	PFI	ACDEFGVW
	Tab	Orl	250 mg		Metronidazole	00545066	AAP	ACDEFGVW
<b>J01XE</b>	<b>NITROFURAN DERIVATIVES</b>							
J01XE01	NITROFURANTOIN							
	Cap	Orl	50 mg		Teva-Nitrofurantoin	02231015	TEV	ACDEFGV
	Cap	Orl	100 mg		pms-Nitrofurantoin	02455676	PMS	ACDEFGV
	Tab	Orl	50 mg		Nitrofurantoin	00319511	AAP	ACDEFGV
	Tab	Orl	100 mg		Nitrofurantoin	00312738	AAP	ACDEFGV
<b>J01XX</b>	<b>OTHER ANTIBACTERIALS</b>							
J01XX01	FOSFOMYCIN							
	Pws	Orl	3 g		Monurol	02240335	PAL	(SA)
					Jamp-Fosfomycin	02473801	JPC	(SA)
J01XX05	METHENAMINE							
	Tab	Orl	500 mg		Mandelamine	00499013	SLP	ACDEFGV
J01XX08	LINEZOLID							
	Tab	Orl	600 mg		Apo-Linezolid	02426552	APX	(SA)
					Jamp Linezolid	02520354	JPC	(SA)
					Sandoz Linezolid	02422689	SDZ	(SA)
J01XX09	DAPTOMYCIN							
	Pws	IV	500 mg / 10 mL		Cubicin RF	02465493	CBP	W (SA)
<b>J02</b>	<b>ANTIMYCOTICS FOR SYSTEMIC USE</b>							
<b>J02A</b>	<b>ANTIMYCOTICS FOR SYSTEMIC USE</b>							
<b>J02AA</b>	<b>ANTIBIOTICS</b>							
J02AA01	AMPHOTERICIN B							
	Pws	Inj	50 mg		AmBisome	02241630	ASL	ACDEFGVW
					Fungizone	00029149	XPI	ACDEFGVW

**J02AB IMIDAZOLE DERIVATIVES**

J02AB02 KETOCONAZOLE

Tab Orl 200 mg

Apo-Ketoconazole 02237235 APX ACDEFGVW

Teva-Ketoconazole 02231061 TEV ACDEFGVW

**J02AC TRIAZOLE DERIVATIVES**

J02AC01 FLUCONAZOLE

Cap Orl 150 mg

Diflucan 02141442 CHC ACDEFGVW

Apo-Fluconazole 02241895 APX ACDEFGVW

Fluconazole-150 02521229 SAS ACDEFGVW

Jamp Fluconazole 02432471 JPC ACDEFGVW

Mar-Fluconazole-150 02428792 MAR ACDEFGVW

Liq Inj 2 mg/mL

Diflucan 00891835 PFI ACDEFGVW

Pws Orl 50 mg / 5 mL

Diflucan 02024152 PFI (SA)

Tab Orl 50 mg

Act Fluconazole 02281260 TEV ACDEFGVW

Apo-Fluconazole 02237370 APX ACDEFGVW

Fluconazole 02517396 SAS ACDEFGVW

Mylan-Fluconazole 02245292 MYL ACDEFGVW

Novo-Fluconazole 02236978 TEV ACDEFGVW

pms-Fluconazole 02245643 PMS ACDEFGVW

Tab Orl 100 mg

Act Fluconazole 02281279 TEV ACDEFGVW

Apo-Fluconazole 02237371 APX ACDEFGVW

Fluconazole 02517418 SAS ACDEFGVW

Mylan-Fluconazole 02245293 MYL ACDEFGVW

Novo-Fluconazole 02236979 TEV ACDEFGVW

pms-Fluconazole 02245644 PMS ACDEFGVW

J02AC02 ITRACONAZOLE

Cap Orl 100 mg

Sporanox 02047454 JAN ACDEFGV

Mint-Itraconazole 02462559 MNT ACDEFGV

Liq Orl 10 mg/mL

Sporanox 02231347 JAN (SA)

Jamp-Itraconazole 02484315 JPC (SA)

Odan-Itraconazole 02495988 ODN (SA)

J02AC03 VORICONAZOLE

Pws Inj 200 mg

Voriconazole for Injection 02381966 SDZ ACDEFGV

J02AC03 VORICONAZOLE

Tab Orl 50 mg

Vfend 02256460 PFI (SA)

Jamp Voriconazole 02525771 JPC (SA)

Sandoz Voriconazole 02399245 SDZ (SA)

Teva-Voriconazole 02396866 TEV (SA)

Tab Orl 200 mg

Vfend 02256479 PFI (SA)

Jamp Voriconazole 02525798 JPC (SA)

Sandoz Voriconazole 02399253 SDZ (SA)

Teva-Voriconazole 02396874 TEV (SA)

J02AC05 ISAVUCONAZOLE

Cap Orl 100 mg

Cresemba 02483971 AVI (SA)

Pws IV 200 mg

Cresemba 02483998 AVI (SA)

**J02AX ANTIMYCOTICS FOR SYSTEMIC USE**

J02AX04 CASPOFUNGIN

Pwd Inj 50 mg

Candida IV 02244265 FRS ACDEFGVW

Caspofungin for Injection 02460947 JNO ACDEFGVW

Pwd Inj 70 mg

Candida IV 02244266 FRS ACDEFGVW

Caspofungin for Injection 02460955 JNO ACDEFGVW

J02AX05 MICAFUNGIN

Pws IV 50 mg

Mycamine 02294222 ASL ACDEFGVW

Pws IV 100 mg

Mycamine 02311054 ASL ACDEFGVW

**J04 ANTIMYCOBACTERIALS**

**J04A DRUGS FOR TREATMENT OF TUBERCULOSIS**

**J04AB ANTIBIOTICS**

J04AB02 RIFAMPICIN

Cap Orl 150 mg

Rofact 00393444 BSL ACDEFGPVW

Cap Orl 300 mg

Rofact 00343617 BSL ACDEFGPVW

J04AB04 RIFABUTIN

Cap Orl 150 mg

Mycobutin 02063786 PFI P (SA)

**J04AC HYDRAZIDES**

J04AC01 ISONIAZID

J04AC01	ISONIAZID							
	Syr	Orl	10 mg/mL		pdp-Isoniazid	00577812	PMS	P
	Tab	Orl	100 mg		pdp-Isoniazid	00577790	PMS	P
	Tab	Orl	300 mg		pdp-Isoniazid	00577804	PMS	P
<b>J04AK</b>	<b>OTHER DRUGS FOR TREATMENT OF TUBERCULOSIS</b>							
J04AK01	PYRAZINAMIDE							
	Tab	Orl	500 mg		pdp-Pyrazinamide	00618810	PMS	P
J04AK02	ETHAMBUTOL							
	Tab	Orl	100 mg		Etibi	00247960	BSL	ACDEFGPV
	Tab	Orl	400 mg		Etibi	00247979	BSL	ACDEFGPV
<b>J04B</b>	<b>DRUGS FOR TREATMENT OF LEPROSY</b>							
<b>J04BA</b>	<b>DRUGS FOR TREATMENT OF LEPROSY</b>							
J04BA02	DAPSONE							
	Tab	Orl	100 mg		Dapsone	02041510	JCB	ACDEFGV
					Mar-Dapsone	02481227	MAR	ACDEFGV
					Riva-Dapsone	02489058	RIV	ACDEFGV
<b>J05</b>	<b>ANTIVIRALS FOR SYSTEMIC USE</b>							
<b>J05A</b>	<b>DIRECT ACTING ANTIVIRALS</b>							
<b>J05AB</b>	<b>NUCLEOSIDES AND NUCLEOTIDES EXCLUDING REVERSE TRANSCRIPTASE INHIBITORS</b>							
J05AB01	ACYCLOVIR							
	Liq	Inj	25 mg/mL		Acyclovir Sodium	02236916	PFI	ACDEFGW
	Liq	Inj	50 mg/mL		Acyclovir Sodium	02236926	FKB	ACDEFGW
	Sus	Orl	200 mg / 5 mL		Zovirax	00886157	GSK	ACDEFGV
	Tab	Orl	200 mg		Apo-Acyclovir	02207621	APX	ACDEFGV
					Mint-Acyclovir	02524708	MNT	ACDEFGV
					Mylan-Acyclovir	02242784	MYL	ACDEFGV
					Teva-Acyclovir	02285959	TEV	ACDEFGV
	Tab	Orl	400 mg		Apo-Acyclovir	02207648	APX	ACDEFGV
					Mint-Acyclovir	02524716	MNT	ACDEFGV
					Mylan-Acyclovir	02242463	MYL	ACDEFGV
					Teva-Acyclovir	02285967	TEV	ACDEFGV

J05AB01	ACYCLOVIR								
Tab	Orl	800 mg							
						Apo-Acyclovir	02207656	APX	ACDEFGV
						Mint-Acyclovir	02524724	MNT	ACDEFGV
						Mylan-Acyclovir	02242464	MYL	ACDEFGV
						Teva-Acyclovir	02285975	TEV	ACDEFGV
J05AB06	GANCICLOVIR								
Pws	Inj	500 mg							
						Cytovene	02162695	MCK	ACDEFGV
J05AB09	FAMCICLOVIR								
Tab	Orl	125 mg							
						Famvir	02229110	NVR	ACDEFGV
						Act Famciclovir	02305682	TEV	ACDEFGV
						Apo-Famciclovir	02292025	APX	ACDEFGV
						pms-Famciclovir	02278081	PMS	ACDEFGV
						Sandoz Famciclovir	02278634	SDZ	ACDEFGV
Tab	Orl	250 mg							
						Famvir	02229129	NVR	ACDEFGV
						Act Famciclovir	02305690	TEV	ACDEFGV
						Apo-Famciclovir	02292041	APX	ACDEFGV
						pms-Famciclovir	02278103	PMS	ACDEFGV
						Sandoz Famciclovir	02278642	SDZ	ACDEFGV
Tab	Orl	500 mg							
						Famvir	02177102	NVR	ACDEFGV
						Act Famciclovir	02305704	TEV	ACDEFGV
						Apo-Famciclovir	02292068	APX	ACDEFGV
						Sandoz Famciclovir	02278650	SDZ	ACDEFGV
J05AB11	VALACYCLOVIR								
Tab	Orl	500 mg							
						Valtrex	02219492	GSK	ACDEFGV
						Apo-Valacyclovir	02295822	APX	ACDEFGV
						Auro-Valacyclovir	02405040	ARO	ACDEFGV
						Jamp Valacyclovir	02440598	JPC	ACDEFGV
						Jamp-Valacyclovir	02441454	JPC	ACDEFGV
						Mylan-Valacyclovir	02351579	MYL	ACDEFGV
						pms-Valacyclovir	02298457	PMS	ACDEFGV
						Sandoz Valacyclovir	02347091	SDZ	ACDEFGV
						Teva-Valacyclovir	02357534	TEV	ACDEFGV
						Valacyclovir	02454645	SAS	ACDEFGV
						Valacyclovir	02442000	SIV	ACDEFGV

J05AB11 VALACYCLOVIR

Tab Orl 1 000 mg

Valtrex 02246559 GSK ACDEFGV  
 Apo-Valacyclovir 02354705 APX ACDEFGV  
 Auro-Valacyclovir 02405059 ARO ACDEFGV  
 Mylan-Valacyclovir 02351560 MYL ACDEFGV  
 pms-Valacyclovir 02381230 PMS ACDEFGV  
 Valacyclovir 02519585 SAS ACDEFGV  
 Valacyclovir 02442019 SIV ACDEFGV

J05AB14 VALGANCICLOVIR

Pws Orl 50 mg/mL

Valcyte 02306085 XPI (SA)

Tab Orl 450 mg

Valcyte 02245777 XPI ACDEFGV  
 Auro-Valganciclovir 02435179 ARO ACDEFGV  
 Mint-Valganciclovir 02495457 MNT ACDEFGV  
 Teva-Valganciclovir 02413825 TEV ACDEFGV

**J05AE PROTEASE INHIBITORS**

J05AE03 RITONAVIR

Tab Orl 100 mg

Norvir 02357593 ABV ACDEFGUV

J05AE07 FOSAMPRENAVIR

Sus Orl 50 mg/mL

Telzir 02261553 VIV ACDEFGUV

Tab Orl 700 mg

Telzir 02261545 VIV ACDEFGUV

J05AE08 ATAZANAVIR

Cap Orl 150 mg

Jamp Atazanavir 02513102 JPC ACDEFGUV  
 Mylan-Atazanavir 02456877 MYL ACDEFGUV  
 Teva-Atazanavir 02443791 TEV ACDEFGUV

Cap Orl 200 mg

Reyataz 02248611 BRI ACDEFGUV  
 Jamp Atazanavir 02513110 JPC ACDEFGUV  
 Mylan-Atazanavir 02456885 MYL ACDEFGUV  
 Teva-Atazanavir 02443813 TEV ACDEFGUV

Cap Orl 300 mg

Reyataz 02294176 BRI ACDEFGUV  
 Jamp Atazanavir 02513129 JPC ACDEFGUV  
 Mylan-Atazanavir 02456893 MYL ACDEFGUV  
 Teva-Atazanavir 02443821 TEV ACDEFGUV

J05AE09 TIPRANAVIR

J05AE09	TIPRANA VIR						
Cap	Orl	250 mg		Aptivus	02273322	BOE	(SA)
J05AE10	DARUNA VIR						
Tab	Orl	75 mg		Prezista	02338432	JAN	ACDEFGUV
Tab	Orl	150 mg		Prezista	02369753	JAN	ACDEFGUV
Tab	Orl	600 mg		Prezista	02324024	JAN	ACDEFGUV
				Apo-Darunavir	02487241	APX	ACDEFGUV
				Auro-Darunavir	02486121	ARO	ACDEFGUV
				Darunavir	02521342	JPC	ACDEFGUV
				M-Darunavir (Disc/non disp Jun 20/24)	02522284	MRA	ACDEFGUV
Tab	Orl	800 mg		Prezista	02393050	JAN	ACDEFGUV
				Apo-Darunavir	02487268	APX	ACDEFGUV
				Auro-Darunavir	02486148	ARO	ACDEFGUV
				Darunavir	02521350	JPC	ACDEFGUV
				M-Darunavir (Disc/non disp Jul 5/24)	02522292	MRA	ACDEFGUV

**J05AF NUCLEOSIDE AND NUCLEOTIDE REVERSE TRANSCRIPTASE INHIBITORS**

J05AF01	ZIDO VUDINE						
Cap	Orl	100 mg		Apo-Zidovudine	01946323	APX	ACDEFGUV
Liq	Inj	10 mg/mL		Retrovir	01902644	VIV	ACDEFGUV
Syr	Orl	50 mg / 5 mL		Retrovir	01902652	VIV	ACDEFGUV
J05AF05	LAMI VUDINE						
Liq	Orl	10 mg/mL		3TC	02192691	VIV	ACDEFGUV
Tab	Orl	100 mg		Apo-Lamivudine HBV	02393239	APX	(SA)
				Jamp-Lamivudine HBV	02512467	JPC	(SA)
Tab	Orl	150 mg		3TC	02192683	VIV	ACDEFGUV
				Apo-Lamivudine	02369052	APX	ACDEFGUV
				Jamp Lamivudine	02507110	JPC	ACDEFGUV
Tab	Orl	300 mg		3TC	02247825	VIV	ACDEFGUV
				Apo-Lamivudine	02369060	APX	ACDEFGUV
				Jamp Lamivudine	02507129	JPC	ACDEFGUV



J05AF06	ABACAVIR					
	Liq	Orl	20 mg/mL		Ziagen	02240358 VIV ACDEFGUV
	Tab	Orl	300 mg		Ziagen	02240357 VIV ACDEFGUV
					Apo-Abacavir	02396769 APX ACDEFGUV
					Mint-Abacavir	02480956 MNT ACDEFGUV

J05AF07	TENOFIVIR DISOPROXIL					
	Tab	Orl	300 mg		Viread	02247128 GIL ACDEFGUV
					Apo-Tenofovir	02451980 APX ACDEFGUV
					Auro-Tenofovir	02460173 ARO ACDEFGUV
					Jamp-Tenofovir	02479087 JPC ACDEFGUV
					Mint-Tenofovir	02512939 MNT ACDEFGUV
					Mylan-Tenofovir Disoproxil	02452634 MYL ACDEFGUV
					Nat-Tenofovir	02472511 NAT ACDEFGUV
					pms-Tenofovir	02453940 PMS ACDEFGUV
					Tenofovir	02523922 SIV ACDEFGUV
					Tenofovir Disoproxil Fumarate	02512327 SAS ACDEFGUV
					Teva-Tenofovir	02403889 TEV ACDEFGUV

J05AF10	ENTECAVIR					
	Tab	Orl	0.5 mg		Baraclude	02282224 BRI ACDEFGV
					Apo-Entecavir	02396955 APX ACDEFGV
					Auro-Entecavir	02448777 ARO ACDEFGV
					Entecavir	02527154 SAS ACDEFGV
					Entecavir	02453797 STD ACDEFGV
					Jamp-Entecavir	02467232 JPC ACDEFGV
					Mint-Entecavir	02485907 MNT ACDEFGV
					pms-Entecavir	02430576 PMS ACDEFGV

**J05AG NON-NUCLEOSIDES REVERSE TRANSCRIPTASE INHIBITORS**

J05AG01	NEVIRAPINE					
	Tab	Orl	200 mg		Auro-Nevirapine	02318601 ARO ACDEFGUV
					Jamp-Nevirapine	02405776 JPC ACDEFGUV
					Mylan-Nevirapine	02387727 MYL ACDEFGUV

J05AG03	EFAVIRENZ					
	Cap	Orl	50 mg		Sustiva (Disc/non disp Mar 31/24)	02239886 BRI ACDEFGUV
	Cap	Orl	200 mg		Sustiva (Disc/non disp Mar 31/24)	02239888 BRI ACDEFGUV

J05AG03	EFAVIRENZ						
Tab	Orl	600 mg					
			Auro-Efavirenz	02418428	ARO	ACDEFGUV	
			Jamp-Efavirenz	02458233	JPC	ACDEFGUV	
			Mylan-Efavirenz	02381524	MYL	ACDEFGUV	
			Teva-Efavirenz	02389762	TEV	ACDEFGUV	

J05AG04	ETRAVIRINE						
Tab	Orl	100 mg					
			Intelence	02306778	JAN	(SA)	
Tab	Orl	200 mg					
			Intelence	02375931	JAN	(SA)	

J05AG05	RILPIVIRINE						
Tab	Orl	25 mg					
			Edurant	02370603	JAN	ACDEFGUV	

J05AG06	DORAVIRINE						
Tab	Orl	100 mg					
			Pifeltro	02481545	FRS	U (SA)	

**J05AH NEURAMINIDASE INHIBITORS**

J05AH01	ZANAMIVIR						
Pwr	Inh	5 mg					
			Relenza	02240863	GSK	(SA)	

J05AH02	OSELTAMIVIR						
Cap	Orl	30 mg					
			Tamiflu	02304848	HLR	(SA)	
			Jamp-Oseltamivir	02497409	JPC	(SA)	
			Mar-Oseltamivir	02497352	MAR	(SA)	
			Mint-Oseltamivir	02497441	MNT	(SA)	
			Nat-Oseltamivir	02472635	NAT	(SA)	
			Osetamivir	02504006	STD	(SA)	

Cap	Orl	45 mg					
			Tamiflu	02304856	HLR	(SA)	
			Mar-Oseltamivir	02497360	MAR	(SA)	
			Nat-Oseltamivir	02472643	NAT	(SA)	
			Osetamivir	02504014	STD	(SA)	

Cap	Orl	75 mg					
			Tamiflu	02241472	HLR	(SA)	
			Jamp-Oseltamivir	02497425	JPC	(SA)	
			Mar-Oseltamivir	02497379	MAR	(SA)	
			Mint-Oseltamivir	02497476	MNT	(SA)	
			Nat-Oseltamivir	02457989	NAT	(SA)	
			Osetamivir	02504022	STD	(SA)	

J05AH02 OSELTAMIVIR

Pws Orl 6 mg/mL

Tamiflu 02381842 HLR (SA)

Nat-Oseltamivir 02499894 NAT (SA)

**J05AJ INTEGRASE INHIBITORS**

J05AJ01 RALTEGRAVIR

Tab Orl 400 mg

Isentress 02301881 FRS ACDEFGUV

J05AJ03 DOLUTEGRAVIR

Tab Orl 50 mg

Tivicay 02414945 VIV ACDEFGUV

J05AJ04 CABOTEGRAVIR

Tab Orl 30 mg

Vocabria 02497204 VIV U (SA)

**J05AP ANTIVIRALS FOR TREATMENT OF HCV INFECTIONS**

J05AP01 RIBAVIRIN

Tab Orl 200 mg

Ibavyr (Disc/non disp Dec 31/23) 02439212 PDP (SA)

J05AP08 SOFOSBUVIR

Tab Orl 400 mg

Sovaldi 02418355 GIL (SA)

J05AP51 SOFOSBUVIR AND LEDIPASVIR

Tab Orl 400 mg / 90 mg

Harvoni 02432226 GIL (SA)

J05AP55 SOFOSBUVIR AND VELPATASVIR

Tab Orl 400 mg / 100 mg

Eplclusa 02456370 GIL (SA)

J05AP56 SOFOSBUVIR, VELPATASVIR AND VOXILAPREVIR

Tab Orl 400 mg / 100 mg / 100mg

Vosevi 02467542 GIL (SA)

J05AP57 GLECAPREVIR AND PIBRENTASVIR

Tab Orl 100 mg / 40 mg

Maviret 02467550 ABV (SA)

**J05AR ANTIVIRALS FOR TREATMENT OF HIV INFECTIONS, COMBINATIONS**

J05AR01 ZIDOVUDINE AND LAMIVUDINE

Tab Orl 300 mg / 150 mg

Combivir 02239213 VIV ACDEFGUV

Apo-Lamivudine/Zidovudine 02375540 APX ACDEFGUV

Auro-Lamivudine/Zidovudine 02414414 ARO ACDEFGUV

Jamp-Lamivudine/Zidovudine 02502801 JPC ACDEFGUV

J05AR02 LAMIVUDINE AND ABACAVIR

J05AR02	LAMIVUDINE AND ABACAVIR								
Tab	Orl	300 mg / 600 mg		Kivexa	02269341	VIV	ACDEFGUV		
				Apo-Abacavir-Lamivudine	02399539	APX	ACDEFGUV		
				Auro-Abacavir/Lamivudine	02454513	ARO	ACDEFGUV		
				Jamp Abacavir/Lamivudine	02497654	JPC	ACDEFGUV		
				Mylan-Abacavir/Lamivudine	02450682	MYL	ACDEFGUV		
				pms-Abacavir-Lamivudine	02458381	PMS	ACDEFGUV		
				Teva-Abacavir/Lamivudine	02416662	TEV	ACDEFGUV		
J05AR03	TENOFIVIR DISOPROXIL AND EMTRICITABINE								
Tab	Orl	300 mg / 200 mg		Truvada	02274906	GIL	ACDEFGUV		
				Apo- Emtricitabine-Tenofovir	02452006	APX	ACDEFGUV		
				Auro-Emtricitabine/Tenofovir	02490684	ARO	ACDEFGUV		
				Jamp-Emtricitabine-Tenofovir Disoproxil Fumarate	02487012	JPC	ACDEFGUV		
				Mint-Emtricitabine/Tenofovir	02521547	MNT	ACDEFGUV		
				Mylan-Emtricitabine/Tenofovir Disoproxil	02443902	MYL	ACDEFGUV		
				pms-Emtricitabine-Tenofovir	02461110	PMS	ACDEFGUV		
				Teva-Emtricitabine/Tenofovir	02399059	TEV	ACDEFGUV		
J05AR04	ZIDOVUDINE, LAMIVUDINE AND ABACAVIR								
Tab	Orl	300 mg / 150 mg / 300 mg		Apo-Abacavir-Lamivudine-Zidovudine	02416255	APX	ACDEFGUV		
J05AR06	EMTRICITABINE, TENOFIVIR DISOPROXIL AND EFAVIRENZ								
Tab	Orl	200 mg / 300 mg / 600 mg		Apo-Efavirenz/Emtricitabine/Tenofovir	02468247	APX	ACDEFGUV		
				Auro-Efavirenz-Emtricitabine-Tenofovir	02478404	ARO	ACDEFGUV		
				Jamp Efavirenz/Emtricitabine/Tenofovir Disoproxil Fumarate	02519461	JPC	ACDEFGUV		
				Mylan-Efavirenz/Emtricitabine/Tenofovir Disoproxil Fumarate	02461412	MYL	ACDEFGUV		
				pms-Efavirenz-Emtricitabine-Tenofovir	02487284	PMS	ACDEFGUV		
				Teva-Efavirenz/Emtricitabine/Tenofovir	02393549	TEV	ACDEFGUV		
J05AR08	EMTRICITABINE, TENOFIVIR DISOPROXIL AND RILPIVIRINE								
Tab	Orl	200 mg / 300 mg / 25 mg		Complera	02374129	GIL	ACDEFGUV		
J05AR09	EMTRICITABINE, TENOFIVIR DISOPROXIL, ELVITEGRAVIR AND COBICSTAT								
Tab	Orl	200 mg / 300 mg / 150 mg / 150 mg		Stribild	02397137	GIL	U (SA)		
J05AR10	LOPINAVIR AND RITONAVIR								
Liq	Orl	80 mg / 20 mg/mL		Kaletra Oral Solution	02243644	ABV	ACDEFGUV		
Tab	Orl	100 mg / 25 mg		Kaletra	02312301	ABV	ACDEFGUV		

J05AR10	LOPINA VIR AND RITONAVIR						
	Tab	Orl	200 mg / 50 mg	Kaletra Tab	02285533	ABV	ACDEFGUV
J05AR13	LAMIVUDINE, ABACAVIR AND DOLUTEGRA VIR						
	Tab	Orl	300 mg / 600 mg / 50 mg	Triumeq	02430932	VIV	ACDEFGUV
J05AR14	DARUNAVIR AND COBICSTAT						
	Tab	Orl	800 mg / 150 mg	Prezcobix	02426501	JAN	U (SA)
J05AR18	EMTRICITABINE, TENOFOVIR ALAFENAMIDE, ELVITEGRA VIR AND COBICSTAT						
	Tab	Orl	200 mg / 10 mg / 150 mg / 150 mg	Genvoya	02449498	GIL	U (SA)
J05AR19	EMTRICITABINE, TENOFOVIR ALAFENAMIDE AND RILPIVIRINE						
	Tab	Orl	200 mg / 25 mg / 25 mg	Odefsey	02461463	GIL	U (SA)
J05AR20	EMTRICITABINE, TENOFOVIR ALAFENAMIDE AND BICTEGRA VIR						
	Tab	Orl	200 mg / 25 mg / 50 mg	Biktarvy	02478579	GIL	U (SA)
J05AR21	DOLUTEGRA VIR AND RILPIVIRINE						
	Tab	Orl	50 mg / 25 mg	Juluca	02475774	VIV	U (SA)
J05AR24	LAMIVUDINE, TENOFOVIR DISOPROXIL AND DORAVIRINE						
	Tab	Orl	300 mg / 300 mg / 100 mg	Delstrigo	02482592	FRS	U (SA)
J05AR25	LAMIVUDINE AND DOLUTEGRA VIR						
	Tab	Orl	50 mg / 300 mg	Dovato	02491753	VIV	U (SA)
J05AR99	CABOTEGRA VIR AND RILPIRIVINE						
	Sus	IM	600 mg / 2 mL, 900 mg / 3mL	Cabenuva	02497220	VIV	U (SA)
				Cabenuva	02497247	VIV	U (SA)

**J05AX OTHER ANTIVIRALS**

J05AX09	MARA VIROC						
	Tab	Orl	150 mg	Celsentri	02299844	VIV	(SA)
	Tab	Orl	300 mg	Celsentri	02299852	VIV	(SA)
J05AX18	LETERMOVIR						
	Liq	IV	240 mg / 12 mL	Prevymis	02469367	FRS	(SA)
	Liq	IV	480 mg / 24 mL	Prevymis	02469405	FRS	(SA)

J05AX18 LETERMOVIR

Tab Orl 240 mg

Prevymis 02469375 FRS (SA)

Tab Orl 480 mg

Prevymis 02469383 FRS (SA)

**L ANTINEOPLASTIC AND IMMUNOMODULATING AGENTS**

**L01 ANTINEOPLASTIC AGENTS**

**L01A ALKYLATING AGENTS**

**L01AA NITROGEN MUSTARD ANALOGUES**

L01AA01 CYCLOPHOSPHAMIDE

Tab Orl 25 mg

Procytox 02241795 BAX ACDEFGV

Tab Orl 50 mg

Procytox 02241796 BAX ACDEFGV

L01AA02 CHLORAMBUCIL

Tab Orl 2 mg

Leukeran 00004626 APN ACDEFGV

L01AA03 MELPHALAN

Tab Orl 2 mg

Alkeran 00004715 APN ACDEFGV

**L01AB ALKYL SULPHONATES**

L01AB01 BUSULFAN

Tab Orl 2 mg

Myleran 00004618 APN ACDEFGV

**L01AD NITROSOUREAS**

L01AD02 LOMUSTINE

Cap Orl 10 mg

CeeNU 00360430 BRI ACDEFGV

Cap Orl 40 mg

CeeNU 00360422 BRI ACDEFGV

**L01AX OTHER ALKYLATING AGENTS**

L01AX03 TEMOZOLOMIDE

Cap Orl 5 mg

Temodal 02241093 FRS ACDEFGV

Jamp Temozolomide 02516799 JPC ACDEFGV

Taro-Temozolomide 02443473 TAR ACDEFGV

Teva-Temozolomide 02441160 TEV ACDEFGV

Cap Orl 20 mg

Temodal 02241094 FRS ACDEFGV

Jamp Temozolomide 02516802 JPC ACDEFGV

Taro-Temozolomide 02443481 TAR ACDEFGV

Teva-Temozolomide 02395274 TEV ACDEFGV

L01AX03 TEMOZOLOMIDE

Cap	Orl	100 mg		Temodal	02241095	FRS	ACDEFGV
				Jamp Temozolomide	02516810	JPC	ACDEFGV
				Taro-Temozolomide	02443511	TAR	ACDEFGV
				Teva-Temozolomide	02395282	TEV	ACDEFGV
Cap	Orl	140 mg		Temodal	02312794	FRS	ACDEFGV
				Jamp Temozolomide	02516829	JPC	ACDEFGV
				Taro-Temozolomide	02443538	TAR	ACDEFGV
				Teva-Temozolomide	02395290	TEV	ACDEFGV
Cap	Orl	250 mg		Temodal	02241096	FRS	ACDEFGV
				Jamp Temozolomide	02516845	JPC	ACDEFGV
				Taro-Temozolomide	02443554	TAR	ACDEFGV
				Teva-Temozolomide	02395312	TEV	ACDEFGV

**L01B ANTIMETABOLITES**

**L01BA FOLIC ACID ANALOGUES**

L01BA01 METHOTREXATE

Liq	IM	7.5 mg / 0.3 mL		Methotrexate Inj BP	02422166	PMS	ACDEFGV
Liq	IM	10 mg / 0.4 mL		Methotrexate Inj BP	02422174	PMS	ACDEFGV
Liq	IM	15 mg / 0.6 mL		Methotrexate Inj BP	02422182	PMS	ACDEFGV
Liq	IM	20 mg / 0.8 mL		Methotrexate Inj BP	02422190	PMS	ACDEFGV
Liq	IM	25 mg/mL		Methotrexate Inj BP	02422204	PMS	ACDEFGV
Liq	Inj	10 mg/mL		Methotrexate Inj USP	02182947	PFI	ACDEFGV
Liq	Inj	25 mg/mL		Methotrexate Inj USP	02182777	PFI	ACDEFGV
				Methotrexate Inj USP (PF)	02182955	PFI	ACDEFGV
				Methotrexate Injection BP	02464365	AHI	ACDEFGV
				Methotrexate Inj USP (PF)	02099705	TEV	ACDEFGV
Liq	SC	10 mg / 0.2 mL		Metोजect Subcutaneous	02454831	MDX	ACDEFGV
Liq	SC	12.5 mg / 0.25 mL		Metोजect Subcutaneous	02454750	MDX	ACDEFGV
Liq	SC	15 mg / 0.3 mL		Metोजect Subcutaneous	02454858	MDX	ACDEFGV
				Methotrexate Subcutaneous	02491311	AHI	ACDEFGV

L01BA01	METHOTREXATE						
Liq	SC	17.5 mg / 0.35 mL					
			Metoject Subcutaneous	02454769	MDX	ACDEFGV	
			Methotrexate Subcutaneous	02491338	AHI	ACDEFGV	
Liq	SC	20 mg / 0.4 mL					
			Metoject Subcutaneous	02454866	MDX	ACDEFGV	
			Methotrexate Subcutaneous	02491346	AHI	ACDEFGV	
Liq	SC	22.5 mg / 0.45 mL					
			Metoject Subcutaneous	02454777	MDX	ACDEFGV	
			Methotrexate Subcutaneous	02491354	AHI	ACDEFGV	
Liq	SC	25 mg / 0.5 mL					
			Metoject Subcutaneous	02454874	MDX	ACDEFGV	
			Methotrexate Subcutaneous	02491362	AHI	ACDEFGV	
Tab	Orl	2.5 mg					
			ACH-Methotrexate	02509067	AHI	ACDEFGV	
			Apo-Methotrexate	02182963	APX	ACDEFGV	
			Auro-Methotrexate	02524023	ARO	ACDEFGV	
			pms-Methotrexate	02170698	PMS	ACDEFGV	
Tab	Orl	10 mg					
			Methotrexate	02182750	PFI	ACDEFGV	

**L01BB PURINE ANALOGUES**

L01BB02	MERCAPTOPYRIMIDINE						
Tab	Orl	50 mg					
			Purinethol	00004723	TEV	ACDEFGV	
			Mercaptopurine	02415275	STR	ACDEFGV	
L01BB03	TIOGUANINE						
Tab	Orl	40 mg					
			Lanvis	00282081	APN	ACDEFGV	
L01BB05	FLUDARABINE						
Tab	Orl	10 mg					
			Fludara	02246226	SAV	(SA)	

**L01BC PYRIMIDINE ANALOGUES**

L01BC02	FLUOROURACIL						
Crm	Top	5%					
			Efudex	00330582	BSL	ACDEFGV	
L01BC06	CAPECITABINE						
Tab	Orl	150 mg					
			Xeloda (Disc/non disp Dec 7/23)	02238453	XPI	ACDEFGV	
			Ach-Capecitabine	02426757	AHI	ACDEFGV	
			Capecitabine	02519879	JPC	ACDEFGV	
			Capecitabine	02514982	SAS	ACDEFGV	
			Sandoz Capecitabine	02421917	SDZ	ACDEFGV	
			Taro-Capecitabine	02457490	TAR	ACDEFGV	



L01BC06	CAPECITABINE							
Tab	Orl	500 mg	Xeloda (Disc/non disp Dec 7/23)	02238454	XPI	ACDEFGV		
			Ach-Capecitabine	02426765	AHI	ACDEFGV		
			Capecitabine	02519887	JPC	ACDEFGV		
			Capecitabine	02514990	SAS	ACDEFGV		
			Mint-Capecitabine	02508028	MNT	ACDEFGV		
			Sandoz Capecitabine	02421925	SDZ	ACDEFGV		
			Taro-Capecitabine	02457504	TAR	ACDEFGV		
L01BC07	AZACITIDINE							
Tab	Orl	200 mg	Onureg	02510197	CEL	(SA)		
Tab	Orl	300 mg	Onureg	02510200	CEL	(SA)		
L01BC08	DECITABINE							
	DECITABINE / CEDAZURIDINE							
Tab	Orl	35 mg / 100 mg	Inqovi	02501600	OTS	(SA)		
L01BC52	FLUOROURACIL, COMBINATION							
	FLUOROURACIL / SALICYLIC ACID							
Liq	Top	0.5% / 10%	Actikerall	02428946	CIP	ACDEFGV		
L01BC59	TRIFLURIDINE, COMBINATION							
	TRIFLURIDINE / TIPIRACIL							
Tab	Orl	15 mg / 6.14 mg	Lonsurf	02472104	TAI	(SA)		
Tab	Orl	20 mg / 8.19 mg	Lonsurf	02472112	TAI	(SA)		
<b>L01C</b>	<b>PLANT ALKALOIDS AND OTHER NATURAL PRODUCTS</b>							
<b>L01CB</b>	<b>PODOPHYLLOTOXIN DERIVATIVES</b>							
L01CB01	ETOPOSIDE							
Cap	Orl	50 mg	Vepesid	00616192	XPI	ACDEFGV		
<b>L01E</b>	<b>PROTEIN KINASE INHIBITORS</b>							
<b>L01EA</b>	<b>BCR-ABL TYROSINE KINASE INHIBITORS</b>							
L01EA01	IMATINIB							

## L01EA01 IMATINIB

Tab Orl 100 mg

Gleevec	02253275	NVR	ACDEFGV
ACH-Imatinib	02490986	AHI	ACDEFGV
Apo-Imatinib	02355337	APX	ACDEFGV
Imatinib	02504596	SAS	ACDEFGV
Jamp Imatinib	02495066	JPC	ACDEFGV
Mint-Imatinib	02492334	MNT	ACDEFGV
Nat-Imatinib	02397285	NAT	ACDEFGV
pms-Imatinib	02431114	PMS	ACDEFGV
Teva-Imatinib	02399806	TEV	ACDEFGV

Tab Orl 400 mg

Gleevec	02253283	NVR	ACDEFGV
ACH-Imatinib	02490994	AHI	ACDEFGV
Apo-Imatinib	02355345	APX	ACDEFGV
Imatinib	02504618	SAS	ACDEFGV
Jamp Imatinib	02495074	JPC	ACDEFGV
Mint-Imatinib	02492342	MNT	ACDEFGV
Nat-Imatinib	02397293	NAT	ACDEFGV
pms-Imatinib	02431122	PMS	ACDEFGV
Teva-Imatinib	02399814	TEV	ACDEFGV

## L01EA02 DASATINIB

Tab Orl 20 mg

Sprycel	02293129	BRI	(SA)
Apo-Dasatinib	02470705	APX	(SA)
Reddy-Dasatinib	02514737	RCH	(SA)
Taro-Dasatinib	02499282	TAR	(SA)
Teva-Dasatinib	02478307	TEV	(SA)

Tab Orl 50 mg

Sprycel	02293137	BRI	(SA)
Apo-Dasatinib	02470713	APX	(SA)
Reddy-Dasatinib	02514745	RCH	(SA)
Taro-Dasatinib	02499304	TAR	(SA)
Teva-Dasatinib	02478315	TEV	(SA)

Tab Orl 70 mg

Sprycel	02293145	BRI	(SA)
Apo-Dasatinib	02481499	APX	(SA)
Reddy-Dasatinib	02514753	RCH	(SA)
Taro-Dasatinib	02499312	TAR	(SA)
Teva-Dasatinib	02478323	TEV	(SA)

L01EA02 DASATINIB

Tab Orl 80 mg

Sprycel 02360810 BRI (SA)  
 Apo-Dasatinib 02481502 APX (SA)  
 Reddy-Dasatinib 02514761 RCH (SA)  
 Taro-Dasatinib 02499320 TAR (SA)  
 Teva-Dasatinib 02478331 TEV (SA)

Tab Orl 100 mg

Sprycel 02320193 BRI (SA)  
 Apo-Dasatinib 02470721 APX (SA)  
 Reddy-Dasatinib 02514788 RCH (SA)  
 Taro-Dasatinib 02499339 TAR (SA)  
 Teva-Dasatinib 02478358 TEV (SA)

Tab Orl 140 mg

Sprycel 02360829 BRI (SA)  
 Reddy-Dasatinib 02514796 RCH (SA)  
 Taro-Dasatinib 02499347 TAR (SA)

L01EA03 NILOTINIB

Cap Orl 150 mg

Tasigna 02368250 NVR (SA)

Cap Orl 200 mg

Tasigna 02315874 NVR (SA)

L01EA04 BOSUTINIB

Tab Orl 100 mg

Bosulif 02419149 PFI (SA)

Tab Orl 500 mg

Bosulif 02419157 PFI (SA)

L01EA05 PONATINIB

Tab Orl 15 mg

Iclusig 02437333 TAK (SA)

Tab Orl 45 mg

Iclusig 02437341 TAK (SA)

L01EA06 ASCIMINIB

Tab Orl 20 mg

Scemblix 02528320 NVR (SA)

Tab Orl 40 mg

Scemblix 02528339 NVR (SA)

**L01EB EPIDERMAL GROWTH FACTOR RECEPTOR (EGFR) TYROSINE KINASE INHIBITORS**

L01EB02 ERLOTINIB

L01EB02	ERLOTINIB						
Tab	Orl	25 mg				Tarceva	02269007 HLR ACDEFGV
						Apo-Erlotinib	02461862 APX ACDEFGV
						Nat-Erlotinib	02483912 NAT ACDEFGV
						Teva-Erlotinib	02377691 TEV ACDEFGV
Tab	Orl	100 mg				Tarceva	02269015 HLR ACDEFGV
						Apo-Erlotinib	02461870 APX ACDEFGV
						Nat-Erlotinib	02483920 NAT ACDEFGV
						Teva-Erlotinib	02377705 TEV ACDEFGV
Tab	Orl	150 mg				Tarceva	02269023 HLR ACDEFGV
						Apo-Erlotinib	02461889 APX ACDEFGV
						Nat-Erlotinib	02483939 NAT ACDEFGV
						Teva-Erlotinib	02377713 TEV ACDEFGV
L01EB03	AFATINIB						
Tab	Orl	20 mg				Giotrif	02415666 BOE (SA)
Tab	Orl	30 mg				Giotrif	02415674 BOE (SA)
Tab	Orl	40 mg				Giotrif	02415682 BOE (SA)
L01EB04	OSIMERTINIB						
Tab	Orl	40 mg				Tagrisso	02456214 AZE (SA)
Tab	Orl	80 mg				Tagrisso	02456222 AZE (SA)
<b>L01EC</b>	<b>B-RAF SERINE-THREONINE KINASE (BRAF) INHIBITORS</b>						
L01EC01	VEMURAFENIB						
Tab	Orl	240 mg				Zelboraf	02380242 HLR (SA)
L01EC02	DABRAFENIB						
Cap	Orl	50 mg				Tafinlar	02409607 NVR (SA)
Cap	Orl	75 mg				Tafinlar	02409615 NVR (SA)
L01EC03	ENCORAFENIB						
Cap	Orl	75 mg				Braftovi	02513099 PFI (SA)
<b>L01ED</b>	<b>ANAPLASTIC LYMPHOMA KINASE (ALK) INHIBITORS</b>						
L01ED01	CRIZOTINIB						

L01ED01	CRIZOTINIB							
Cap	Orl	200 mg		Xalkori	02384256	PFI	(SA)	
Cap	Orl	250 mg		Xalkori	02384264	PFI	(SA)	
L01ED02	CERITINIB							
Cap	Orl	150 mg		Zykadia	02436779	NVR	(SA)	
L01ED03	ALECTINIB							
Cap	Orl	150 mg		Alecensaro	02458136	HLR	(SA)	
L01ED04	BRIGATINIB							
Tab	Orl	30 mg		Alunbrig	02479206	TAK	(SA)	
Tab	Orl	90 mg		Alunbrig	02479214	TAK	(SA)	
Tab	Orl	180 mg		Alunbrig (initiation pack)	02479230	TAK	(SA)	
Tab	Orl	180 mg		Alunbrig	02479222	TAK	(SA)	

**L01EE MITOGEN-ACTIVATED PROTEIN KINASE (MEK) INHIBITORS**

L01EE01	TRAMETINIB							
Tab	Orl	0.5 mg		Mekinist	02409623	NVR	(SA)	
Tab	Orl	2 mg		Mekinist	02409658	NVR	(SA)	
L01EE02	COBIMETINIB							
Tab	Orl	20 mg		Cotellic	02452340	HLR	(SA)	
L01EE03	BINIMETINIB							
Tab	Orl	15 mg		Mektovi	02513080	PFI	(SA)	

**L01EF CYCLIN-DEPENDENT KINASE (CDK) INHIBITORS**

L01EF01	PALBOCICLIB							
Cap	Orl	75 mg		Ibrance	02453150	PFI	(SA)	
Cap	Orl	100 mg		Ibrance	02453169	PFI	(SA)	
Cap	Orl	125 mg		Ibrance	02453177	PFI	(SA)	
Tab	Orl	75 mg		Ibrance	02493535	PFI	(SA)	
Tab	Orl	100 mg		Ibrance	02493543	PFI	(SA)	

L01EF01 PALBOCICLIB  
Tab Orl 125 mg Ibrance 02493551 PFI (SA)

L01EF02 RIBOCICLIB  
Tab Orl 200 mg Kisqali 02473569 NVR (SA)

**L01EG MAMMALIAN TARGET OF RAPAMYCIN (MTOR) KINASE INHIBITORS**

L01EG02 EVEROLIMUS  
Tab Orl 2.5 mg Afinitor 02369257 NVR (SA)  
Nat-Everolimus 02530090 NAT (SA)  
pms-Everolimus 02504677 PMS (SA)  
Sandoz Everolimus 02492911 SDZ (SA)  
Teva-Everolimus 02463229 TEV (SA)

Tab Orl 5 mg Afinitor 02339501 NVR (SA)  
Nat-Everolimus 02530104 NAT (SA)  
pms-Everolimus 02504685 PMS (SA)  
Sandoz Everolimus 02492938 SDZ (SA)  
Teva-Everolimus 02463237 TEV (SA)

Tab Orl 10 mg Afinitor 02339528 NVR (SA)  
Nat-Everolimus 02530120 NAT (SA)  
pms-Everolimus 02504693 PMS (SA)  
Sandoz Everolimus 02492946 SDZ (SA)  
Teva-Everolimus 02463253 TEV (SA)

**L01EH HUMAN EPIDERMAL GROWTH FACTOR RECEPTOR 2 (HER2) TYROSINE KINASE INHIBITORS**

L01EH01 LAPATINIB  
Tab Orl 250 mg Tykerb 02326442 NVR (SA)

L01EH03 TUCATINIB  
Tab Orl 50 mg Tukysa 02499827 SGC (SA)

Tab Orl 150 mg Tukysa 02499835 SGC (SA)

**L01EJ JANUS-ASSOCIATED KINASE (JAK) INHIBITORS**

L01EJ01 RUXOLITINIB  
Tab Orl 5 mg Jakavi 02388006 NVR (SA)

Tab Orl 10 mg Jakavi 02434814 NVR (SA)

Tab Orl 15 mg Jakavi 02388014 NVR (SA)

L01EJ01	RUXOLITINIB							
Tab	Orl	20 mg			Jakavi	02388022	NVR	(SA)
L01EJ02	FEDRATINIB							
Tab	Orl	100 mg			Inrebic	02502445	CEL	(SA)
<b>L01EK</b>	<b>VASCULAR ENDOTHELIAL GROWTH FACTOR RECEPTOR (VEGFR) TYROSINE KINASE INHIBITORS</b>							
L01EK01	AXITINIB							
Tab	Orl	1 mg			Inlyta	02389630	PFI	(SA)
Tab	Orl	5 mg			Inlyta	02389649	PFI	(SA)
<b>L01EL</b>	<b>BRUTON'S TYROSINE KINASE (BTK) INHIBITORS</b>							
L01EL01	IBRUTINIB							
Cap	Orl	140 mg			Imbruvica	02434407	JAN	(SA)
L01EL02	ACALABRUTINIB							
Cap	Orl	100 mg			Calquence	02491788	AZE	(SA)
L01EL03	ZANUBRUTINIB							
Cap	Orl	80 mg			Brukinsa	02512963	BGN	(SA)
<b>L01EM</b>	<b>PHOSPHATIDYLINOSITOL-3-KINASE (PI3K) INHIBITORS</b>							
L01EM01	IDELALISIB							
Tab	Orl	100 mg			Zydelig	02438798	GIL	(SA)
Tab	Orl	150 mg			Zydelig	02438801	GIL	(SA)
<b>L01EX</b>	<b>OTHER PROTEIN KINASE INHIBITORS</b>							
L01EX01	SUNITINIB							
Cap	Orl	12.5 mg			Sutent	02280795	PFI	(SA)
					Sandoz Sunitinib	02532840	SDZ	(SA)
					Taro-Sunitinib	02524058	TAR	(SA)
Cap	Orl	25 mg			Sutent	02280809	PFI	(SA)
					Sandoz Sunitinib	02532867	SDZ	(SA)
					Taro-Sunitinib	02524066	TAR	(SA)
Cap	Orl	50 mg			Sutent	02280817	PFI	(SA)
					Sandoz Sunitinib	02532883	SDZ	(SA)
					Taro-Sunitinib	02524082	TAR	(SA)

L01EX02	SORAFENIB							
	Tab	Orl	200 mg		Nexavar	02284227	BAY	(SA)
L01EX03	PAZOPANIB							
	Tab	Orl	200 mg		Votrient	02352303	NVR	(SA)
					pms-Pazopanib	02525666	PMS	(SA)
L01EX04	VANDETANIB							
	Tab	Orl	100 mg		Caprelsa	02378582	GZM	(SA)
	Tab	Orl	300 mg		Caprelsa	02378590	GZM	(SA)
L01EX05	REGORAFENIB							
	Tab	Orl	40 mg		Stivarga	02403390	BAY	(SA)
L01EX07	CABOZANTINIB							
	Tab	Orl	20 mg		Cabometyx	02480824	IPS	(SA)
	Tab	Orl	40 mg		Cabometyx	02480832	IPS	(SA)
	Tab	Orl	60 mg		Cabometyx	02480840	IPS	(SA)
L01EX08	LENVATINIB							
	Cap	Orl	4 mg/dose		Lenvima	02484056	EIS	(SA)
	Cap	Orl	8 mg/dose		Lenvima	02468220	EIS	(SA)
	Cap	Orl	10 mg/dose		Lenvima	02450321	EIS	(SA)
	Cap	Orl	12 mg/dose		Lenvima	02484129	EIS	(SA)
	Cap	Orl	14 mg/dose		Lenvima	02450313	EIS	(SA)
	Cap	Orl	20 mg/dose		Lenvima	02450305	EIS	(SA)
	Cap	Orl	24 mg/dose		Lenvima	02450291	EIS	(SA)
L01EX09	NINTEDANIB							
	Cap	Orl	100 mg		Ofev	02443066	BOE	(SA)
	Cap	Orl	150 mg		Ofev	02443074	BOE	(SA)
L01EX10	MIDOSTAURIN							



L01EX10	MIDOSTAURIN								
Cap	Orl	25 mg				Rydapt	02466236	NVR	(SA)
L01EX12	LAROTRECTINIB								
Cap	Orl	25 mg				Vittrakvi	02490315	BAY	(SA)
Cap	Orl	100 mg				Vittrakvi	02490323	BAY	(SA)
Liq	Orl	20 mg/mL				Vittrakvi	02490331	BAY	(SA)
<b>L01F</b>	<b>MONOCLONAL ANTIBODIES AND ANTIBODY DRUG CONJUGATES</b>								
<b>L01FA</b>	<b>CD20 (CLUSTERS OF DIFFERENTIATION 20) INHIBITORS</b>								
L01FA01	RITUXIMAB								
Liq	IV	10 mg/mL				Riximyo	02498316	SDZ	(SA)
						Ruxience	02495724	PFI	(SA)
						Truxima (10 mL)	02478382	TMP	(SA)
						Truxima (50 mL)	02478390	TMP	(SA)
<b>L01X</b>	<b>OTHER ANTINEOPLASTIC AGENTS</b>								
<b>L01XB</b>	<b>METHYLHYDRAZINES</b>								
L01XB01	PROCARBAZINE								
Cap	Orl	50 mg				Matulane	00012750	LDN	ACDEFGV
<b>L01XE</b>	<b>PROTEIN KINASE INHIBITORS</b>								
L01XE54	GILTERITINIB								
Tab	Orl	40 mg				Xospata	02495058	ASL	(SA)
L01XE56	ENTRECTINIB								
Cap	Orl	100 mg				Rozlytrek	02495007	HLR	(SA)
Cap	Orl	200 mg				Rozlytrek	02495015	HLR	(SA)
<b>L01XF</b>	<b>RETINOIDS FOR CANCER TREATMENT</b>								
L01XF01	TRETINOIN								
Cap	Orl	10 mg				Vesanoid	02145839	XPI	ACDEFGV
						Jamp Tretinoin	02520036	JPC	ACDEFGV
<b>L01XJ</b>	<b>HEDGEHOG PATHWAY INHIBITORS</b>								
L01XJ01	VISMODEGIB								
Cap	Orl	150 mg				Erivedge	02409267	HLR	(SA)
<b>L01XK</b>	<b>POLY (ADP-RIBOSE) POLYMERASE (PARP) INHIBITORS</b>								

L01XK01	OLAPARIB							
Tab	Orl	100 mg			Lynparza	02475200	AZE	(SA)
Tab	Orl	150 mg			Lynparza	02475219	AZE	(SA)
L01XK02	NIRAPARIB							
Cap	Orl	100 mg			Zejula	02489783	GSK	(SA)

**L01XX OTHER ANTINEOPLASTIC AGENTS**

L01XX05	HYDROXYCARBAMIDE (HYDROXYUREA)							
Cap	Orl	500 mg			Hydrea	00465283	XPI	ACDEFGV
					Apo-Hydroxyurea	02247937	APX	ACDEFGV
					Mylan-Hydroxyurea	02242920	MYL	ACDEFGV

L01XX35	ANAGRELIDE							
Cap	Orl	0.5 mg			Agrylin	02236859	TAK	ACDEFGV
					pms-Anagrelide	02274949	PMS	ACDEFGV

L01XX52	VENETOCLAX							
Kit	Orl	10 mg, 50 mg, 100 mg			Venclexta Starter Kit	02458063	ABV	(SA)
Tab	Orl	10 mg			Venclexta	02458039	ABV	(SA)
Tab	Orl	50 mg			Venclexta	02458047	ABV	(SA)
Tab	Orl	100 mg			Venclexta	02458055	ABV	(SA)

**L02 ENDOCRINE THERAPY**

**L02A HORMONES AND RELATED AGENTS**

**L02AB PROGESTOGENS**

L02AB01	MEGESTROL							
Tab	Orl	40 mg			Megestrol	02195917	AAP	ACDEFGVW
Tab	Orl	160 mg			Megestrol	02195925	AAP	ACDEFGVW

**L02AE GONADOTROPHIN RELEASING HORMONE ANALOGUES**

L02AE01	BUSERELIN							
Imp	Inj	6.3 mg			Suprefact Depot	02228955	XPI	ACDEFV
Imp	Inj	9.45 mg			Suprefact Depot	02240749	XPI	ACDEFV

L02AE02	LEUPRORELIN (LEUPROLIDE)							
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L02AE02 LEUPRORELIN (LEUPROLIDE)

Pws	Inj	3.75 mg	Lupron Depot	00884502	ABV	ACDEFGV
			Zeulide Depot	02429977	VRT	ACDEFV
Pws	Inj	7.5 mg	Lupron Depot	00836273	ABV	ACDEFGV
Pws	Inj	11.25 mg	Lupron Depot	02239834	ABV	ACDEFGV
Pws	Inj	22.5 mg	Lupron Depot	02230248	ABV	ACDEFGV
			Zeulide Depot	02462699	VRT	ACDEFV
Pws	Inj	30 mg	Lupron Depot	02239833	ABV	ACDEFGV
Sus	Inj	7.5 mg	Eligard	02248239	TOL	ACDEFV
Sus	Inj	22.5 mg	Eligard	02248240	TOL	ACDEFV
Sus	Inj	30 mg	Eligard	02248999	TOL	ACDEFV
Sus	Inj	45 mg	Eligard	02268892	TOL	ACDEFV

L02AE03 GOSERELIN

Imp	Inj	3.6 mg	Zoladex	02049325	AZE	ACDEFV
Imp	Inj	10.8 mg	Zoladex LA	02225905	AZE	ACDEFV

L02AE04 TRIPTORELIN

Pws	Inj	3.75 mg	Trelstar	02240000	KNI	ACDEFV
Pws	Inj	11.25 mg	Trelstar	02243856	KNI	ACDEFV
Pws	Inj	22.5 mg	Trelstar	02412322	KNI	ACDEFV

**L02B HORMONE ANTAGONISTS AND RELATED AGENTS**

**L02BA ANTI-ESTROGENS**

L02BA01 TAMOXIFEN

Tab	Orl	10 mg	Apo-Tamox	00812404	APX	ACDEFGV
			Teva-Tamoxifen	00851965	TEV	ACDEFGV
Tab	Orl	20 mg	Nolvadex-d (Disc/non disp Mar 22/24)	02048485	AZE	ACDEFGV
			Apo-Tamox	00812390	APX	ACDEFGV
			Teva-Tamoxifen	00851973	TEV	ACDEFGV

L02BA03 FULVESTRANT

Liq IM 50 mg/mL

Fulvestrant Injection 02483610 SDZ ACDEFGV

Teva-Fulvestrant 02460130 TEV ACDEFGV

**L02BB ANTI-ANDROGENS**

L02BB01 FLUTAMIDE

Tab Orl 250 mg

Flutamide 02238560 AAP ACDEFV

L02BB02 NILUTAMIDE

Tab Orl 50 mg

Anandron 02221861 XPI ACDEFV

L02BB03 BICALUTAMIDE

Tab Orl 50 mg

Casodex 02184478 AZE ACDEFV

Apo-Bicalutamide 02296063 APX ACDEFV

Bicalutamide 02325985 AHI ACDEFV

Bicalutamide 02519178 SAS ACDEFV

Jamp-Bicalutamide 02357216 JPC ACDEFV

pms-Bicalutamide 02275589 PMS ACDEFV

Teva-Bicalutamide 02270226 TEV ACDEFV

L02BB04 ENZALUTAMIDE

Cap Orl 40 mg

Xtandi 02407329 ASL (SA)

L02BB05 APALUTAMIDE

Tab Orl 60 mg

Erleada 02478374 JAN (SA)

L02BB06 DAROLUTAMIDE

Tab Orl 300 mg

Nubeqa 02496348 BAY (SA)

**L02BG AROMATASE INHIBITORS**

L02BG03 ANASTROZOLE

## L02BG03 ANASTROZOLE

Tab Orl 1 mg

Arimidex	02224135	AZE	ACDEFV
Act Anastrozole	02394898	TEV	ACDEFV
Anastrozole	02351218	AHI	ACDEFV
Anastrozole	02442736	SAS	ACDEFV
Anastrozole	02529904	SIV	ACDEFV
Apo-Anastrozole	02374420	APX	ACDEFV
CCP-Anastrozole	02458799	CCM	ACDEFV
Jamp-Anastrozole	02339080	JPC	ACDEFV
Mar-Anastrozole	02379562	MAR	ACDEFV
Mint-Anastrozole	02393573	MNT	ACDEFV
Nat-Anastrozole	02417855	NAT	ACDEFV
pms-Anastrozole	02320738	PMS	ACDEFV
Riva-Anastrozole	02392259	RIV	ACDEFV
Sandoz Anastrozole	02338467	SDZ	ACDEFV
Taro-Anastrozole	02365650	TAR	ACDEFV

## L02BG04 LETROZOLE

Tab Orl 2.5 mg

Femara	02231384	NVR	ACDEFV
Apo-Letrozole	02358514	APX	ACDEFV
CCP-Letrozole	02459884	CCM	ACDEFV
Jamp-Letrozole	02373009	JPC	ACDEFV
Letrozole	02504472	SAS	ACDEFV
Letrozole	02524244	SIV	ACDEFV
Letrozole tablets usp	02338459	AHI	ACDEFV
Mar-Letrozole	02373424	MAR	ACDEFV
Med-Letrozole	02322315	GMP	ACDEFV
Mint-Letrozole	02508109	MNT	ACDEFV
Nat-Letrozole	02421585	NAT	ACDEFV
pms-Letrozole	02309114	PMS	ACDEFV
Riva-Letrozole	02398656	RIV	ACDEFV
Sandoz Letrozole	02344815	SDZ	ACDEFV
Teva-Letrozole	02343657	TEV	ACDEFV
Zinda-Letrozole	02378213	MCK	ACDEFV

## L02BG06 EXEMESTANE

Tab Orl 25 mg

Aromasin	02242705	PFI	ACDEFV
Act Exemestane	02390183	TEV	ACDEFV
Med-Exemestane	02407841	GMP	ACDEFV
Teva-Exemestane	02408473	TEV	ACDEFV

**L02BX OTHER HORMONE ANTAGONISTS AND RELATED AGENTS****L02BX02 DEGARELIX**

Pws	Inj	80 mg/vial	Firmagon	02337029	FEI	ACDEFV
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Pws	Inj	120 mg/vial	Firmagon	02337037	FEI	ACDEFV
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**L02BX03 ABIRATERONE**

Tab	Orl	250 mg	Zytiga	02371065	JAN	(SA)
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Apo-Abiraterone	02491397	APX	(SA)
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Jamp Abiraterone	02502305	JPC	(SA)
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Mar-Abiraterone	02503980	MAR	(SA)
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Nat-Abiraterone	02494132	NAT	(SA)
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pms-Abiraterone	02492601	PMS	(SA)
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Reddy-Abiraterone	02477114	RCH	(SA)
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Sandoz Abiraterone	02486393	SDZ	(SA)
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Tab	Orl	500 mg	Zytiga	02457113	JAN	(SA)
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Abiraterone	02525380	JPC	(SA)
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Apo-Abiraterone	02491400	APX	(SA)
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Jamp Abiraterone	02529629	JPC	(SA)
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Mar-Abiraterone	02503999	MAR	(SA)
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pms-Abiraterone	02501503	PMS	(SA)
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Reddy-Abiraterone	02533251	RCH	(SA)
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Sandoz Abiraterone	02521644	SDZ	(SA)
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**L03 IMMUNOSTIMULANTS****L03A IMMUNOSTIMULANTS****L03AA COLONY STIMULATING FACTORS****L03AA02 FILGRASTIM**

Liq	SC	300 mcg / 0.5 mL	Grastofil	02441489	APO	(SA)
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Nivestym (prefilled syringe)	02485575	PFI	(SA)
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Liq	SC	300 mcg/mL	Nivestym	02485591	PFI	(SA)
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Liq	SC	480 mcg / 0.8 mL	Grastofil	02454548	APO	(SA)
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Nivestym (prefilled syringe)	02485583	PFI	(SA)
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Liq	SC	480 mcg / 1.6 mL	Nivestym	02485656	PFI	(SA)
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**L03AA13 PEGFILGRASTIM**

L03AA13 PEGFILGRASTIM

Liq SC 6 mg / 0.6 mL

Fulphila 02484153 BGP (SA)

Lapelga 02474565 APO (SA)

Nyvepria 02506238 PFI (SA)

Ziextenzo 02497395 SDZ (SA)

**L03AB INTERFERONS**

L03AB07 INTERFERON BETA-1A

Liq Inj 22 mcg / 0.5 mL

Rebif 02237319 EMD (SA)

Liq Inj 30 mcg / 0.5 mL

Avonex PS 02269201 BIG (SA)

Liq Inj 44 mcg / 0.5 mL

Rebif 02237320 EMD (SA)

Liq Inj 66 mcg / 1.5 mL

Rebif Cartridge 02318253 EMD (SA)

Liq Inj 132 mcg / 1.5 mL

Rebif Cartridge 02318261 EMD (SA)

L03AB08 INTERFERON BETA-1B

Pws SC 0.3 mg

Betaseron 02169649 BAY (SA)

L03AB11 PEGINTERFERON ALFA-2A

PEGINTERFERON ALFA-2A

Liq SC 180 mcg / 0.5 mL

Pegasys 02248077 ARN ACDEFGV

L03AB13 PEGINTERFERON BETA-1A

Kit SC 63 mcg / 0.5 mL,  
94 mcg / 0.5 mL

Plegridy (starter pack) 02444402 BIG (SA)

Liq SC 125 mcg / 0.5 mL

Plegridy 02444399 BIG (SA)

**L03AX OTHER IMMUNOSTIMULANTS**

L03AX13 GLATIRAMER ACETATE

Liq Inj 20 mg/mL

Glatect 02460661 PMS ACDEFGV

L03AX16 PLERIXAFOR

Liq Inj 24 mg / 1.2 mL

Mozobil 02377225 SAV (SA)

**L04 IMMUNOSUPPRESSANTS**

**L04A IMMUNOSUPPRESSANTS**

**L04AA SELECTIVE IMMUNOSUPPRESSANTS**

L04AA06 MYCOPHENOLIC ACID

## L04AA06 MYCOPHENOLIC ACID

Cap Orl 250 mg

Cellcept	02192748	HLR	ACDEFGRV
Apo-Mycophenolate	02352559	APX	ACDEFGRV
Jamp-Mycophenolate	02386399	JPC	ACDEFGRV
Mycophenolate Mofetil	02383780	AHI	ACDEFGRV
Mycophenolate Mofetil	02457369	SAS	ACDEFGRV
Sandoz Mycophenolate	02320630	SDZ	ACDEFGRV
Teva-Mycophenolate	02364883	TEV	ACDEFGRV

ECT Orl 180 mg

Myfortic	02264560	NVR	ACDEFGRV
Apo-Mycophenolic Acid	02372738	APX	ACDEFGRV
Mar-Mycophenolic Acid	02511673	MAR	ACDEFGRV

ECT Orl 360 mg

Myfortic	02264579	NVR	ACDEFGRV
Apo-Mycophenolic Acid	02372746	APX	ACDEFGRV
Mar-Mycophenolic Acid	02511681	MAR	ACDEFGRV

Pws Orl 200 mg/mL

Cellcept	02242145	HLR	ACDEFGRV
Mar-Mycophenolate Mofetil	02522233	MAR	ACDEFGRV

Tab Orl 500 mg

Cellcept	02237484	HLR	ACDEFGRV
Apo-Mycophenolate	02352567	APX	ACDEFGRV
Jamp-Mycophenolate	02380382	JPC	ACDEFGRV
Mycophenolate Mofetil	02378574	AHI	ACDEFGRV
Mycophenolate Mofetil	02457377	SAS	ACDEFGRV
Sandoz Mycophenolate	02313855	SDZ	ACDEFGRV
Teva-Mycophenolate	02348675	TEV	ACDEFGRV

## L04AA10 SIROLIMUS

Liq Orl 1 mg/mL

Rapamune	02243237	PFI	ACDEFGRV
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Tab Orl 1 mg

Rapamune	02247111	PFI	ACDEFGRV
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## L04AA13 LEFLUNOMIDE

Tab Orl 10 mg

Arava	02241888	SAV	ACDEFGRV
Apo-Leflunomide	02256495	APX	ACDEFGRV
Leflunomide	02351668	SAS	ACDEFGRV
Novo-Leflunomide	02261251	TEV	ACDEFGRV
pms-Leflunomide	02288265	PMS	ACDEFGRV
Sandoz Leflunomide	02283964	SDZ	ACDEFGRV



L04AA13	LEFLUNOMIDE							
Tab	Orl	20 mg			Arava	02241889	SAV	ACDEFGV
					Apo-Leflunomide	02256509	APX	ACDEFGV
					Leflunomide	02351676	SAS	ACDEFGV
					Novo-Leflunomide	02261278	TEV	ACDEFGV
					pms-Leflunomide	02288273	PMS	ACDEFGV
					Sandoz Leflunomide	02283972	SDZ	ACDEFGV
L04AA23	NATALIZUMAB							
Liq	IV	300 mg / 15 mL			Tysabri	02286386	BIG	(SA)
L04AA24	ABATACEPT							
Liq	SC	125 mg/mL			Orencia	02402475	BRI	(SA)
Pws	IV	250 mg / 15 mL			Orencia	02282097	BRI	(SA)
L04AA25	ECULIZUMAB							
Liq	IV	300 mg / 30 mL			Soliris	02322285	ALX	(SA)
L04AA27	FINGOLIMOD							
Cap	Orl	0.5 mg			Gilenya	02365480	NVR	(SA)
					Apo-Fingolimod	02469936	APX	(SA)
					Jamp-Fingolimod	02487772	JPC	(SA)
					Mar-Fingolimod	02474743	MAR	(SA)
					Mylan-Fingolimod	02469715	MYL	(SA)
					pms-Fingolimod	02469782	PMS	(SA)
					Sandoz Fingolimod	02482606	SDZ	(SA)
					Taro-Fingolimod	02469618	TAR	(SA)
					Teva-Fingolimod	02469561	TEV	(SA)
L04AA29	TOFACITINIB							
ERT	Orl	11 mg			Xeljanz XR	02470608	PFI	(SA)
Tab	Orl	5 mg			Xeljanz	02423898	PFI	(SA)
					Auro-Tofacitinib	02530007	ARO	(SA)
					pms-Tofacitinib	02522799	PMS	(SA)
					Taro-Tofacitinib	02511304	TAR	(SA)
Tab	Orl	10 mg			Xeljanz	02480786	PFI	(SA)
					Auro-Tofacitinib	02530015	ARO	(SA)
					Taro-Tofacitinib	02511312	TAR	(SA)

## L04AA31 TERIFLUNOMIDE

Tab Orl 14 mg

Aubagio	02416328	GZM	(SA)
ACH-Teriflunomide	02502933	AHI	(SA)
Apo-Teriflunomide	02500639	APX	(SA)
Jamp Teriflunomide	02504170	JPC	(SA)
M-Teriflunomide	02523833	MRA	(SA)
Mar-Teriflunomide	02500469	MAR	(SA)
Nat-Teriflunomide	02500310	NAT	(SA)
pms-Teriflunomide	02500434	PMS	(SA)
Sandoz Teriflunomide	02505843	SDZ	(SA)
Teva-Teriflunomide	02501090	TEV	(SA)

## L04AA33 VEDOLIZUMAB

Liq SC 108 mg / 0.68 mL

Entyvio (autoinjector)	02497867	TAK	(SA)
Entyvio (prefilled syringe)	02497875	TAK	(SA)

Pws IV 300 mg

Entyvio 02436841 TAK (SA)

## L04AA34 ALEMTUZUMAB

Liq IV 12 mg / 1.2 mL

Lemtrada 02418320 GZM (SA)

## L04AA36 OCRELIZUMAB

Liq IV 30 mg/mL

Ocrevus 02467224 HLR (SA)

## L04AA37 BARICITINIB

Tab Orl 2 mg

Olumiant 02480018 LIL (SA)

## L04AA40 CLADRIBINE

Tab Orl 10 mg

Mavenclad 02470179 EMD (SA)

## L04AA42 SIPONIMOD

Tab Orl 0.25 mg

Mayzent 02496429 NVR (SA)

Tab Orl 2 mg

Mayzent 02496437 NVR (SA)

## L04AA44 UPADACITINIB

ERT Orl 15 mg

Rinvoq 02495155 ABV (SA)

## L04AA52 OFATUMUMAB

Liq SC 20 mg / 0.4 mL

Kesimpta 02511355 NVR (SA)

**L04AB TUMOR NECROSIS FACTOR ALPHA (TNF-A) INHIBITORS**

L04AB01	ETANERCEPT							
	Liq	SC	25 mg / 0.5 mL		Erelzi (syringe)	02462877	SDZ	(SA)
	Liq	SC	50 mg/mL		Brenzys (autoinjector)	02455331	ORG	(SA)
					Brenzys (syringe)	02455323	ORG	(SA)
					Erelzi (autoinjector)	02462850	SDZ	(SA)
					Erelzi (syringe)	02462869	SDZ	(SA)
L04AB02	INFLIXIMAB							
	Pws	IV	100 mg		Avsola	02496933	AGA	(SA)
					Inflectra	02419475	HOS	(SA)
					Renflexis	02470373	ORG	(SA)
L04AB04	ADALIMUMAB							
	Liq	SC	20 mg / 0.2 mL		Amgevita (prefilled syringe)	02459310	AGA	(SA)
					Hyrimoz (prefilled syringe)	02505258	SDZ	(SA)
	Liq	SC	20 mg / 0.4 mL		Abrilada (prefilled syringe)	02511061	PFI	(SA)
					Hulio (prefilled syringe)	02502380	BGP	(SA)
	Liq	SC	40 mg / 0.4 mL		Simlandi (autoinjector)	02523957	JPC	(SA)
					Simlandi (prefilled syringe)	02523949	JPC	(SA)
					Yuflyma (autoinjector)	02523779	CTL	(SA)
	Liq	SC	40 mg / 0.8 mL		Abrilada (autoinjector)	02511045	PFI	(SA)
					Abrilada (prefilled syringe)	02511053	PFI	(SA)
					Amgevita (autoinjector)	02459302	AGA	(SA)
					Amgevita (prefilled syringe)	02459299	AGA	(SA)
					Hadlima (autoinjector)	02473100	ORG	(SA)
					Hadlima (prefilled syringe)	02473097	ORG	(SA)
					Hulio (autoinjector)	02502402	BGP	(SA)
					Hulio (prefilled syringe)	02502399	BGP	(SA)
					Hyrimoz (autoinjector)	02492156	SDZ	(SA)
					Hyrimoz (prefilled syringe)	02492164	SDZ	(SA)
					Idacio (autoinjector)	02502674	FKB	(SA)
	Liq	SC	80 mg / 0.8 mL		Simlandi (prefilled syringe)	02523965	JPC	(SA)
L04AB05	CERTOLIZUMAB PEGOL							
	Liq	SC	200 mg/mL		Cimzia	02331675	UCB	(SA)
					Cimzia (autoinjector)	02465574	UCB	(SA)

L04AB06	GOLIMUMAB						
Liq	SC	50 mg / 0.5 mL			Simponi (autoinjector)	02324784	JAN (SA)
					Simponi (prefilled syringe)	02324776	JAN (SA)
Liq	SC	100 mg/mL			Simponi (autoinjector)	02413183	JAN (SA)
					Simponi (prefilled syringe)	02413175	JAN (SA)

**L04AC INTERLEUKIN INHIBITORS**

L04AC05	USTEKINUMAB						
Liq	SC	45 mg / 0.5 mL			Stelara	02320673	JAN (SA)
Liq	SC	90 mg/mL			Stelara	02320681	JAN (SA)
L04AC07	TOCILIZUMAB						
Liq	IV	80 mg / 4 mL			Actemra	02350092	HLR (SA)
Liq	IV	200 mg / 10 mL			Actemra	02350106	HLR (SA)
Liq	IV	400 mg / 20 mL			Actemra	02350114	HLR (SA)
Liq	SC	162 mg / 0.9 mL			Actemra (autoinjector)	02483327	HLR (SA)
					Actemra (prefilled syringe)	02424770	HLR (SA)
L04AC08	CANAKINUMAB						
Liq	SC	150 mg/mL			Ilaris	02460351	NVR (SA)
L04AC10	SECUKINUMAB						
Liq	SC	150 mg/mL			Cosentyx	02438070	NVR (SA)
L04AC12	BRODALUMAB						
Liq	SC	210 mg			Siliq	02473623	BSL (SA)
L04AC13	IXEKIZUMAB						
Liq	SC	80 mg/mL			Taltz (autoinjector)	02455102	LIL (SA)
					Taltz (prefilled syringe)	02455110	LIL (SA)
L04AC14	SARILUMAB						
Liq	SC	150 mg / 1.14 mL			Kevzara (autoinjector)	02472961	SAV (SA)
Liq	SC	200 mg / 1.14 mL			Kevzara (autoinjector)	02472988	SAV (SA)
					Kevzara (prefilled syringe)	02460548	SAV (SA)
L04AC17	TILDRAKIZUMAB						

L04AC17 TILDRAKIZUMAB  
Liq SC 100 mg/mL Ilumya 02516098 SUN (SA)

L04AC18 RISANKIZUMAB  
Liq SC 75 mg / 0.83 mL Skyrizi 02487454 ABV (SA)

Liq SC 150 mg/mL Skyrizi (autoinjector) 02519291 ABV (SA)  
Skyrizi (prefilled syringe) 02519283 ABV (SA)

L04AC19 SATRALIZUMAB  
Liq SC 120 mg/mL Enspryng 02499681 HLR (SA)

L04AC21 BIMEKIZUMAB  
Liq SC 160 mg/mL Bimzelx (autoinjector) 02525275 UCB (SA)  
Bimzelx (prefilled syringe) 02525267 UCB (SA)

**L04AD CALCINEURIN INHIBITORS**

L04AD01 CYCLOSPORINE  
Cap Orl 10 mg Neoral 02237671 NVR ACDEFGRV

Cap Orl 25 mg Neoral 02150689 NVR ACDEFGRV  
Cyclosporine Capsules 02495805 STD ACDEFGRV  
Sandoz Cyclosporine 02247073 SDZ ACDEFGRV

Cap Orl 50 mg Neoral 02150662 NVR ACDEFGRV  
Cyclosporine Capsules 02495821 STD ACDEFGRV  
Sandoz Cyclosporine 02247074 SDZ ACDEFGRV

Cap Orl 100 mg Neoral 02150670 NVR ACDEFGRV  
Cyclosporine Capsules 02495813 STD ACDEFGRV  
Sandoz Cyclosporine 02242821 SDZ ACDEFGRV

Liq Orl 100 mg/mL Neoral 02150697 NVR ACDEFGRV

L04AD02 TACROLIMUS  
Cap Orl 0.5 mg Prograf 02243144 ASL ACDEFGRV  
Sandoz Tacrolimus 02416816 SDZ ACDEFGRV

Cap Orl 1 mg Prograf 02175991 ASL ACDEFGRV  
Sandoz Tacrolimus 02416824 SDZ ACDEFGRV

L04AD02 TACROLIMUS

Cap	Orl	5 mg	Prograf	02175983	ASL	ACDEFGRV
			Sandoz Tacrolimus	02416832	SDZ	ACDEFGRV
ERC	Orl	0.5 mg	Advagraf	02296462	ASL	ACDEFGRV
ERC	Orl	1 mg	Advagraf	02296470	ASL	ACDEFGRV
ERC	Orl	3 mg	Advagraf	02331667	ASL	ACDEFGRV
ERC	Orl	5 mg	Advagraf	02296489	ASL	ACDEFGRV
ERT	Orl	0.75 mg	Envarsus PA	02485877	EDO	ACDEFGRV
ERT	Orl	1 mg	Envarsus PA	02485885	EDO	ACDEFGRV
ERT	Orl	4 mg	Envarsus PA	02485893	EDO	ACDEFGRV

**L04AX OTHER IMMUNOSUPPRESSANTS**

L04AX01 AZATHIOPRINE

Tab	Orl	50 mg	Imuran	00004596	APN	ACDEFGRV
			Apo-Azathioprine	02242907	APX	ACDEFGRV
			Teva-Azathioprine	02236819	TEV	ACDEFGRV

L04AX04 LENALIDOMIDE

Cap	Orl	2.5 mg	Revlimid	02459418	CEL	(SA)
			Apo-Lenalidomide	02507927	APX	(SA)
			Jamp Lenalidomide	02506130	JPC	(SA)
			Nat-Lenalidomide	02493837	NAT	(SA)
			Reddy-Lenalidomide	02484714	RCH	(SA)
			Sandoz Lenalidomide	02518562	SDZ	(SA)
			Taro-Lenalidomide	02507862	TAR	(SA)
Cap	Orl	5 mg	Revlimid	02304899	CEL	(SA)
			Apo-Lenalidomide	02507935	APX	(SA)
			Jamp Lenalidomide	02506149	JPC	(SA)
			Nat-Lenalidomide	02493845	NAT	(SA)
			Reddy-Lenalidomide	02483017	RCH	(SA)
			Sandoz Lenalidomide	02518570	SDZ	(SA)
			Taro-Lenalidomide	02507870	TAR	(SA)

## L04AX04 LENALIDOMIDE

Cap Orl 10 mg

Revlimid	02304902	CEL	(SA)
Apo-Lenalidomide	02507943	APX	(SA)
Jamp Lenalidomide	02506157	JPC	(SA)
Nat-Lenalidomide	02493861	NAT	(SA)
Reddy-Lenalidomide	02483025	RCH	(SA)
Sandoz Lenalidomide	02518589	SDZ	(SA)
Taro-Lenalidomide	02507889	TAR	(SA)

Cap Orl 15 mg

Revlimid	02317699	CEL	(SA)
Apo-Lenalidomide	02507951	APX	(SA)
Jamp Lenalidomide	02506165	JPC	(SA)
Nat-Lenalidomide	02493888	NAT	(SA)
Reddy-Lenalidomide	02483033	RCH	(SA)
Sandoz Lenalidomide	02518597	SDZ	(SA)
Taro-Lenalidomide	02507897	TAR	(SA)

Cap Orl 20 mg

Revlimid	02440601	CEL	(SA)
Apo-Lenalidomide	02507978	APX	(SA)
Jamp Lenalidomide	02506173	JPC	(SA)
Nat-Lenalidomide	02493896	NAT	(SA)
Reddy-Lenalidomide	02483041	RCH	(SA)
Sandoz Lenalidomide	02518600	SDZ	(SA)
Taro-Lenalidomide	02507900	TAR	(SA)

Cap Orl 25 mg

Revlimid	02317710	CEL	(SA)
Apo-Lenalidomide	02507986	APX	(SA)
Jamp Lenalidomide	02506181	JPC	(SA)
Nat-Lenalidomide	02493918	NAT	(SA)
Reddy-Lenalidomide	02483068	RCH	(SA)
Sandoz Lenalidomide	02518619	SDZ	(SA)
Taro-Lenalidomide	02507919	TAR	(SA)

## L04AX05 PIRFENIDONE

Cap Orl 267 mg

Esbriet	02393751	HLR	(SA)
Jamp Pirfenidone	02509938	JPC	(SA)
Sandoz Pirfenidone	02488833	SDZ	(SA)

## L04AX05 PIRFENIDONE

Tab Orl 267 mg

Esbriet	02464489	HLR	(SA)
Jamp Pirfenidone	02514702	JPC	(SA)
pms-Pirfenidone	02531526	PMS	(SA)
Sandoz Pirfenidone	02488507	SDZ	(SA)

Tab Orl 801 mg

Esbriet	02464500	HLR	(SA)
Jamp Pirfenidone	02514710	JPC	(SA)
pms-Pirfenidone	02531534	PMS	(SA)
Sandoz Pirfenidone	02488515	SDZ	(SA)

## L04AX06 POMALIDOMIDE

Cap Orl 1 mg

Pomalyst	02419580	CEL	(SA)
Apo-Pomalidomide	02520427	APX	(SA)
Jamp Pomalidomide	02538059	JPC	(SA)
Nat-Pomalidomide	02506394	NAT	(SA)
Reddy-Pomalidomide	02504073	RCH	(SA)
Sandoz Pomalidomide	02523973	SDZ	(SA)

Cap Orl 2 mg

Pomalyst	02419599	CEL	(SA)
Apo-Pomalidomide	02520435	APX	(SA)
Jamp Pomalidomide	02538075	JPC	(SA)
Nat-Pomalidomide	02506408	NAT	(SA)
Reddy-Pomalidomide	02504081	RCH	(SA)
Sandoz Pomalidomide	02523981	SDZ	(SA)

Cap Orl 3 mg

Pomalyst	02419602	CEL	(SA)
Apo-Pomalidomide	02520443	APX	(SA)
Jamp Pomalidomide	02538083	JPC	(SA)
Nat-Pomalidomide	02506416	NAT	(SA)
Reddy-Pomalidomide	02504103	RCH	(SA)
Sandoz Pomalidomide	02524007	SDZ	(SA)

Cap Orl 4 mg

Pomalyst	02419610	CEL	(SA)
Apo-Pomalidomide	02520451	APX	(SA)
Jamp Pomalidomide	02538091	JPC	(SA)
Nat-Pomalidomide	02506424	NAT	(SA)
Reddy-Pomalidomide	02504111	RCH	(SA)
Sandoz Pomalidomide	02524015	SDZ	(SA)

## N07XX OTHER NERVOUS SYSTEM DRUGS



L04AX07 DIMETHYL FUMARATE

CDR Orl 120 mg

Tecfidera 02404508 BIG (SA)  
 ACH-Dimethyl Fumarate 02495341 AHI (SA)  
 Apo-Dimethyl Fumarate 02505762 APX (SA)  
 GLN-Dimethyl Fumarate 02494809 GLM (SA)  
 Jamp-Dimethyl Fumarate 02516047 JPC (SA)  
 Mar-Dimethyl Fumarate 02502690 MAR (SA)  
 pms-Dimethyl Fumarate 02497026 PMS (SA)  
 Sandoz Dimethyl Fumarate 02513781 SDZ (SA)

CDR Orl 240 mg

Tecfidera 02420201 BIG (SA)  
 ACH-Dimethyl Fumarate 02495368 AHI (SA)  
 Apo-Dimethyl Fumarate 02505770 APX (SA)  
 GLN-Dimethyl Fumarate 02494817 GLM (SA)  
 Jamp-Dimethyl Fumarate 02516055 JPC (SA)  
 Mar-Dimethyl Fumarate 02502704 MAR (SA)  
 pms-Dimethyl Fumarate 02497034 PMS (SA)  
 Sandoz Dimethyl Fumarate 02513803 SDZ (SA)

**M MUSCULO-SKELETAL SYSTEM**

**M01 ANTIINFLAMMATORY AND ANTIRHEUMATIC PRODUCTS**

**M01A ANTIINFLAMMATORY AND ANTIRHEUMATIC PRODUCTS, NON-STEROIDS**

**M01AB ACETIC ACID DERIVATIVES AND RELATED SUBSTANCES**

M01AB01 INDOMETHACIN

Cap Orl 25 mg

Mint-Indomethacin 02461811 MNT ACDEFGV  
 Teva-Indomethacin 00337420 TEV ACDEFGV

Cap Orl 50 mg

Auro-Indomethacin 02499223 ARO ACDEFGV  
 Mint-Indomethacin 02461536 MNT ACDEFGV  
 Teva-Indomethacin 00337439 TEV ACDEFGV

Sup Rt 50 mg

Sandoz Indomethacin 02231799 SDZ ACDEFGV

Sup Rt 100 mg

Sandoz Indomethacin 02231800 SDZ ACDEFGV

M01AB02 SULINDAC

Tab Orl 150 mg

Teva-Sundac 00745588 TEV ACDEFGV

Tab Orl 200 mg

Teva-Sundac 00745596 TEV ACDEFGV

M01AB05 DICLOFENAC

M01AB05 DICLOFENAC

ECT Orl 25 mg

Apo-Diclo 00839175 APX ACDEFGV  
 pms-Diclofenac 02302616 PMS ACDEFGV  
 Teva-Difenac 00808539 TEV ACDEFGV

ECT Orl 50 mg

Apo-Diclo 00839183 APX ACDEFGV  
 pms-Diclofenac 02302624 PMS ACDEFGV  
 Teva-Difenac 00808547 TEV ACDEFGV

SRT Orl 75 mg

Apo-Diclo SR 02162814 APX ACDEFGV  
 Teva-Difenac SR 02158582 TEV ACDEFGV

SRT Orl 100 mg

Voltaren SR 00590827 NVR ACDEFGV  
 Apo-Diclo SR 02091194 APX ACDEFGV  
 Sandoz Diclofenac SR 02261944 SDZ ACDEFGV  
 Teva-Difenac SR 02048698 TEV ACDEFGV

Sup Rt 50 mg

Voltaren 00632724 NVR ACDEFGV  
 Sandoz Diclofenac 02261928 SDZ ACDEFGV

M01AB15 KETOROLAC

Liq Inj 10 mg

Toradol 02162644 MTP W

M01AB55 DICLOFENAC COMBINATIONS  
 DICLOFENAC / MISOPROSTOL

Tab Orl 50 mg / 200 mcg

Arthrotec 01917056 PFI ACDEFGV  
 GD-Diclofenac/Misoprostol 02341689 GMD ACDEFGV  
 pms-Diclofenac/Misoprostol 02413469 PMS ACDEFGV

Tab Orl 75 mg / 200 mcg

Arthrotec 02229837 PFI ACDEFGV  
 GD-Diclofenac/Misoprostol 02341697 GMD ACDEFGV  
 pms-Diclofenac/Misoprostol 02413477 PMS ACDEFGV

**M01AC OXICAMS**

M01AC01 PIROXICAM

Cap Orl 10 mg

Novo-Pirocam 00695718 TEV ACDEFGV

Cap Orl 20 mg

Novo-Pirocam 00695696 TEV ACDEFGV

M01AC06 MELOXICAM

M01AC06 MELOXICAM

Tab Orl 7.5 mg

Apo-Meloxicam 02248973 APX ACDEFGV  
 Auro-Meloxicam 02390884 ARO ACDEFGV  
 Meloxicam 02353148 SAS ACDEFGV  
 pms-Meloxicam 02248267 PMS ACDEFGV  
 Teva-Meloxicam 02258315 TEV ACDEFGV

Tab Orl 15 mg

Apo-Meloxicam 02248974 APX ACDEFGV  
 Auro-Meloxicam 02390892 ARO ACDEFGV  
 Meloxicam 02353156 SAS ACDEFGV  
 pms-Meloxicam 02248268 PMS ACDEFGV  
 Teva-Meloxicam 02258323 TEV ACDEFGV

**M01AE PROPIONIC ACID DERIVATIVES**

M01AE01 IBUPROFEN

Tab Orl 300 mg

Apo-Ibuprofen 00441651 APX AEFVG

Tab Orl 400 mg

Motrin IB 02242658 JNJ AEFVG

Tab Orl 600 mg

Apo-Ibuprofen 00585114 APX ACDEFGV  
 Novo-Profen 00629359 TEV ACDEFGV

M01AE02 NAPROXEN

ECT Orl 250 mg

Naproxen EC (Disc/non disp Jul 4/24) 02350785 SAS ACDEFGV  
 Teva-Naprox EC 02243312 TEV ACDEFGV

ECT Orl 375 mg

Naprosyn E 02162415 MTP ACDEFGV  
 Apo-Naproxen EC 02246700 APX ACDEFGV  
 Naproxen EC (Disc/non disp Jul 4/24) 02350793 SAS ACDEFGV  
 Teva-Naprox EC 02243313 TEV ACDEFGV

ECT Orl 500 mg

Naprosyn E 02162423 MTP ACDEFGV  
 Apo-Naproxen EC 02246701 APX ACDEFGV  
 Naproxen EC (Disc/non disp Jul 4/24) 02350807 SAS ACDEFGV  
 pms-Naproxen EC 02294710 PMS ACDEFGV  
 Teva-Naprox EC 02243314 TEV ACDEFGV

SRT Orl 750 mg

Naprosyn SR 02162466 MTP ACDEFGV

Sus Orl 25 mg/mL

Pediapharm Naproxen 02162431 MDX ACDEFGV

M01AE02	NAPROXEN						
Tab	Orl	250 mg	Apo-Naproxen	00522651	APX	ACDEFGV	
			Naproxen (Disc/non disp Jul 4/24)	02350750	SAS	ACDEFGV	
			Teva-Naproxen	00565350	TEV	ACDEFGV	
Tab	Orl	275 mg	Anaprox	02162725	MTP	ACDEFGV	
			Apo-Napro-Na	00784354	APX	ACDEFGV	
			Naproxen Sodium	02351013	SAS	ACDEFGV	
			Teva-Naproxen Sodium	00778389	TEV	ACDEFGV	
Tab	Orl	375 mg	Apo-Naproxen	00600806	APX	ACDEFGV	
			Naproxen (Disc/non disp Jul 4/24)	02350769	SAS	ACDEFGV	
			Teva-Naproxen	00627097	TEV	ACDEFGV	
Tab	Orl	500 mg	Apo-Naproxen	00592277	APX	ACDEFGV	
			Naproxen (Disc/non disp Jul 4/24)	02350777	SAS	ACDEFGV	
			Teva-Naproxen	00589861	TEV	ACDEFGV	
Tab	Orl	550 mg	Anaprox DS	02162717	MTP	ACDEFGV	
			Apo-Napro-Na DS	01940309	APX	ACDEFGV	
			Naproxen Sodium DS (Disc/non disp Jul 4/24)	02351021	SAS	ACDEFGV	
			Teva-Naproxen Sodium DS	02026600	TEV	ACDEFGV	
M01AE03	KETOPROFEN						
Cap	Orl	50 mg	Keto	00790427	AAP	ACDEFGV	
ECT	Orl	50 mg	Keto-E	00790435	AAP	ACDEFGV	
ECT	Orl	100 mg	Keto-E	00842664	AAP	ACDEFGV	
SRT	Orl	200 mg	Keto SR	02172577	AAP	ACDEFGV	
M01AE09	FLURBIPROFEN						
Tab	Orl	50 mg	Flurbiprofen	01912046	AAP	ACDEFGV	
Tab	Orl	100 mg	Flurbiprofen	01912038	AAP	ACDEFGV	
M01AE11	TIAPROFENIC ACID						
Tab	Orl	200 mg	Teva-Tiaprofenic	02179679	TEV	ACDEFGV	
Tab	Orl	300 mg	Teva-Tiaprofenic	02179687	TEV	ACDEFGV	

**M01AG FENEMATES**

M01AG01 MEFENAMIC ACID

Cap Orl 250 mg

Ponstan 00155225 AAP ACDEFGV

Mefenamic 02229452 AAP ACDEFGV

**M01AH COXIBS**

M01AH01 CELECOXIB

Cap Orl 100 mg

Celebrex 02239941 UJC ACDEFGV

Apo-Celecoxib 02418932 APX ACDEFGV

Auro-Celecoxib 02445670 ARO ACDEFGV

Celecoxib 02436299 SAS ACDEFGV

Celecoxib 02429675 SIV ACDEFGV

Jamp-Celecoxib 02424533 JPC ACDEFGV

M-Celecoxib 02495465 MRA ACDEFGV

Mar-Celecoxib 02420058 MAR ACDEFGV

Mint-Celecoxib 02412497 MNT ACDEFGV

NRA-Celecoxib 02479737 NRA ACDEFGV

pms-Celecoxib 02355442 PMS ACDEFGV

pmsc-Celecoxib 02517116 PMS ACDEFGV

Cap Orl 200 mg

Celebrex 02239942 UJC ACDEFGV

Apo-Celecoxib 02418940 APX ACDEFGV

Auro-Celecoxib 02445689 ARO ACDEFGV

Celecoxib 02436302 SAS ACDEFGV

Celecoxib 02429683 SIV ACDEFGV

Jamp-Celecoxib 02424541 JPC ACDEFGV

M-Celecoxib 02495473 MRA ACDEFGV

Mar-Celecoxib 02420066 MAR ACDEFGV

Mint-Celecoxib 02412500 MNT ACDEFGV

NRA-Celecoxib 02479745 NRA ACDEFGV

pms-Celecoxib 02355450 PMS ACDEFGV

pmsc-Celecoxib 02517124 PMS ACDEFGV

**M01AX OTHER ANTIINFLAMMATORY AND ANTIRHEUMATIC AGENTS, NON STEROIDS**

M01AX01 NABUMETONE

Tab Orl 500 mg

Nabumetone 02238639 AAP ACDEFGV

**M01C SPECIFIC ANTIRHEUMATIC AGENTS****M01CC PENICILLAMINE AND SIMILAR AGENTS**

M01CC01 PENICILLAMINE

Cap Orl 250 mg

Cuprimine 00016055 BSL ACDEFGV

**M03 MUSCLE RELAXANTS****M03A PERIPHERALLY ACTING AGENTS, MUSCLE RELAXANTS****M03AX OTHER MUSCLE RELAXANTS, PERIPHERALLY ACTING**

M03AX01 BOTULINUM TOXIN  
 ABOBOTULINUMTOXINA

Pws IM 300 Unit Dysport Therapeutic 02460203 IPS (SA)

Pws IM 500 Unit Dysport Therapeutic 02456117 IPS (SA)

INCOBOTULINUMTOXINA

Pws IM 50 Unit Xeomin 02371081 MRZ (SA)

Pws IM 100 Unit Xeomin 02324032 MRZ (SA)

ONABOTULINUMTOXINA

Pws IM 50 Unit Botox 00903741 ABV (SA)

Pws IM 100 Unit Botox 01981501 ABV (SA)

Pws IM 200 Unit Botox 00999505 ABV (SA)

**M03B MUSCLE RELAXANTS, CENTRALLY ACTING AGENTS****M03BA CARBAMIC ACID ESTERS**

M03BA03 METHOCARBAMOL

Tab Orl 500 mg Robaxin 01930990 GCH AEEFGV

Tab Orl 750 mg Robaxin 01932187 GCH AEEFGV

**M03BC ETHERS, CHEMICALLY CLOSE TO ANTIHISTAMINES**

M03BC01 ORPHENADRINE

SRT Orl 100 mg Sandoz Orphenadrine Citrate 02243559 SDZ AEEFGV

**M03BX OTHER CENTRALLY ACTING AGENTS**

M03BX01 BACLOFEN

Liq Int 0.05 mg/mL Lioresal 02131048 NVR ACDEFGV

Baclofen 02457059 HIK ACDEFGV

Liq Int 0.5 mg/mL Lioresal 02131056 NVR ACDEFGV

Baclofen 02457067 HIK ACDEFGV

Liq Int 2 mg/mL Lioresal 02131064 NVR ACDEFGV

Baclofen 02457075 HIK ACDEFGV

**M03BX01 BACLOFEN**

Tab Orl 10 mg

Apo-Baclofen	02139332	APX	ACDEFGV
Baclofen	02287021	SAS	ACDEFGV
Mylan-Baclofen	02088398	MYL	ACDEFGV
pms-Baclofen	02063735	PMS	ACDEFGV

Tab Orl 20 mg

Apo-Baclofen	02139391	APX	ACDEFGV
Baclofen	02287048	SAS	ACDEFGV
Mylan-Baclofen	02088401	MYL	ACDEFGV
pms-Baclofen	02063743	PMS	ACDEFGV

**M03BX02 TIZANIDINE**

Tab Orl 4 mg

Tizanidine	02259893	AAP	ACDEFGV
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**M03BX08 CYCLOBENZAPRINE**

Tab Orl 10 mg

Apo-Cycloprine	02177145	APX	ACDEFGV
Auro-Cyclobenzaprine	02348853	ARO	ACDEFGV
Cyclobenzaprine	02287064	SAS	ACDEFGV
Cyclobenzaprine	02424584	SIV	ACDEFGV
Flexeril	02495422	ORI	ACDEFGV
Jamp-Cyclobenzaprine	02357127	JPC	ACDEFGV
Novo-Cycloprine	02080052	TEV	ACDEFGV
pms-Cyclobenzaprine	02212048	PMS	ACDEFGV

**M03C MUSCLE RELAXANTS, DIRECTLY ACTING AGENTS****M03CA DANTROLENE AND DERIVATIVES****M03CA01 DANTROLENE**

Cap Orl 25 mg

Dantrium	01997602	PAL	ACDEFGV
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**M04 ANTIGOUT PREPARATIONS****M04A ANTIGOUT PREPARATIONS****M04AA PREPARATIONS INHIBITING URIC ACID PRODUCTION****M04AA01 ALLOPURINOL**

Tab Orl 100 mg

Zyloprim	00402818	AAP	ACDEFGV
Apo-Allopurinol	02402769	APX	ACDEFGV
Mar-Allopurinol	02396327	MAR	ACDEFGV

Tab Orl 200 mg

Zyloprim	00479799	AAP	ACDEFGV
Apo-Allopurinol	02402777	APX	ACDEFGV
Mar-Allopurinol	02396335	MAR	ACDEFGV

M04AA01 ALLOPURINOL

Tab Orl 300 mg

Zyloprim 00402796 AAP ACDEFGV  
Apo-Allopurinol 02402785 APX ACDEFGV  
Mar-Allopurinol 02396343 MAR ACDEFGV

M04AA03 FEBUXOSTAT

Tab Orl 80 mg

Auro-Febuxostat 02533243 ARO (SA)  
Jamp-Febuxostat 02490870 JPC (SA)  
Mar-Febuxostat 02473607 MAR (SA)  
Teva-Febuxostat 02466198 TEV (SA)

**M04AC PREPARATION WITH NO EFFECT ON URIC ACID METABOLISM**

M04AC01 COLCHICINE

Tab Orl 0.6 mg

Colchicine 00572349 ODN ACDEFGV  
Jamp-Colchicine 02373823 JPC ACDEFGV  
pms-Colchicine 02402181 PMS ACDEFGV  
Sandoz Colchicine 00287873 SDZ ACDEFGV

**M05 DRUGS FOR TREATMENT OF BONE DISEASES**

**M05B DRUGS AFFECTING BONE STRUCTURE AND MINERALIZATION**

**M05BA BISPHOSPHONATES**

M05BA02 CLODRONIC ACID

Cap Orl 400 mg

Clasteon 02245828 SNV ACDEFGV

M05BA04 ALENDRONIC ACID

Tab Orl 10 mg

Alendronate Sodium 02381486 AHI ACDEFGV  
Auro-Alendronate 02388545 ARO ACDEFGV



M05BA04 ALENDRONIC ACID

Tab Orl 70 mg

Fosamax 02245329 ORG ACDEFGV  
 Alendronate 02352966 SAS ACDEFGV  
 Alendronate 02299712 SIV ACDEFGV  
 Alendronate Sodium 02381494 AHI ACDEFGV  
 Alendronate-70 02303078 PDL ACDEFGV  
 Apo-Alendronate 02248730 APX ACDEFGV  
 Auro-Alendronate 02388553 ARO ACDEFGV  
 Jamp Alendronate Sodium 02500175 JPC ACDEFGV  
 Jamp-Alendronate 02385031 JPC ACDEFGV  
 M-Alendronate 02529394 MRA ACDEFGV  
 Mint-Alendronate 02394871 MNT ACDEFGV  
 pms-Alendronate FC 02284006 PMS ACDEFGV  
 Riva-Alendronate 02270889 RIV ACDEFGV  
 Sandoz Alendronate 02288109 SDZ ACDEFGV  
 Teva-Alendronate 02261715 TEV ACDEFGV

M05BA07 RISEDRONIC ACID

Tab Orl 5 mg

Teva-Risedronate 02298376 TEV ACDEFGV

Tab Orl 30 mg

Teva-Risedronate 02298384 TEV (SA)

Tab Orl 35 mg

Actonel 02246896 ABV ACDEFGV  
 Apo-Risedronate 02353687 APX ACDEFGV  
 Auro-Risedronate 02406306 ARO ACDEFGV  
 pms-Risedronate 02302209 PMS ACDEFGV  
 Risedronate 02347474 PDL ACDEFGV  
 Risedronate 02370255 SAS ACDEFGV  
 Risedronate 02411407 SIV ACDEFGV  
 Sandoz Risedronate 02327295 SDZ ACDEFGV  
 Teva-Risedronate 02298392 TEV ACDEFGV

M05BA08 ZOLEDRONIC ACID

Liq IV 5 mg / 100 mL

Aclasta 02269198 SDZ ACDEFGV  
 Taro-Zoledronic Acid 02415100 TAR ACDEFGV  
 Zoledronic Acid 02422433 RCH ACDEFGV

**M05BB BISPHOSPHONATES, COMBINATIONS**

M05BB03 ALENDRONIC ACID AND COLECALCIFEROL

**M05BB03 ALENDRONIC ACID AND COLECALCIFEROL**

Tab Orl 70 mg / 5 600 IU

Fosavance 02314940 ORG ACDEFGV

Apo-Alendronate/Vitamin D3 02454475 APX ACDEFGV

Jamp Alendronate/Vitamin D3 02519836 JPC ACDEFGV

Teva-Alendronate/Cholecalciferol 02403641 TEV ACDEFGV

**M05BX OTHER DRUGS AFFECTING BONE STRUCTURE AND MINERALIZATION****M05BX04 DENOSUMAB**

Liq SC 60 mg/mL

Prolia 02343541 AGA (SA)

Liq SC 120 mg / 1.7 mL

Xgeva 02368153 AGA (SA)

**M05BX05 BUROSUMAB**

Liq SC 10 mg/mL

Crysvita 02483629 UGX (SA)

Liq SC 20 mg/mL

Crysvita 02483637 UGX (SA)

Liq SC 30 mg/mL

Crysvita 02483645 UGX (SA)

**M09 OTHER DRUGS FOR DISORDERS OF THE MUSCULO-SKELETAL SYSTEM****M09A OTHER DRUGS FOR DISORDERS OF THE MUSCULO-SKELETAL SYSTEM****M09AX OTHER DRUGS FOR DISORDERS OF THE MUSCULO-SKELETAL SYSTEM****M09AX07 NUSINERSEN**

Liq INT 2.4 mg/mL

Spinraza 02465663 BIG (SA)

**M09AX09 ONASEMNOGENE ABEPARVOVEC**Liq IV 2 x 10<sup>13</sup> vg/mL

Zolgensma 02509695 NVR (SA)

**M09AX10 RISDIPLAM**

Pws Orl 0.75 mg/mL

Evrysdi 02514931 HLR (SA)

**N NERVOUS SYSTEM****N01 ANAESTHETICS****N01B LOCAL ANAESTHETICS****N01BB AMIDES****N01BB02 LIDOCAINE**

Liq Orl 2%

Lidodan Viscous 01968823 ODN ACDEFGV

**N01BB09 ROPIVACAINE**

Liq Prt 5 mg/mL

Naropin 02229415 APN ACDEFGV

Liq Prt 10 mg/mL

Naropin 02229418 APN ACDEFGV

**N01BX OTHER LOCAL ANAESTHETICS**

N01BX04 CAPSAICIN

Crm Top 0.025%

Zostrix 00740306 BSH AEEFGV

Capsaicin 02157101 BSL AEEFGV

Crm Top 0.075%

Zostrix H.P. 02004240 BSH AEEFGV

Capsaicin Crm 02157128 BSL AEEFGV

**N02 ANALGESICS****N02A OPIOIDS****N02AA NATURAL OPIUM ALKALOIDS**

N02AA01 MORPHINE

MORPHINE SULFATE

Liq Inj 10 mg/mL

Morphine Sulfate 00392588 SDZ ACDEFGVW

Liq Inj 15 mg/mL

Morphine Sulfate 00392561 SDZ ACDEFGVW

Liq Inj 50 mg/mL

Morphine HP 50 00617288 SDZ ACDEFGVW

SRC Orl 10 mg

Kadian 02242163 BGP ACDEFGVW

M-Eslon 02019930 ETH ACDEFGVW

SRC Orl 15 mg

M-Eslon 15 02177749 SAV ACDEFGVW

SRC Orl 20 mg

Kadian 02184435 BGP ACDEFGVW

SRC Orl 30 mg

M-Eslon 02019949 SAV ACDEFGVW

SRC Orl 50 mg

Kadian 02184443 BGP ACDEFGVW

SRC Orl 60 mg

M-Eslon 02019957 SAV ACDEFGVW

SRC Orl 100 mg

Kadian 02184451 BGP ACDEFGVW

M-Eslon 02019965 SAV ACDEFGVW

SRC Orl 200 mg

M-Eslon 02177757 SAV ACDEFGVW

SRT Orl 15 mg

MS Contin 02015439 PFR ACDEFGVW

Sandoz Morphine SR 02244790 SDZ ACDEFGVW

Teva-Morphine SR 02302764 TEV ACDEFGVW

N02AA01	MORPHINE							
	MORPHINE SULFATE							
SRT	Orl	30 mg	MS Contin	02014297	PFR	ACDEFGVW		
			Sandoz Morphine SR	02244791	SDZ	ACDEFGVW		
			Teva-Morphine SR	02302772	TEV	ACDEFGVW		
SRT	Orl	60 mg	MS Contin	02014300	PFR	ACDEFGVW		
			Sandoz Morphine SR	02244792	SDZ	ACDEFGVW		
			Teva-Morphine SR	02302780	TEV	ACDEFGVW		
SRT	Orl	100 mg	MS Contin	02014319	PFR	ACDEFGVW		
			Sandoz Morphine SR	02478889	SDZ	ACDEFGVW		
			Teva-Morphine SR	02302799	TEV	ACDEFGVW		
SRT	Orl	200 mg	MS Contin	02014327	PFR	ACDEFGVW		
			Sandoz Morphine SR	02478897	SDZ	ACDEFGVW		
			Teva-Morphine SR	02302802	TEV	ACDEFGVW		
Syr	Orl	1 mg/mL	Doloral	00614491	ATL	ACDEFGVW		
Syr	Orl	5 mg/mL	Doloral	00614505	ATL	ACDEFGVW		
Tab	Orl	5 mg	MS IR	02014203	PFR	ACDEFGVW		
			Statex	00594652	PAL	ACDEFGVW		
Tab	Orl	10 mg	MS IR	02014211	PFR	ACDEFGVW		
			Statex	00594644	PAL	ACDEFGVW		
Tab	Orl	20 mg	MS IR	02014238	PFR	ACDEFGVW		
Tab	Orl	30 mg	MS IR	02014254	PFR	ACDEFGVW		
Tab	Orl	50 mg	Statex (Disc/non disp Sep 2/23)	00675962	PAL	ACDEFGVW		
N02AA03	HYDROMORPHONE							
Liq	Inj	2 mg/mL	Dilaudid	00627100	PFR	ACDEFGVW		
			Hydromorphone Hydrochloride	02145901	SDZ	ACDEFGVW		
Liq	Inj	10 mg/mL	Dilaudid HP	00622133	PFR	ACDEFGVW		
			Hydromorphone HP 10	02145928	SDZ	ACDEFGVW		
Liq	Inj	20 mg/mL	Hydromorphone HP 20	02145936	SDZ	ACDEFGVW		

N02AA03 HYDROMORPHONE

Liq	Inj	50 mg/mL	Hydromorphone HP 50	02146126	SDZ	ACDEFGVW
			Hydromorphone Hydrochloride	02469413	STR	ACDEFGVW
SRC	Orl	3 mg	Hydromorph Contin	02125323	PFR	ACDEFGVW
SRC	Orl	4.5 mg	Hydromorph Contin	02359502	PFR	ACDEFGVW
SRC	Orl	6 mg	Hydromorph Contin	02125331	PFR	ACDEFGVW
SRC	Orl	9 mg	Hydromorph Contin	02359510	PFR	ACDEFGVW
SRC	Orl	12 mg	Hydromorph Contin	02125366	PFR	ACDEFGVW
SRC	Orl	18 mg	Hydromorph Contin	02243562	PFR	ACDEFGVW
SRC	Orl	24 mg	Hydromorph Contin	02125382	PFR	ACDEFGVW
SRC	Orl	30 mg	Hydromorph Contin	02125390	PFR	ACDEFGVW
Syr	Orl	1 mg/mL	pms-Hydromorphone	01916386	PMS	ACDEFGVW
Tab	Orl	1 mg	Dilaudid	00705438	PFR	ACDEFGVW
			Apo-Hydromorphone	02364115	APX	ACDEFGVW
			pms-Hydromorphone	00885444	PMS	ACDEFGVW
Tab	Orl	2 mg	Dilaudid	00125083	PFR	ACDEFGVW
			Apo-Hydromorphone	02364123	APX	ACDEFGVW
			pms-Hydromorphone	00885436	PMS	ACDEFGVW
Tab	Orl	4 mg	Dilaudid	00125121	PFR	ACDEFGVW
			Apo-Hydromorphone	02364131	APX	ACDEFGVW
			pms-Hydromorphone	00885401	PMS	ACDEFGVW
Tab	Orl	8 mg	Dilaudid	00786543	PFR	ACDEFGVW
			Apo-Hydromorphone	02364158	APX	ACDEFGVW
			pms-Hydromorphone	00885428	PMS	ACDEFGVW

N02AA05 OXYCODONE

ERT	Orl	10 mg	Oxyneo	02372525	PFR	W
ERT	Orl	15 mg	Oxyneo	02372533	PFR	W

N02AA05 OXYCODONE

ERT	Orl	20 mg	Oxyneo	02372797	PFR	W
ERT	Orl	30 mg	Oxyneo	02372541	PFR	W
ERT	Orl	40 mg	Oxyneo	02372568	PFR	W
ERT	Orl	60 mg	Oxyneo	02372576	PFR	W
ERT	Orl	80 mg	Oxyneo	02372584	PFR	W
Sup	Rt	10 mg	Supeudol	00392480	SDZ	ACDEFGV
Tab	Orl	5 mg	Oxy-IR	02231934	PFR	W (SA)
			Supeudol	00789739	SDZ	W (SA)
			pms-Oxycodone IR	02319977	PMS	W (SA)
Tab	Orl	10 mg	Oxy-IR	02240131	PFR	W (SA)
			Supeudol	00443948	SDZ	W (SA)
			pms-Oxycodone IR	02319985	PMS	W (SA)
Tab	Orl	20 mg	Oxy-IR	02240132	PFR	W (SA)
			Supeudol	02262983	SDZ	W (SA)
			pms-Oxycodone IR	02319993	PMS	W (SA)

N02AA59 CODEINE, COMBINATIONS, EXCLUDING PSYCHOLEPTICS  
ACETAMINOPHEN / CAFFEINE / CODEINE

Tab	Orl	300 mg / 15 mg / 30 mg	Teva-Lenoltec #3	00653276	TEV	ACDEFGVW
Tab	Orl	300 mg / 30 mg	Teva-Emtec-30	00608882	TEV	ACDEFGVW
Tab	Orl	300 mg / 60 mg	Teva-Lenoltec #4	00621463	TEV	ACDEFGVW

**N02AB PHENYLPIPERIDINE DERIVATIVES**

N02AB03 FENTANYL

Pth	Trd	12 mcg/hr	Sandoz Fentanyl patch	02327112	SDZ	W (SA)
			Teva-Fentanyl	02311925	TEV	W (SA)
Pth	Trd	25 mcg/hr	Sandoz Fentanyl	02327120	SDZ	W (SA)
			Teva-Fentanyl	02282941	TEV	W (SA)
Pth	Trd	37 mcg/hr	Sandoz Fentanyl	02327139	SDZ	W

N02AB03	FENTANYL							
Pth	Trd	50 mcg/hr	Sandoz Fentanyl	02327147	SDZ	W	(SA)	
			Teva-Fentanyl	02282968	TEV	W	(SA)	
Pth	Trd	75 mcg/hr	Sandoz Fentanyl	02327155	SDZ	W	(SA)	
			Teva-Fentanyl	02282976	TEV	W	(SA)	
Pth	Trd	100 mcg/hr	Sandoz Fentanyl	02327163	SDZ	W	(SA)	
			Teva-Fentanyl	02282984	TEV	W	(SA)	

**N02B OTHER ANALGESICS AND ANTIPYRETICS**

**N02BA SALICYLIC ACID AND DERIVATIVES**

N02BA01 ACETYLSALICYLIC ACID

ECT	Orl	81 mg	ASA	02433044	PMS	EV	
			ASA	02449277	TLI	EV	
			ASA EC	02244993	PMS	EV	
			ASA EC	02426811	SAS	EV	
			Equate daily low-dose EC	02243801	PMS	EV	
			Exact Coated daily low dose ASA	02243896	PMS	EV	
			Jamp-ASA EC	02427206	JPC	EV	
			Praxis ASA	02283700	PMS	EV	
ECT	Orl	325 mg	ASATAB EC	02352427	ODN	AEFGV	
			Enteric Coated ASA	02010526	VTH	AEFGV	

**N02BE ANILIDES**

N02BE01 PARACETAMOL (ACETAMINOPHEN)

Sup	Rt	120 mg	Acet - 120	02230434	PDP	G	
Tab	Orl	325 mg	Acetaminophen	02252805	CCM	G	
			Novo-Gesic	00389218	TEV	G	
Tab	Orl	500 mg	Acetaminophen	02252813	CCM	G	
			Novo-Gesic	00482323	TEV	G	

N02BE51 PARACETAMOL (ACETAMINOPHEN), COMBINATIONS EXCLUDING PSYCHOLEPTICS

ACETAMINOPHEN / CAFFEINE / CODEINE

Tab	Orl	300 mg / 15 mg / 15 mg	Teva-Lenoltec #2	00653241	TEV	ACDEFGVW	
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ACETAMINOPHEN / OXYCODONE

N02BE51 PARACETAMOL (ACETAMINOPHEN), COMBINATIONS EXCLUDING PSYCHOLEPTICS

ACETAMINOPHEN / OXYCODONE

Tab	Orl	325 mg / 5 mg	Apo-Oxycodone/Acet	02324628	APX	ACDEFGVW
			Sandoz Oxycodone/Acetaminophen	02307898	SDZ	ACDEFGVW
			Teva-Oxycocet	00608165	TEV	ACDEFGVW

**N02C ANTIMIGRAINE PREPARATIONS**

**N02CA ERGOT ALKALOIDS**

N02CA01 DIHYDROERGOTAMINE

Liq	Nas	4 mg/mL	Migranal	02228947	STR	ACDEFGV
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**N02CC SELECTIVE 5HT1-RECEPTOR AGONISTS**

N02CC01 SUMATRIPTAN

Liq	SC	6 mg / 0.5 mL	Imitrex	02212188	GSK	(SA)
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Taro-Sumatriptan	02361698	TAR	(SA)
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Spr	Nas	5 mg	Imitrex	02230418	GSK	(SA)
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Spr	Nas	20 mg	Imitrex	02230420	GSK	(SA)
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Tab	Orl	50 mg	Imitrex DF	02212153	GSK	ACDEFGV
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Apo-Sumatriptan	02268388	APX	ACDEFGV
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Mylan-Sumatriptan	02268914	MYL	ACDEFGV
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pms-Sumatriptan	02256436	PMS	ACDEFGV
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Sumatriptan	02286521	SAS	ACDEFGV
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Sumatriptan DF	02385570	SIV	ACDEFGV
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Teva-Sumatriptan DF	02286823	TEV	ACDEFGV
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Tab	Orl	100 mg	Imitrex DF	02212161	GSK	ACDEFGV
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Apo-Sumatriptan	02268396	APX	ACDEFGV
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Mylan-Sumatriptan	02268922	MYL	ACDEFGV
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pms-Sumatriptan	02256444	PMS	ACDEFGV
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Sumatriptan	02286548	SAS	ACDEFGV
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Sumatriptan DF	02385589	SIV	ACDEFGV
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Teva-Sumatriptan	02239367	TEV	ACDEFGV
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Teva-Sumatriptan DF	02286831	TEV	ACDEFGV
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N02CC02 NARATRIPTAN

Tab	Orl	1 mg	Teva-Naratriptan	02314290	TEV	(SA)
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N02CC02 NARATRIPTAN

Tab Orl 2.5 mg

Amerge (Disc/non disp Dec 14/23) 02237821 GSK (SA)  
 Sandoz Naratriptan 02322323 SDZ (SA)  
 Teva-Naratriptan 02314304 TEV (SA)

N02CC03 ZOLMITRIPTAN

ODT Orl 2.5 mg

Zomig Rapimelt 02243045 XPI ACDEFGV  
 Jamp-Zolmitriptan ODT 02428237 JPC ACDEFGV  
 pms-Zolmitriptan ODT 02324768 PMS ACDEFGV  
 Sandoz Zolmitriptan ODT 02362996 SDZ ACDEFGV  
 Septa-Zolmitriptan ODT 02428474 SPT ACDEFGV  
 Teva-Zolmitriptan OD 02342545 TEV ACDEFGV  
 Zolmitriptan ODT 02442671 SAS ACDEFGV

Spr Nas 2.5 mg

Zomig 02248992 XPI (SA)

Spr Nas 5 mg

Zomig Nasal 02248993 XPI (SA)

Tab Orl 2.5 mg

Zomig 02238660 XPI ACDEFGV  
 Auro-Zolmitriptan 02481030 ARO ACDEFGV  
 CCP-Zolmitriptan 02458780 CCM ACDEFGV  
 Jamp-Zolmitriptan 02421623 JPC ACDEFGV  
 Jamp-Zolmitriptan 02477106 JPC ACDEFGV  
 Mar-Zolmitriptan 02399458 MAR ACDEFGV  
 Nat-Zolmitriptan 02421534 NAT ACDEFGV  
 pms-Zolmitriptan 02324229 PMS ACDEFGV  
 Sandoz Zolmitriptan 02362988 SDZ ACDEFGV  
 Teva-Zolmitriptan 02313960 TEV ACDEFGV  
 Zolmitriptan 02442655 SAS ACDEFGV

N02CC04 RIZATRIPTAN

N02CC04 RIZATRIPTAN

ODT Orl 5 mg

Maxalt RPD 02240518 ORG ACDEFGV  
 CCP-Rizatriptan ODT 02458764 CCM ACDEFGV  
 Jamp-Rizatriptan ODT 02465086 JPC ACDEFGV  
 Mar-Rizatriptan ODT 02462788 MAR ACDEFGV  
 Mylan-Rizatriptan ODT 02379198 MYL ACDEFGV  
 Nat-Rizatriptan ODT 02436604 NAT ACDEFGV  
 pms-Rizatriptan RDT 02393360 PMS ACDEFGV  
 Rizatriptan ODT 02442906 SAS ACDEFGV  
 Rizatriptan ODT 02446111 SIV ACDEFGV  
 Sandoz Rizatriptan ODT 02351870 SDZ ACDEFGV  
 Teva-Rizatriptan ODT 02396661 TEV ACDEFGV

ODT Orl 10 mg

Maxalt RPD 02240519 ORG ACDEFGV  
 CCP-Rizatriptan ODT 02458772 CCM ACDEFGV  
 Jamp-Rizatriptan ODT 02465094 JPC ACDEFGV  
 Mar-Rizatriptan ODT 02462796 MAR ACDEFGV  
 Mylan-Rizatriptan ODT 02379201 MYL ACDEFGV  
 Nat-Rizatriptan ODT 02436612 NAT ACDEFGV  
 pms-Rizatriptan RDT 02393379 PMS ACDEFGV  
 Rizatriptan ODT 02442914 SAS ACDEFGV  
 Rizatriptan ODT 02446138 SIV ACDEFGV  
 Sandoz Rizatriptan ODT 02351889 SDZ ACDEFGV  
 Teva-Rizatriptan ODT 02396688 TEV ACDEFGV

Tab Orl 5 mg

Apo-Rizatriptan 02393468 APX ACDEFGV

Tab Orl 10 mg

Maxalt 02240521 ORG ACDEFGV  
 Act Rizatriptan 02381702 TEV ACDEFGV  
 Apo-Rizatriptan 02393476 APX ACDEFGV  
 Auro-Rizatriptan 02441144 ARO ACDEFGV  
 Jamp-Rizatriptan 02380463 JPC ACDEFGV  
 Mar-Rizatriptan 02379678 MAR ACDEFGV  
 Rizatriptan 02516756 SAS ACDEFGV

N02CC05 ALMOTRIPTAN

Tab Orl 12.5 mg

Almotriptan 02466821 SAS ACDEFGV  
 Mylan-Almotriptan 02398443 MYL ACDEFGV  
 Sandoz Almotriptan 02405334 SDZ ACDEFGV  
 Teva-Almotriptan 02434849 TEV ACDEFGV

**N02CC06 ELETRIPTAN**

Tab Orl 20 mg

Relpax 02256290 UJC ACDEFGV

Apo-Eletriptan 02386054 APX ACDEFGV

Apo-Eletriptan Tablets 02518015 APX ACDEFGV

Auro-Eletriptan 02479451 ARO ACDEFGV

Eletriptan 02511266 SAS ACDEFGV

Jamp Eletriptan 02493683 JPC ACDEFGV

Teva-Eletriptan 02382091 TEV ACDEFGV

Tab Orl 40 mg

Relpax 02256304 UJC ACDEFGV

Apo-Eletriptan 02386062 APX ACDEFGV

Apo-Eletriptan Tablets 02518023 APX ACDEFGV

Auro-Eletriptan 02479478 ARO ACDEFGV

Eletriptan 02511274 SAS ACDEFGV

Jamp Eletriptan 02493691 JPC ACDEFGV

Teva-Eletriptan 02382105 TEV ACDEFGV

**N02CD CALCITONIN GENE-RELATED PEPTIDE (CGRP) ANTAGONISTS****N02CD02 GALCANEZUMAB**

Liq SC 120 mg / mL

Emgality (autoinjector) 02491087 LIL (SA)

Emgality (prefilled syringe) 02491060 LIL (SA)

**N02CD03 FREMANEZUMAB**

Liq SC 225 mg / 1.5 mL

Ajovy (autoinjector) 02509474 TEV (SA)

Ajovy (prefilled syringe) 02497859 TEV (SA)

**N02CX OTHER ANTIMIGRAINE PREPARATIONS****N02CX01 PIZOTIFEN**

Tab Orl 1 mg

Sandomigran DS 00511552 PAL ACDEFGV

**N03 ANTIEPILEPTICS****N02B OTHER ANALGESICS AND ANTIPYRETICS****N02BF GABAPENTINOIDS****N02BF01 GABAPENTIN**

N02BF01 GABAPENTIN

Cap Orl 100 mg

Neurontin 02084260 BGP ACDEFGVW  
 Apo-Gabapentin 02244304 APX ACDEFGVW  
 Auro-Gabapentin 02321203 ARO ACDEFGVW  
 Gabapentin 02416840 AHI ACDEFGVW  
 Gabapentin 02353245 SAS ACDEFGVW  
 Gabapentin 02246314 SIV ACDEFGVW  
 Jamp-Gabapentin 02361469 JPC ACDEFGVW  
 Mar-Gabapentin 02391473 MAR ACDEFGVW  
 pms-Gabapentin 02243446 PMS ACDEFGVW  
 Teva-Gabapentin 02244513 TEV ACDEFGVW

Cap Orl 300 mg

Neurontin 02084279 BGP ACDEFGVW  
 Apo-Gabapentin 02244305 APX ACDEFGVW  
 Auro-Gabapentin 02321211 ARO ACDEFGVW  
 Gabapentin 02416859 AHI ACDEFGVW  
 Gabapentin 02353253 SAS ACDEFGVW  
 Gabapentin 02246315 SIV ACDEFGVW  
 Jamp-Gabapentin 02361485 JPC ACDEFGVW  
 Mar-Gabapentin 02391481 MAR ACDEFGVW  
 pms-Gabapentin 02243447 PMS ACDEFGVW  
 Teva-Gabapentin 02244514 TEV ACDEFGVW

Cap Orl 400 mg

Neurontin 02084287 BGP ACDEFGVW  
 Apo-Gabapentin 02244306 APX ACDEFGVW  
 Auro-Gabapentin 02321238 ARO ACDEFGVW  
 Gabapentin 02416867 AHI ACDEFGVW  
 Gabapentin 02353261 SAS ACDEFGVW  
 Gabapentin 02246316 SIV ACDEFGVW  
 Jamp-Gabapentin 02361493 JPC ACDEFGVW  
 Mar-Gabapentin 02391503 MAR ACDEFGVW  
 pms-Gabapentin 02243448 PMS ACDEFGVW  
 Teva-Gabapentin 02244515 TEV ACDEFGVW

N02BF01 GABAPENTIN

Tab Orl 600 mg

Neurontin 02239717 BGP ACDEFGVW  
 Apo-Gabapentin 02293358 APX ACDEFGVW  
 Auro-Gabapentin 02428334 ARO ACDEFGVW  
 Gabapentin 02392526 AHI ACDEFGVW  
 Gabapentin 02410990 GLM ACDEFGVW  
 Gabapentin 02432072 JPC ACDEFGVW  
 Gabapentin 02431289 SAS ACDEFGVW  
 Gabapentin 02388200 SIV ACDEFGVW  
 Jamp-Gabapentin 02402289 JPC ACDEFGVW  
 Teva-Gabapentin 02248457 TEV ACDEFGVW

Tab Orl 800 mg

Neurontin 02239718 BGP ACDEFGVW  
 Apo-Gabapentin 02293366 APX ACDEFGVW  
 Auro-Gabapentin 02428342 ARO ACDEFGVW  
 Gabapentin 02392534 AHI ACDEFGVW  
 Gabapentin 02411008 GLM ACDEFGVW  
 Gabapentin 02432080 JPC ACDEFGVW  
 Gabapentin 02431297 SAS ACDEFGVW  
 Gabapentin 02388219 SIV ACDEFGVW  
 Jamp-Gabapentin 02402297 JPC ACDEFGVW  
 Teva-Gabapentin 02247346 TEV ACDEFGVW

**N03A ANTIEPILEPTICS**

**N03AA BARBITURATES AND DERIVATIVES**

N03AA02 PHENOBARBITAL

Elx Orl 5 mg/mL

Phenobarbital 00645575 PDP ACDEFGV

Liq Inj 30 mg/mL

Phenobarbital Sodium 02304082 SDZ ACDEFGVW

Liq Inj 120 mg/mL

Phenobarbital Sodium 02304090 SDZ ACDEFGVW

Tab Orl 15 mg

Phenobarbital 00178799 PDP ACDEFGV

Tab Orl 30 mg

Phenobarbital 00178802 PDP ACDEFGV

Tab Orl 60 mg

Phenobarbital 00178810 PDP ACDEFGV

Tab Orl 100 mg

Phenobarbital 00178829 PDP ACDEFGV

N03AA03 PRIMIDONE

Tab Orl 125 mg

Primidone 00399310 AAP ACDEFGV

N03AA03 PRIMIDONE

Tab Orl 250 mg

Primidone 00396761 AAP ACDEFGV

**N03AB HYDANTOIN DERIVATIVES**

N03AB02 PHENYTOIN

Cap Orl 30 mg

Dilantin 00022772 BGP ACDEFGV

Cap Orl 100 mg

Dilantin 00022780 BGP ACDEFGV

Phenytoin Sodium 02460912 AAP ACDEFGV

Liq Inj 50 mg/mL

Phenytoin Sodium 00780626 SDZ V

Sus Orl 30 mg / 5 mL

Dilantin 30 00023442 BGP ACDEFGV

Sus Orl 125 mg / 5 mL

Dilantin 125 00023450 BGP ACDEFGV

Taro-Phenytoin 02250896 TAR ACDEFGV

Tab Orl 50 mg

Dilantin infatabs 00023698 BGP ACDEFGV

**N03AD SUCCINIMIDE DERIVATIVES**

N03AD01 ETHOSUXIMIDE

Cap Orl 250 mg

Zarontin 00022799 ERF ACDEFGV

Syr Orl 50 mg/mL

Zarontin 00023485 ERF ACDEFGV

**N03AE BENZODIAZEPINE DERIVATIVES**

N03AE01 CLONAZEPAM

Tab Orl 0.25 mg

pms-Clonazepam 02179660 PMS ACDEFGV

Tab Orl 0.5 mg

Rivotril 00382825 XPI ACDEFGV

Apo-Clonazepam 02177889 APX ACDEFGV

pms-Clonazepam R 02207818 PMS ACDEFGV

Tab Orl 1 mg

pms-Clonazepam 02048728 PMS ACDEFGV

Tab Orl 2 mg

Rivotril 00382841 XPI ACDEFGV

Apo-Clonazepam 02177897 APX ACDEFGV

pms-Clonazepam 02048736 PMS ACDEFGV

**N03AF CARBOXAMIDE DERIVATIVES**

N03AF01 CARBAMAZEPINE

N03AF01		CARBAMAZEPINE							
SRT	Orl	200 mg		Tegretol CR	00773611	NVR	ACDEFGV		
				Sandoz Carbamazepine CR	02261839	SDZ	ACDEFGV		
SRT	Orl	400 mg		Tegretol CR	00755583	NVR	ACDEFGV		
				Sandoz Carbamazepine CR	02261847	SDZ	ACDEFGV		
Sus	Orl	100 mg / 5 mL		Tegretol	02194333	NVR	ACDEFGV		
				Taro-Carbamazepine	02367394	TAR	ACDEFGV		
Tab	Orl	200 mg		Tegretol	00010405	NVR	ACDEFGV		
				Teva-Carbamazepine	00782718	TEV	ACDEFGV		
TabC	Orl	100 mg		Taro-Carbamazepine Chewable	02244403	TAR	ACDEFGV		
TabC	Orl	200 mg		Taro-Carbamazepine Chewable	02244404	TAR	ACDEFGV		
N03AF02		OXCARBAZEPINE							
Sus	Orl	60 mg/mL		Trileptal	02244673	NVR	(SA)		
Tab	Orl	150 mg		Apo-Oxcarbazepine	02284294	APX	(SA)		
Tab	Orl	300 mg		Trileptal	02242068	NVR	(SA)		
				Apo-Oxcarbazepine	02284308	APX	(SA)		
Tab	Orl	600 mg		Trileptal	02242069	NVR	(SA)		
				Apo-Oxcarbazepine	02284316	APX	(SA)		
N03AF03		RUFINAMIDE							
Tab	Orl	100 mg		Banzel	02369613	EIS	(SA)		
Tab	Orl	200 mg		Banzel	02369621	EIS	(SA)		
Tab	Orl	400 mg		Banzel	02369648	EIS	(SA)		
N03AF04		ESLICARBAZEPINE							
Tab	Orl	200 mg		Aptiom	02426862	SNV	(SA)		
Tab	Orl	400 mg		Aptiom	02426870	SNV	(SA)		
Tab	Orl	600 mg		Aptiom	02426889	SNV	(SA)		
Tab	Orl	800 mg		Aptiom	02426897	SNV	(SA)		

**N03AG FATTY ACID DERIVATIVES**

## N03AG01 VALPROIC ACID

Cap	Orl	250 mg	Apo-Valproic	02238048	APX	ACDEFGV
			pms-Valproic Acid	02230768	PMS	ACDEFGV
ECC	Orl	500 mg	pms-Valproic Acid	02229628	PMS	ACDEFGV
ECT	Orl	125 mg	Epival	00596418	BGP	ACDEFGV
			Apo-Divalproex	02239698	APX	ACDEFGV
			Mylan-Divalproex	02458926	MYL	ACDEFGV
ECT	Orl	250 mg	Epival	00596426	BGP	ACDEFGV
			Apo-Divalproex	02239699	APX	ACDEFGV
			Mylan-Divalproex	02458934	MYL	ACDEFGV
ECT	Orl	500 mg	Epival	00596434	BGP	ACDEFGV
			Apo-Divalproex	02239700	APX	ACDEFGV
			Mylan-Divalproex	02459019	MYL	ACDEFGV
Syr	Orl	250 mg / 5 mL	Depakene	00443832	BGP	ACDEFGV
			Apo-Valproic Acid	02238370	APX	ACDEFGV
			pms-Valproic	02236807	PMS	ACDEFGV

## N03AG04 VIGABATRIN

Pws	Orl	500 mg	Sabril	02068036	LBK	(SA)
Tab	Orl	500 mg	Sabril	02065819	LBK	(SA)

**N03AX OTHER ANTIEPILEPTICS**

## N03AX09 LAMOTRIGINE

Tab	Orl	25 mg	Lamictal	02142082	GSK	ACDEFGV
			Apo-Lamotrigine	02245208	APX	ACDEFGV
			Auro-Lamotrigine	02381354	ARO	ACDEFGV
			Lamotrigine	02343010	SAS	ACDEFGV
			Lamotrigine	02428202	SIV	ACDEFGV
			Mylan-Lamotrigine	02265494	MYL	ACDEFGV
			pms-Lamotrigine	02246897	PMS	ACDEFGV



N03AX09 LAMOTRIGINE

Tab Orl 100 mg

Lamictal 02142104 GSK ACDEFGV  
 Apo-Lamotrigine 02245209 APX ACDEFGV  
 Auro-Lamotrigine 02381362 ARO ACDEFGV  
 Lamotrigine 02343029 SAS ACDEFGV  
 Lamotrigine 02428210 SIV ACDEFGV  
 Mylan-Lamotrigine 02265508 MYL ACDEFGV  
 pms-Lamotrigine 02246898 PMS ACDEFGV

Tab Orl 150 mg

Lamictal 02142112 GSK ACDEFGV  
 Apo-Lamotrigine 02245210 APX ACDEFGV  
 Auro-Lamotrigine 02381370 ARO ACDEFGV  
 Lamotrigine 02343037 SAS ACDEFGV  
 Lamotrigine 02428229 SIV ACDEFGV  
 Mylan-Lamotrigine 02265516 MYL ACDEFGV  
 pms-Lamotrigine 02246899 PMS ACDEFGV

TabC Orl 2 mg

Lamictal Chewtabs 02243803 GSK ACDEFGV

TabC Orl 5 mg

Lamictal Chewtabs 02240115 GSK ACDEFGV

N03AX11 TOPIRAMATE

Cap Orl 15 mg

Topamax 02239907 JAN (SA)

Cap Orl 25 mg

Topamax 02239908 JAN (SA)

Tab Orl 25 mg

Topamax 02230893 JAN ACDEFGV  
 Apo-Topiramate 02279614 APX ACDEFGV  
 Auro-Topiramate 02345803 ARO ACDEFGV  
 GLN-Topiramate 02287765 GLM ACDEFGV  
 Jamp Topiramate Tablets 02345250 JPC ACDEFGV  
 Jamp-Topiramate 02435608 JPC ACDEFGV  
 Mint-Topiramate 02315645 MNT ACDEFGV  
 Mylan-Topiramate 02263351 MYL ACDEFGV  
 pms-Topiramate 02262991 PMS ACDEFGV  
 Sandoz Topiramate Tablets (Disc/non disp Apr 11/24) 02431807 SDZ ACDEFGV  
 Teva-Topiramate 02248860 TEV ACDEFGV  
 Topiramate 02395738 AHI ACDEFGV  
 Topiramate 02356856 SAS ACDEFGV  
 Topiramate 02389460 SIV ACDEFGV

N03AX11 TOPIRAMATE

Tab Orl 100 mg

Topamax	02230894	JAN	ACDEFGV
Apo-Topiramate	02279630	APX	ACDEFGV
Auro-Topiramate	02345838	ARO	ACDEFGV
GLN-Topiramate	02287773	GLM	ACDEFGV
Jamp-Topiramate	02435616	JPC	ACDEFGV
Mint-Topiramate	02315653	MNT	ACDEFGV
Mylan-Topiramate	02263378	MYL	ACDEFGV
pms-Topiramate	02263009	PMS	ACDEFGV
Sandoz Topiramate Tablets (Disc/non disp Apr 11/24)	02431815	SDZ	ACDEFGV
Teva-Topiramate	02248861	TEV	ACDEFGV
Topiramate	02395746	AHI	ACDEFGV
Topiramate	02356864	SAS	ACDEFGV
Topiramate	02389487	SIV	ACDEFGV

Tab Orl 200 mg

Topamax	02230896	JAN	ACDEFGV
Apo-Topiramate	02279649	APX	ACDEFGV
Auro-Topiramate	02345846	ARO	ACDEFGV
GLN-Topiramate	02287781	GLM	ACDEFGV
Jamp-Topiramate	02435624	JPC	ACDEFGV
Mint-Topiramate	02315661	MNT	ACDEFGV
Mylan-Topiramate	02263386	MYL	ACDEFGV
pms-Topiramate	02263017	PMS	ACDEFGV
Sandoz Topiramate Tablets (Disc/non disp Apr 11/24)	02431823	SDZ	ACDEFGV
Teva-Topiramate	02248862	TEV	ACDEFGV
Topiramate	02395754	AHI	ACDEFGV
Topiramate	02356872	SAS	ACDEFGV

N03AX14 LEVETIRACETAM

Liq Orl 100 mg/mL

pdp-Levetiracetam	02490447	PDP	(SA)
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N03AX14 LEVETIRACETAM

Tab Orl 250 mg

	Keppra	02247027	UCB	ACDEFGV
	Act Levetiracetam	02274183	TEV	ACDEFGV
	Apo-Levetiracetam	02285924	APX	ACDEFGV
	Auro-Levetiracetam	02375249	ARO	ACDEFGV
Jamp	Levetiracetam Tablets	02504553	JPC	ACDEFGV
	Jamp-Levetiracetam	02403005	JPC	ACDEFGV
	Levetiracetam	02454653	PMS	ACDEFGV
	Levetiracetam	02353342	SAS	ACDEFGV
	Levetiracetam	02442531	SIV	ACDEFGV
	Levetiracetam Tablets	02399776	AHI	ACDEFGV
	M-Levetiracetam	02524562	MRA	ACDEFGV
	Mint-Levetiracetam	02442388	MNT	ACDEFGV
	Nat-Levetiracetam	02440202	NAT	ACDEFGV
	NRA-Levetiracetam	02499193	NRA	ACDEFGV
	pms-Levetiracetam	02296101	PMS	ACDEFGV
	Riva-Levetiracetam	02482274	RIV	ACDEFGV
	Sandoz Levetiracetam	02461986	SDZ	ACDEFGV

Tab Orl 500 mg

	Keppra	02247028	UCB	ACDEFGV
	Act Levetiracetam	02274191	TEV	ACDEFGV
	Apo-Levetiracetam	02285932	APX	ACDEFGV
	Auro-Levetiracetam	02375257	ARO	ACDEFGV
Jamp	Levetiracetam Tablets	02504561	JPC	ACDEFGV
	Jamp-Levetiracetam	02403021	JPC	ACDEFGV
	Levetiracetam	02454661	PMS	ACDEFGV
	Levetiracetam	02353350	SAS	ACDEFGV
	Levetiracetam	02442558	SIV	ACDEFGV
	Levetiracetam Tablets	02399784	AHI	ACDEFGV
	M-Levetiracetam	02524570	MRA	ACDEFGV
	Mint-Levetiracetam	02442396	MNT	ACDEFGV
	Nat-Levetiracetam	02440210	NAT	ACDEFGV
	NRA-Levetiracetam	02499207	NRA	ACDEFGV
	pms-Levetiracetam	02296128	PMS	ACDEFGV
	Pro-Levetiracetam	02311380	PDL	ACDEFGV
	Riva-Levetiracetam	02482282	RIV	ACDEFGV
	Sandoz Levetiracetam	02461994	SDZ	ACDEFGV

N03AX14 LEVETIRACETAM

Tab Orl 750 mg

Keppra	02247029	UCB	ACDEFGV
Act Levetiracetam	02274205	TEV	ACDEFGV
Apo-Levetiracetam	02285940	APX	ACDEFGV
Auro-Levetiracetam	02375265	ARO	ACDEFGV
Jamp Levetiracetam Tablets	02504588	JPC	ACDEFGV
Jamp-Levetiracetam	02403048	JPC	ACDEFGV
Levetiracetam	02454688	PMS	ACDEFGV
Levetiracetam	02353369	SAS	ACDEFGV
Levetiracetam	02442566	SIV	ACDEFGV
Levetiracetam Tablets	02399792	AHI	ACDEFGV
M-Levetiracetam	02524589	MRA	ACDEFGV
Mint-Levetiracetam	02442418	MNT	ACDEFGV
Nat-Levetiracetam	02440229	NAT	ACDEFGV
NRA-Levetiracetam	02499215	NRA	ACDEFGV
pms-Levetiracetam	02296136	PMS	ACDEFGV
Pro-Levetiracetam	02311399	PDL	ACDEFGV
Riva-Levetiracetam	02482290	RIV	ACDEFGV
Sandoz Levetiracetam	02462001	SDZ	ACDEFGV

Tab Orl 1000 mg

Sandoz Levetiracetam	02462028	SDZ	ACDEFGV
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N03AX16 PREGABALIN

Cap Orl 25 mg

Lyrice	02268418	BGP	ACDEFGVW
Ach-Pregabalin	02449838	AHI	ACDEFGVW
Apo-Pregabalin	02394235	APX	ACDEFGVW
Auro-Pregabalin	02433869	ARO	ACDEFGVW
Jamp-Pregabalin	02435977	JPC	ACDEFGVW
M-Pregabalin	02467291	MRA	ACDEFGVW
Mint-Pregabalin	02423804	MNT	ACDEFGVW
Nat-Pregabalin	02494841	NAT	ACDEFGVW
NRA-Pregabalin	02479117	NRA	ACDEFGVW
pms-Pregabalin	02359596	PMS	ACDEFGVW
Pregabalin	02396483	PDL	ACDEFGVW
Pregabalin	02405539	SAS	ACDEFGVW
Pregabalin	02403692	SIV	ACDEFGVW
Sandoz Pregabalin	02390817	SDZ	ACDEFGVW
Taro-Pregabalin	02392801	SUN	ACDEFGVW
Teva-Pregabalin	02361159	TEV	ACDEFGVW

N03AX16 PREGABALIN

Cap Orl 50 mg

Lyrica	02268426	BGP	ACDEFGVW
Ach-Pregabalin	02449846	AHI	ACDEFGVW
Apo-Pregabalin	02394243	APX	ACDEFGVW
Auro-Pregabalin	02433877	ARO	ACDEFGVW
Jamp-Pregabalin	02435985	JPC	ACDEFGVW
M-Pregabalin	02467305	MRA	ACDEFGVW
Mint-Pregabalin	02423812	MNT	ACDEFGVW
Nat-Pregabalin	02494868	NAT	ACDEFGVW
NRA-Pregabalin	02479125	NRA	ACDEFGVW
pms-Pregabalin	02359618	PMS	ACDEFGVW
Pregabalin	02396505	PDL	ACDEFGVW
Pregabalin	02405547	SAS	ACDEFGVW
Pregabalin	02403706	SIV	ACDEFGVW
Sandoz Pregabalin	02390825	SDZ	ACDEFGVW
Taro-Pregabalin	02392828	SUN	ACDEFGVW
Teva-Pregabalin	02361175	TEV	ACDEFGVW

Cap Orl 75 mg

Lyrica	02268434	BGP	ACDEFGVW
Ach-Pregabalin	02449854	AHI	ACDEFGVW
Apo-Pregabalin	02394251	APX	ACDEFGVW
Auro-Pregabalin	02433885	ARO	ACDEFGVW
Jamp-Pregabalin	02435993	JPC	ACDEFGVW
M-Pregabalin	02467313	MRA	ACDEFGVW
Mint-Pregabalin	02424185	MNT	ACDEFGVW
Nat-Pregabalin	02494876	NAT	ACDEFGVW
NRA-Pregabalin	02479133	NRA	ACDEFGVW
pms-Pregabalin	02359626	PMS	ACDEFGVW
Pregabalin	02396513	PDL	ACDEFGVW
Pregabalin	02405555	SAS	ACDEFGVW
Pregabalin	02403714	SIV	ACDEFGVW
Sandoz Pregabalin	02390833	SDZ	ACDEFGVW
Taro-Pregabalin	02392836	SUN	ACDEFGVW
Teva-Pregabalin	02361183	TEV	ACDEFGVW

N03AX16 PREGABALIN

Cap Orl 150 mg

Lyricea 02268450 BGP ACDEFGVW  
 Apo-Pregabalin 02394278 APX ACDEFGVW  
 Auro-Pregabalin 02433907 ARO ACDEFGVW  
 Jamp-Pregabalin 02436000 JPC ACDEFGVW  
 M-Pregabalin 02467321 MRA ACDEFGVW  
 Mint-Pregabalin 02424207 MNT ACDEFGVW  
 Nat-Pregabalin 02494884 NAT ACDEFGVW  
 NRA-Pregabalin 02479168 NRA ACDEFGVW  
 pms-Pregabalin 02359634 PMS ACDEFGVW  
 Pregabalin 02396521 PDL ACDEFGVW  
 Pregabalin 02405563 SAS ACDEFGVW  
 Pregabalin 02403722 SIV ACDEFGVW  
 Sandoz Pregabalin 02390841 SDZ ACDEFGVW  
 Taro-Pregabalin 02392844 SUN ACDEFGVW  
 Teva-Pregabalin 02361205 TEV ACDEFGVW

Cap Orl 225 mg

Lyricea 02268477 BGP ACDEFGVW  
 Ach-Pregabalin 02449897 AHI ACDEFGVW  
 Apo-Pregabalin 02394286 APX ACDEFGVW  
 Nat-Pregabalin 02494892 NAT ACDEFGVW  
 pms-Pregabalin 02398079 PMS ACDEFGVW  
 Teva-Pregabalin 02361221 TEV ACDEFGVW

Cap Orl 300 mg

Lyricea 02268485 BGP ACDEFGVW  
 Ach-Pregabalin 02449900 AHI ACDEFGVW  
 Apo-Pregabalin 02394294 APX ACDEFGVW  
 Jamp-Pregabalin 02436019 JPC ACDEFGVW  
 Nat-Pregabalin 02494906 NAT ACDEFGVW  
 pms-Pregabalin 02359642 PMS ACDEFGVW  
 Pregabalin 02396548 PDL ACDEFGVW  
 Pregabalin 02405598 SAS ACDEFGVW  
 Pregabalin 02403730 SIV ACDEFGVW  
 Sandoz Pregabalin 02390868 SDZ ACDEFGVW  
 Taro-Pregabalin 02392860 SUN ACDEFGVW  
 Teva-Pregabalin 02361248 TEV ACDEFGVW

N03AX17 STIRIPENTOL

Cap Orl 250 mg

Diacomit 02398958 BOX (SA)

Cap Orl 500 mg

Diacomit 02398966 BOX (SA)

N03AX17 STIRIPENTOL

Pws Orl 250 mg

Diacomit 02398974 BOX (SA)

Pws Orl 500 mg

Diacomit 02398982 BOX (SA)

N03AX18 LACOSAMIDE

Tab Orl 50 mg

Vimpat 02357615 UCB ACDEFGV  
 ACH-Lacosamide 02489287 AHI ACDEFGV  
 Auro-Lacosamide 02475332 ARO ACDEFGV  
 Jamp-Lacosamide 02488388 JPC ACDEFGV  
 Lacosamide 02512874 SAS ACDEFGV  
 Mar-Lacosamide 02487802 MAR ACDEFGV  
 Mint-Lacosamide 02490544 MNT ACDEFGV  
 NRA-Lacosamide 02499568 NRA ACDEFGV  
 pharma-Lacosamide 02478196 PMS ACDEFGV  
 Sandoz-Lacosamide 02474670 SDZ ACDEFGV  
 Teva-Lacosamide 02472902 TEV ACDEFGV

Tab Orl 100 mg

Vimpat 02357623 UCB ACDEFGV  
 ACH-Lacosamide 02489295 AHI ACDEFGV  
 Auro-Lacosamide 02475340 ARO ACDEFGV  
 Jamp-Lacosamide 02488396 JPC ACDEFGV  
 Lacosamide 02512882 SAS ACDEFGV  
 Mar-Lacosamide 02487810 MAR ACDEFGV  
 Mint-Lacosamide 02490552 MNT ACDEFGV  
 NRA-Lacosamide 02499576 NRA ACDEFGV  
 pharma-Lacosamide 02478218 PMS ACDEFGV  
 Sandoz-Lacosamide 02474689 SDZ ACDEFGV  
 Teva-Lacosamide 02472910 TEV ACDEFGV

Tab Orl 150 mg

Vimpat 02357631 UCB ACDEFGV  
 ACH-Lacosamide 02489309 AHI ACDEFGV  
 Auro-Lacosamide 02475359 ARO ACDEFGV  
 Jamp-Lacosamide 02488418 JPC ACDEFGV  
 Lacosamide 02512890 SAS ACDEFGV  
 Mar-Lacosamide 02487829 MAR ACDEFGV  
 Mint-Lacosamide 02490560 MNT ACDEFGV  
 NRA-Lacosamide 02499584 NRA ACDEFGV  
 pharma-Lacosamide 02478226 PMS ACDEFGV  
 Sandoz-Lacosamide 02474697 SDZ ACDEFGV  
 Teva-Lacosamide 02472929 TEV ACDEFGV

N03AX18 LACOSAMIDE

Tab Orl 200 mg

Vimpat 02357658 UCB ACDEFGV  
 ACH-Lacosamide 02489317 AHI ACDEFGV  
 Auro-Lacosamide 02475367 ARO ACDEFGV  
 Jamp-Lacosamide 02488426 JPC ACDEFGV  
 Lacosamide 02512904 SAS ACDEFGV  
 Mar-Lacosamide 02487837 MAR ACDEFGV  
 Mint-Lacosamide 02490579 MNT ACDEFGV  
 NRA-Lacosamide 02499592 NRA ACDEFGV  
 pharma-Lacosamide 02478234 PMS ACDEFGV  
 Sandoz-Lacosamide 02474700 SDZ ACDEFGV  
 Teva-Lacosamide 02472937 TEV ACDEFGV

N03AX22 PERAMPANEL

Tab Orl 2 mg

Fycompa 02404516 EIS (SA)

Tab Orl 4 mg

Fycompa 02404524 EIS (SA)

Tab Orl 6 mg

Fycompa 02404532 EIS (SA)

Tab Orl 8 mg

Fycompa 02404540 EIS (SA)

Tab Orl 10 mg

Fycompa 02404559 EIS (SA)

Tab Orl 12 mg

Fycompa 02404567 EIS (SA)

N03AX23 BRIVARACETAM

Tab Orl 10 mg

Brivlera 02452936 UCB (SA)

Tab Orl 25 mg

Brivlera 02452944 UCB (SA)

Tab Orl 50 mg

Brivlera 02452952 UCB (SA)

Tab Orl 75 mg

Brivlera 02452960 UCB (SA)

Tab Orl 100 mg

Brivlera 02452979 UCB (SA)

**N04 ANTI-PARKINSON DRUGS**

**N04A ANTI-CHOLINERGIC AGENTS**

**N04AA TERTIARY AMINES**

N04AA01 TRIHEXYPHENIDYL

Tab Orl 2 mg

Trihex 00545058 AAP ACDEFGV



N04AA01	TRIHEXYPHENIDYL							
Tab	Orl	5 mg		Trihex	00545074	AAP	ACDEFGV	
N04AA04	PROCYCLIDINE							
Elx	Orl	2.5 mg / 5 mL		pdp-Procyclidine	00587362	PDP	ACDEFGV	
Tab	Orl	2.5 mg		pdp-Procyclidine	00649392	PDP	ACDEFGV	
Tab	Orl	5 mg		pdp-Procyclidine	00587354	PDP	ACDEFGV	
N04AA05	PROFENAMINE (ETHOPROPAZINE)							
Tab	Orl	50 mg		Parsitan	01927744	SLP	ACDEFGV	

**N04AC ETHERS OF TROPINE OR TROPINE DERIVATIVES**

N04AC01	BENZATROPINE							
Liq	Inj	1 mg/mL		Benztropine Omega	02238903	OMG	ACDEFGV	
Tab	Orl	1 mg		pdp-Benztropine	00706531	PDP	ACDEFGV	
Tab	Orl	2 mg		pdp-Benztropine	00426857	PDP	ACDEFGV	

**N04B DOPAMINERGIC AGENTS**

**N04BA DOPA AND DOPA DERIVATIVES**

N04BA02	LEVODOPA AND DECARBOXYLASE INHIBITOR							
	LEVODOPA / BENSERAZIDE							
Cap	Orl	50 mg / 12.5 mg		Prolopa	00522597	HLR	ACDEFGV	
Cap	Orl	100 mg / 25 mg		Prolopa	00386464	HLR	ACDEFGV	
Cap	Orl	200 mg / 50 mg		Prolopa	00386472	HLR	ACDEFGV	
	LEVODOPA / CARBIDOPA							
Gel	Itt	20 mg / 5 mg/mL		Duodopa	02292165	ABV	(SA)	
SRT	Orl	100 mg / 25 mg		AA-Levocarb CR	02272873	AAP	ACDEFGV	
SRT	Orl	200 mg / 50 mg		AA-Levocarb CR	02245211	AAP	ACDEFGV	
Tab	Orl	100 mg / 10 mg		Apo-Levocarb	02195933	APX	ACDEFGV	
				Auro-Levocarb	02531593	ARO	ACDEFGV	
				Mint-Levocarb	02457954	MNT	ACDEFGV	
				Teva-Levocarbido	02244494	TEV	ACDEFGV	

N04BA02 LEVODOPA AND DECARBOXYLASE INHIBITOR

LEVODOPA / CARBIDOPA

Tab	Orl	100 mg / 25 mg	Apo-Levocarb	02195941	APX	ACDEFGV
			Auro-Levocarb	02531607	ARO	ACDEFGV
			Mint-Levocarb	02457962	MNT	ACDEFGV
			Teva-Levocarbido	02244495	TEV	ACDEFGV
Tab	Orl	250 mg / 25 mg	Apo-Levocarb	02195968	APX	ACDEFGV
			Auro-Levocarb	02531615	ARO	ACDEFGV
			Mint-Levocarb	02457970	MNT	ACDEFGV
			Teva-Levocarbido	02244496	TEV	ACDEFGV

N04BA03 LEVODOPA, DECARBOXYLASE INHIBITOR AND COMT INHIBITOR

LEVODOPA, CARBIDOPA, ENTACAPONE

Tab	Orl	50 mg / 12.5 mg / 200 mg	Stalevo	02305933	SDZ	(SA)
Tab	Orl	75 mg / 18.75 mg / 200 mg	Stalevo	02337827	SDZ	(SA)
Tab	Orl	100 mg / 25 mg / 200 mg	Stalevo	02305941	SDZ	(SA)
Tab	Orl	125 mg / 31.25 mg / 200 mg	Stalevo	02337835	SDZ	(SA)
Tab	Orl	150 mg / 37.5 mg / 200 mg	Stalevo	02305968	SDZ	(SA)

**N04BB ADAMANTANE DERIVATIVES**

N04BB01 AMANTADINE

Cap	Orl	100 mg	pdp-Amantadine Hydrochloride	01990403	PDP	ACDEFGV
Syr	Orl	10 mg/mL	Odan-Amantadine Syrup	02538601	ODN	ACDEFGV
			pdp-Amantadine	02022826	PDP	ACDEFGV

**N04BC DOPAMINE AGONISTS**

N04BC04 ROPINIROLE

Tab	Orl	0.25 mg	Jamp-Ropinirole	02352338	JPC	ACDEFV
			Ran-Ropinirole	02314037	RAN	ACDEFV
			Ropinirole (Disc/non disp Dec 6/23)	02353040	SAS	ACDEFV
			Teva-Ropinirole	02316846	TEV	ACDEFV
Tab	Orl	1 mg	Jamp-Ropinirole	02352346	JPC	ACDEFV
			Ran-Ropinirole	02314053	RAN	ACDEFV
			Teva-Ropinirole	02316854	TEV	ACDEFV

N04BC04		ROPINIROLE						
Tab	Orl	2 mg		Jamp-Ropinirole	02352354	JPC	ACDEFV	
				Ran-Ropinirole	02314061	RAN	ACDEFV	
				Teva-Ropinirole	02316862	TEV	ACDEFV	
Tab	Orl	5 mg		Ran-Ropinirole	02314088	RAN	ACDEFV	
				Teva-Ropinirole	02316870	TEV	ACDEFV	
N04BC05		PRAMIPEXOLE						
Tab	Orl	0.25 mg		Mirapex	02237145	BOE	ACDEFV	
				Act Pramipexole	02297302	TEV	ACDEFV	
				Apo-Pramipexole	02292378	APX	ACDEFV	
				Auro-Pramipexole	02424061	ARO	ACDEFV	
				Pramipexole	02367602	SAS	ACDEFV	
				Pramipexole	02309122	SIV	ACDEFV	
				Sandoz Pramipexole	02315262	SDZ	ACDEFV	
Tab	Orl	0.5 mg		Act Pramipexole	02297310	TEV	ACDEFV	
				Apo-Pramipexole	02292386	APX	ACDEFV	
				Auro-Pramipexole	02424088	ARO	ACDEFV	
				Pramipexole	02367610	SAS	ACDEFV	
				Pramipexole	02309130	SIV	ACDEFV	
				Sandoz Pramipexole	02315270	SDZ	ACDEFV	
Tab	Orl	1 mg		Act Pramipexole	02297329	TEV	ACDEFV	
				Apo-Pramipexole	02292394	APX	ACDEFV	
				Auro-Pramipexole	02424096	ARO	ACDEFV	
				Pramipexole	02367629	SAS	ACDEFV	
				Pramipexole	02309149	SIV	ACDEFV	
				Sandoz Pramipexole	02315289	SDZ	ACDEFV	
Tab	Orl	1.5 mg		Act Pramipexole	02297337	TEV	ACDEFV	
				Apo-Pramipexole	02292408	APX	ACDEFV	
				Auro-Pramipexole	02424118	ARO	ACDEFV	
				Pramipexole	02367645	SAS	ACDEFV	
				Pramipexole	02309157	SIV	ACDEFV	
				Sandoz Pramipexole	02315297	SDZ	ACDEFV	
N04BC07		APOMORPHINE						
Liq	SC	30 mg / 3 mL		Movapo (Disc/non disp Jun 30/24)	02459132	PAL	(SA)	

**N04BC07 APOMORPHINE**

ODF	Orl	10 mg	Kynmobi	02500264	SNV	(SA)
ODF	Orl	15 mg	Kynmobi	02500272	SNV	(SA)
ODF	Orl	20 mg	Kynmobi	02500280	SNV	(SA)
ODF	Orl	25 mg	Kynmobi	02500299	SNV	(SA)
ODF	Orl	30 mg	Kynmobi	02500302	SNV	(SA)

**N04BC09 ROTIGOTINE**

Pth	Trd	2 mg	Neupro	02403900	UCB	(SA)
Pth	Trd	4 mg	Neupro	02403927	UCB	(SA)
Pth	Trd	6 mg	Neupro	02403935	UCB	(SA)
Pth	Trd	8 mg	Neupro	02403943	UCB	(SA)

**N04BD MONOAMINE OXIDASE TYPE B INHIBITORS****N04BD01 SELEGILINE**

Tab	Orl	5 mg	Novo-Selegiline	02068087	TEV	ACDEFV
			Selegiline	02230641	AAP	ACDEFV

**N04BX OTHER DOPAMINERGIC AGENTS****N04BX02 ENTACAPONE**

Tab	Orl	200 mg	Comtan	02243763	SDZ	ACDEFGV
			Sandoz Entacapone	02380005	SDZ	ACDEFGV
			Teva-Entacapone	02375559	TEV	ACDEFGV

**N05 PSYCHOLEPTICS****N05A ANTIPSYCHOTICS****N05AA PHENOTHIAZINE WITH ALIPHATIC SIDE CHAIN****N05AA01 CHLORPROMAZINE**

Tab	Orl	25 mg	Teva-Chlorpromazine	00232823	TEV	ACDEFGVW
Tab	Orl	50 mg	Teva-Chlorpromazine	00232807	TEV	ACDEFGVW
Tab	Orl	100 mg	Teva-Chlorpromazine	00232831	TEV	ACDEFGVW

**N05AA02 LEVOMEPRMAZINE (METHOTRIMEPRAZINE)**

Liq	Inj	25 mg/mL	Nozinan	01927698	SAV	ACDEFVW
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**N05AA02 LEVOMEPRMAZINE (METHOTRIMEPRAZINE)**

Tab	Orl	2 mg	Methoprazine	02238403	AAP	ACDEFGVW
Tab	Orl	5 mg	Methoprazine	02238404	AAP	ACDEFGVW
Tab	Orl	25 mg	Methoprazine	02238405	AAP	ACDEFGVW
Tab	Orl	50 mg	Methoprazine	02238406	AAP	ACDEFGVW

**N05AB PHENOTHIAZINE WITH PIPERAZINE STRUCTURE****N05AB02 FLUPHENAZINE**

Tab	Orl	1 mg	Fluphenazine	00405345	AAP	ACDEFGV
Tab	Orl	2 mg	Fluphenazine	00410632	AAP	ACDEFGV
Tab	Orl	5 mg	Fluphenazine	00405361	AAP	ACDEFGV

**N05AB03 PERPHENAZINE**

Tab	Orl	2 mg	Perphenazine	00335134	AAP	ACDEFGV
Tab	Orl	4 mg	Perphenazine	00335126	AAP	ACDEFGV
Tab	Orl	8 mg	Perphenazine	00335118	AAP	ACDEFGV
Tab	Orl	16 mg	Perphenazine	00335096	AAP	ACDEFGV

**N05AB04 PROCHLORPERAZINE**

Sup	Rt	10 mg	Sandoz Prochlorperazine	00789720	SDZ	ACDEFGV
Tab	Orl	5 mg	Prochlorazine	00886440	AAP	ACDEFGV
Tab	Orl	10 mg	Prochlorazine	00886432	AAP	ACDEFGV

**N05AB06 TRIFLUOPERAZINE**

Tab	Orl	1 mg	Trifluoperazine	00345539	AAP	ACDEFGV
Tab	Orl	2 mg	Trifluoperazine	00312754	AAP	ACDEFGV
Tab	Orl	5 mg	Trifluoperazine	00312746	AAP	ACDEFGV
Tab	Orl	10 mg	Trifluoperazine	00326836	AAP	ACDEFGV

**N05AC PHENOTHIAZINE WITH PIPERIDINE STRUCTURE**

N05AC01	PERICYAZINE					
Cap	Orl	5 mg	Neuleptil	01926780	SLP	ACDEFGV
Cap	Orl	10 mg	Neuleptil	01926772	SLP	ACDEFGV
Cap	Orl	20 mg	Neuleptil	01926764	SLP	ACDEFGV
Dps	Orl	10 mg/mL	Neuleptil	01926756	SLP	ACDEFGV

**N05AD BUTYROPHENONE DERIVATIVES**

N05AD01	HALOPERIDOL					
Liq	Inj	5 mg/mL	Haloperidol	00808652	SDZ	ACDEFGVW
			Haloperidol Injection	02366010	OMG	ACDEFGVW
Liq	Inj	100 mg/mL	Haloperidol LA	02130300	SDZ	ACDEFGVW
Tab	Orl	0.5 mg	Teva-Haloperidol	00363685	TEV	ACDEFGVW
Tab	Orl	1 mg	Teva-Haloperidol	00363677	TEV	ACDEFGVW
Tab	Orl	2 mg	Teva-Haloperidol	00363669	TEV	ACDEFGVW
Tab	Orl	5 mg	Teva-Haloperidol	00363650	TEV	ACDEFGVW
Tab	Orl	10 mg	Teva-Haloperidol	00713449	TEV	ACDEFGVW

**N05AE INDOLE DERIVATIVES**

N05AE04	ZIPRASIDONE					
Cap	Orl	20 mg	Zeldox	02298597	UJC	ACDEFGV
			Auro-Ziprasidone	02449544	ARO	ACDEFGV
Cap	Orl	40 mg	Zeldox	02298600	UJC	ACDEFGV
			Auro-Ziprasidone	02449552	ARO	ACDEFGV
Cap	Orl	60 mg	Zeldox	02298619	UJC	ACDEFGV
			Auro-Ziprasidone	02449560	ARO	ACDEFGV
Cap	Orl	80 mg	Zeldox	02298627	UJC	ACDEFGV
			Auro-Ziprasidone	02449579	ARO	ACDEFGV

N05AE05 LURASIDONE

N05AE05 LURASIDONE

Tab	Orl	20 mg	Latuda	02422050	SNV	ACDEFGV
			Jamp Lurasidone	02516438	JPC	ACDEFGV
			pms-Lurasidone	02505878	PMS	ACDEFGV
			Sandoz Lurasidone	02521075	SDZ	ACDEFGV
			Taro-Lurasidone	02504499	TAR	ACDEFGV
Tab	Orl	40 mg	Latuda	02387751	SNV	ACDEFGV
			Jamp Lurasidone	02516446	JPC	ACDEFGV
			pms-Lurasidone	02505886	PMS	ACDEFGV
			Sandoz Lurasidone	02521091	SDZ	ACDEFGV
			Taro-Lurasidone	02504502	TAR	ACDEFGV
Tab	Orl	60 mg	Latuda	02413361	SNV	ACDEFGV
			Jamp Lurasidone	02516454	JPC	ACDEFGV
			pms-Lurasidone	02505894	PMS	ACDEFGV
			Sandoz Lurasidone	02521105	SDZ	ACDEFGV
			Taro-Lurasidone	02504510	TAR	ACDEFGV
Tab	Orl	80 mg	Latuda	02387778	SNV	ACDEFGV
			Jamp Lurasidone	02516462	JPC	ACDEFGV
			pms-Lurasidone	02505908	PMS	ACDEFGV
			Sandoz Lurasidone	02521113	SDZ	ACDEFGV
			Taro-Lurasidone	02504529	TAR	ACDEFGV
Tab	Orl	120 mg	Latuda	02387786	SNV	ACDEFGV
			Jamp Lurasidone	02516470	JPC	ACDEFGV
			pms-Lurasidone	02505916	PMS	ACDEFGV
			Sandoz Lurasidone	02521121	SDZ	ACDEFGV
			Taro-Lurasidone	02504537	TAR	ACDEFGV

**N05AF THIOXANTHENE DERIVATIVES**

N05AF01 FLUPENTHIXOL

Liq	Inj	20 mg/mL	Fluanxol Depot	02156032	VLH	ACDEFGV
Liq	Inj	100 mg/mL	Fluanxol Depot	02156040	VLH	ACDEFGV
Tab	Orl	0.5 mg	Fluanxol	02156008	VLH	ACDEFGV
Tab	Orl	3 mg	Fluanxol	02156016	VLH	ACDEFGV

N05AF05 ZUCLOPENTHIXOL

N05AF05 ZUCLOPENTHIXOL

Liq Inj 200 mg/mL

Clopixol Depot 02230406 VLH ACDEFGV

Tab Orl 10 mg

Clopixol 02230402 VLH ACDEFGV

Tab Orl 25 mg

Clopixol 02230403 VLH ACDEFGV

**N05AG DIPHENYLBUTYLPIPERIDINE DERIVATIVES**

N05AG02 PIMOZIDE

Tab Orl 2 mg

Pimozide 02245432 AAP ACDEFGV

Tab Orl 4 mg

Pimozide 02245433 AAP ACDEFGV

**N05AH DIAZEPINES, OXAZEPINES, THIAZEPINES AND OXEPINES**

N05AH01 LOXAPINE

Tab Orl 2.5 mg

Xylac 02242868 PDP ACDEFGV

Tab Orl 10 mg

Xylac 02230838 PDP ACDEFGV

Tab Orl 25 mg

Xylac 02230839 PDP ACDEFGV

N05AH02 CLOZAPINE

Tab Orl 25 mg

Clozaril 00894737 HLS ACDEFGV

AA-Clozapine 02248034 AAP ACDEFGV

Gen-Clozapine 02247243 MYL ACDEFGV

Tab Orl 50 mg

Clozaril 02490668 HLS ACDEFGV

AA-Clozapine 02458748 AAP ACDEFGV

Gen-Clozapine 02305003 MYL ACDEFGV

Tab Orl 100 mg

Clozaril 00894745 HLS ACDEFGV

AA-Clozapine 02248035 AAP ACDEFGV

Gen-Clozapine 02247244 MYL ACDEFGV

Tab Orl 200 mg

Clozaril 02490676 HLS ACDEFGV

AA-Clozapine 02458756 AAP ACDEFGV

Gen-Clozapine 02305011 MYL ACDEFGV

N05AH03 OLANZAPINE



N05AH03 OLANZAPINE

ODT Orl 5 mg

Zyprexa Zydis	02243086	LIL	ACDEFGVW
Apo-Olanzapine ODT	02360616	APX	ACDEFGVW
Auro-Olanzapine ODT	02448726	ARO	ACDEFGVW
Jamp-Olanzapine ODT	02406624	JPC	ACDEFGVW
Mint-Olanzapine ODT	02436965	MNT	ACDEFGVW
Olanzapine ODT	02338645	PDL	ACDEFGVW
Olanzapine ODT	02352974	SAS	ACDEFGVW
Olanzapine ODT	02343665	SIV	ACDEFGVW
pms-Olanzapine ODT	02303191	PMS	ACDEFGVW
Sandoz Olanzapine ODT	02327775	SDZ	ACDEFGVW

ODT Orl 10 mg

Zyprexa Zydis	02243087	LIL	ACDEFGVW
Apo-Olanzapine ODT	02360624	APX	ACDEFGVW
Auro-Olanzapine ODT	02448734	ARO	ACDEFGVW
Jamp-Olanzapine ODT	02406632	JPC	ACDEFGVW
Mint-Olanzapine ODT	02436973	MNT	ACDEFGVW
Olanzapine ODT	02338653	PDL	ACDEFGVW
Olanzapine ODT	02352982	SAS	ACDEFGVW
Olanzapine ODT	02343673	SIV	ACDEFGVW
pms-Olanzapine ODT	02303205	PMS	ACDEFGVW
Sandoz Olanzapine ODT	02327783	SDZ	ACDEFGVW

ODT Orl 15 mg

Zyprexa Zydis	02243088	LIL	ACDEFGVW
Apo-Olanzapine ODT	02360632	APX	ACDEFGVW
Auro-Olanzapine ODT	02448742	ARO	ACDEFGVW
Jamp-Olanzapine ODT	02406640	JPC	ACDEFGVW
Mint-Olanzapine ODT	02436981	MNT	ACDEFGVW
Olanzapine ODT	02338661	PDL	ACDEFGVW
Olanzapine ODT	02352990	SAS	ACDEFGVW
Olanzapine ODT	02343681	SIV	ACDEFGVW
pms-Olanzapine ODT	02303213	PMS	ACDEFGVW
Sandoz Olanzapine ODT	02327791	SDZ	ACDEFGVW

N05AH03 OLANZAPINE

ODT Orl 20 mg

Zyprexa Zydis 02243089 LIL ACDEFGVW  
 Apo-Olanzapine ODT 02360640 APX ACDEFGVW  
 Auro-Olanzapine ODT 02448750 ARO ACDEFGVW  
 Jamp-Olanzapine ODT 02406659 JPC ACDEFGVW  
 Olanzapine ODT 02425114 PDL ACDEFGVW  
 Olanzapine ODT 02343703 SIV ACDEFGVW  
 Sandoz Olanzapine ODT 02327805 SDZ ACDEFGVW

Tab Orl 2.5 mg

Zyprexa 02229250 LIL ACDEFGVW  
 Apo-Olanzapine 02281791 APX ACDEFGVW  
 Jamp-Olanzapine FC 02417243 JPC ACDEFGVW  
 Mint-Olanzapine 02410141 MNT ACDEFGVW  
 Olanzapine 02311968 PDL ACDEFGVW  
 Olanzapine 02372819 SAS ACDEFGVW  
 Olanzapine 02385864 SIV ACDEFGVW  
 pms-Olanzapine 02303116 PMS ACDEFGVW  
 Sandoz Olanzapine 02310341 SDZ ACDEFGVW  
 Teva-Olanzapine 02276712 TEV ACDEFGVW

Tab Orl 5 mg

Zyprexa 02229269 LIL ACDEFGVW  
 Apo-Olanzapine 02281805 APX ACDEFGVW  
 Jamp-Olanzapine FC 02417251 JPC ACDEFGVW  
 Mint-Olanzapine 02410168 MNT ACDEFGVW  
 Olanzapine 02311976 PDL ACDEFGVW  
 Olanzapine 02372827 SAS ACDEFGVW  
 Olanzapine 02385872 SIV ACDEFGVW  
 pms-Olanzapine 02303159 PMS ACDEFGVW  
 Sandoz Olanzapine 02310368 SDZ ACDEFGVW  
 Teva-Olanzapine 02276720 TEV ACDEFGVW

N05AH03 OLANZAPINE

Tab Orl 7.5 mg

Zyprexa	02229277	LIL	ACDEFGVW
Apo-Olanzapine	02281813	APX	ACDEFGVW
Jamp-Olanzapine FC	02417278	JPC	ACDEFGVW
Mint-Olanzapine	02410176	MNT	ACDEFGVW
Olanzapine	02311984	PDL	ACDEFGVW
Olanzapine	02372835	SAS	ACDEFGVW
Olanzapine	02385880	SIV	ACDEFGVW
pms-Olanzapine	02303167	PMS	ACDEFGVW
Sandoz Olanzapine	02310376	SDZ	ACDEFGVW
Teva-Olanzapine	02276739	TEV	ACDEFGVW

Tab Orl 10 mg

Zyprexa	02229285	LIL	ACDEFGVW
Apo-Olanzapine	02281821	APX	ACDEFGVW
Jamp-Olanzapine FC	02417286	JPC	ACDEFGVW
Mint-Olanzapine	02410184	MNT	ACDEFGVW
Olanzapine	02311992	PDL	ACDEFGVW
Olanzapine	02372843	SAS	ACDEFGVW
Olanzapine	02385899	SIV	ACDEFGVW
pms-Olanzapine	02303175	PMS	ACDEFGVW
Sandoz Olanzapine	02310384	SDZ	ACDEFGVW
Teva-Olanzapine	02276747	TEV	ACDEFGVW

Tab Orl 15 mg

Zyprexa	02238850	LIL	ACDEFGVW
Apo-Olanzapine	02281848	APX	ACDEFGVW
Jamp-Olanzapine FC	02417294	JPC	ACDEFGVW
Mint-Olanzapine	02410192	MNT	ACDEFGVW
Olanzapine	02312018	PDL	ACDEFGVW
Olanzapine	02372851	SAS	ACDEFGVW
Olanzapine	02385902	SIV	ACDEFGVW
pms-Olanzapine	02303183	PMS	ACDEFGVW
Sandoz Olanzapine	02310392	SDZ	ACDEFGVW
Teva-Olanzapine	02276755	TEV	ACDEFGVW

N05AH03 OLANZAPINE

Tab Orl 20 mg

Zyprexa	02238851	LIL	ACDEFGVW
Apo-Olanzapine	02333015	APX	ACDEFGVW
Jamp-Olanzapine FC	02417308	JPC	ACDEFGVW
Olanzapine	02421704	PDL	ACDEFGVW
Olanzapine	02385910	SIV	ACDEFGVW
pms-Olanzapine	02367483	PMS	ACDEFGVW
Teva-Olanzapine	02359707	TEV	ACDEFGVW

N05AH04 QUETIAPINE

ERT Orl 50 mg

Seroquel XR	02300184	AZE	ACDEFGVW
ACH-Quetiapine Fumarate XR	02450860	AHI	ACDEFGVW
Apo-Quetiapine XR	02457229	APX	ACDEFGVW
M-Quetiapine Fumarate XR	02527928	MRA	ACDEFGVW
Mint-Quetiapine XR	02522187	MNT	ACDEFGVW
NRA-Quetiapine XR	02510677	NRA	ACDEFGVW
Quetiapine Fumarate XR	02516616	SAS	ACDEFGVW
Quetiapine XR	02519607	JPC	ACDEFGVW
Quetiapine XR	02417359	SIV	ACDEFGVW
Sandoz Quetiapine XR	02407671	SDZ	ACDEFGVW
Teva-Quetiapine XR	02395444	TEV	ACDEFGVW

ERT Orl 150 mg

Seroquel XR	02321513	AZE	ACDEFGVW
ACH-Quetiapine Fumarate XR	02450879	AHI	ACDEFGVW
Apo-Quetiapine XR	02457237	APX	ACDEFGVW
M-Quetiapine Fumarate XR	02527936	MRA	ACDEFGVW
Mint-Quetiapine XR	02522195	MNT	ACDEFGVW
NRA-Quetiapine XR	02510685	NRA	ACDEFGVW
Quetiapine Fumarate XR	02516624	SAS	ACDEFGVW
Quetiapine XR	02519615	JPC	ACDEFGVW
Quetiapine XR	02417367	SIV	ACDEFGVW
Sandoz Quetiapine XR	02407698	SDZ	ACDEFGVW
Teva-Quetiapine XR	02395452	TEV	ACDEFGVW

N05AH04 QUETIAPINE

ERT Orl 200 mg

Seroquel XR	02300192	AZE	ACDEFGVW
ACH-Quetiapine Fumarate XR	02450887	AHI	ACDEFGVW
Apo-Quetiapine XR	02457245	APX	ACDEFGVW
M-Quetiapine Fumarate XR	02527944	MRA	ACDEFGVW
Mint-Quetiapine XR	02522209	MNT	ACDEFGVW
NRA-Quetiapine XR	02510693	NRA	ACDEFGVW
Quetiapine Fumarate XR	02516632	SAS	ACDEFGVW
Quetiapine XR	02519623	JPC	ACDEFGVW
Quetiapine XR	02417375	SIV	ACDEFGVW
Sandoz Quetiapine XR	02407701	SDZ	ACDEFGVW
Teva-Quetiapine XR	02395460	TEV	ACDEFGVW

ERT Orl 300 mg

Seroquel XR	02300206	AZE	ACDEFGVW
ACH-Quetiapine Fumarate XR	02450895	AHI	ACDEFGVW
Apo-Quetiapine XR	02457253	APX	ACDEFGVW
M-Quetiapine Fumarate XR	02527952	MRA	ACDEFGVW
Mint-Quetiapine XR	02522217	MNT	ACDEFGVW
NRA-Quetiapine XR	02510707	NRA	ACDEFGVW
Quetiapine Fumarate XR	02516640	SAS	ACDEFGVW
Quetiapine XR	02519747	JPC	ACDEFGVW
Quetiapine XR	02417383	SIV	ACDEFGVW
Sandoz Quetiapine XR	02407728	SDZ	ACDEFGVW
Teva-Quetiapine XR	02395479	TEV	ACDEFGVW

ERT Orl 400 mg

Seroquel XR	02300214	AZE	ACDEFGVW
ACH-Quetiapine Fumarate XR	02450909	AHI	ACDEFGVW
Apo-Quetiapine XR	02457261	APX	ACDEFGVW
M-Quetiapine Fumarate XR	02527960	MRA	ACDEFGVW
Mint-Quetiapine XR	02522225	MNT	ACDEFGVW
NRA-Quetiapine XR	02510715	NRA	ACDEFGVW
Quetiapine Fumarate XR	02516659	SAS	ACDEFGVW
Quetiapine XR	02519763	JPC	ACDEFGVW
Quetiapine XR	02417391	SIV	ACDEFGVW
Sandoz Quetiapine XR	02407736	SDZ	ACDEFGVW
Teva-Quetiapine XR	02395487	TEV	ACDEFGVW

N05AH04 QUETIAPINE

Tab Orl 25 mg

Seroquel	02236951	AZE	ACDEFGVW
Act Quetiapine	02316080	TEV	ACDEFGVW
Apo-Quetiapine	02313901	APX	ACDEFGVW
Apo-Quetiapine Fumarate	02501635	APX	ACDEFGVW
Auro-Quetiapine	02390205	ARO	ACDEFGVW
Jamp Quetiapine Fumarate	02390140	JPC	ACDEFGVW
Jamp-Quetiapine	02330415	JPC	ACDEFGVW
Mar-Quetiapine	02399822	MAR	ACDEFGVW
Mint-Quetiapine	02438003	MNT	ACDEFGVW
Nat-Quetiapine	02439158	NAT	ACDEFGVW
pms-Quetiapine	02296551	PMS	ACDEFGVW
Pro-Quetiapine	02317346	PDL	ACDEFGVW
Quetiapine	02387794	AHI	ACDEFGVW
Quetiapine	02353164	SAS	ACDEFGVW
Quetiapine	02317893	SIV	ACDEFGVW

Tab Orl 100 mg

Seroquel	02236952	AZE	ACDEFGVW
Act Quetiapine	02316099	TEV	ACDEFGVW
Apo-Quetiapine	02313928	APX	ACDEFGVW
Apo-Quetiapine Fumarate	02501643	APX	ACDEFGVW
Auro-Quetiapine	02390213	ARO	ACDEFGVW
Jamp Quetiapine Fumarate	02390159	JPC	ACDEFGVW
Jamp-Quetiapine	02330423	JPC	ACDEFGVW
Mar-Quetiapine	02399830	MAR	ACDEFGVW
Mint-Quetiapine	02438011	MNT	ACDEFGVW
Nat-Quetiapine	02439166	NAT	ACDEFGVW
pms-Quetiapine	02296578	PMS	ACDEFGVW
Pro-Quetiapine	02317354	PDL	ACDEFGVW
Quetiapine	02387808	AHI	ACDEFGVW
Quetiapine	02353172	SAS	ACDEFGVW
Quetiapine	02317907	SIV	ACDEFGVW

Tab Orl 150 mg

Nat-Quetiapine	02439174	NAT	AEFGVW
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N05AH04 QUETIAPINE

Tab Orl 200 mg

Seroquel	02236953	AZE	ACDEFGVW
Act Quetiapine	02316110	TEV	ACDEFGVW
Apo-Quetiapine	02313936	APX	ACDEFGVW
Apo-Quetiapine Fumarate	02501651	APX	ACDEFGVW
Auro-Quetiapine	02390248	ARO	ACDEFGVW
Jamp Quetiapine Fumarate	02390167	JPC	ACDEFGVW
Jamp-Quetiapine	02330458	JPC	ACDEFGVW
Mar-Quetiapine	02399849	MAR	ACDEFGVW
Mint-Quetiapine	02438046	MNT	ACDEFGVW
Nat-Quetiapine	02439182	NAT	ACDEFGVW
pms-Quetiapine	02296594	PMS	ACDEFGVW
Pro-Quetiapine	02317362	PDL	ACDEFGVW
Quetiapine	02387824	AHI	ACDEFGVW
Quetiapine	02353199	SAS	ACDEFGVW
Quetiapine	02317923	SIV	ACDEFGVW

Tab Orl 300 mg

Seroquel	02244107	AZE	ACDEFGVW
Act Quetiapine	02316129	TEV	ACDEFGVW
Apo-Quetiapine	02313944	APX	ACDEFGVW
Apo-Quetiapine Fumarate	02501678	APX	ACDEFGVW
Auro-Quetiapine	02390256	ARO	ACDEFGVW
Jamp Quetiapine Fumarate	02390175	JPC	ACDEFGVW
Jamp-Quetiapine	02330466	JPC	ACDEFGVW
Mar-Quetiapine	02399857	MAR	ACDEFGVW
Mint-Quetiapine	02438054	MNT	ACDEFGVW
Nat-Quetiapine	02439190	NAT	ACDEFGVW
pms-Quetiapine	02296608	PMS	ACDEFGVW
Pro-Quetiapine	02317370	PDL	ACDEFGVW
Quetiapine	02387832	AHI	ACDEFGVW
Quetiapine	02353202	SAS	ACDEFGVW
Quetiapine	02317931	SIV	ACDEFGVW

N05AH05 ASENAPINE

Slit Orl 5 mg

Saphris (Sublingual) 02374803 ORG (SA)

Slit Orl 10 mg

Saphris (Sublingual) 02374811 ORG (SA)

**N05AN LITHIUM**

N05AN01 LITHIUM

N05AN01 LITHIUM

Cap Orl 150 mg

Carbolith 00461733 BSL ACDEFGV  
 Lithane 02013231 SLP ACDEFGV  
 Apo-Lithium Carbonate 02242837 APX ACDEFGV  
 pms-Lithium Carbonate 02216132 PMS ACDEFGV

Cap Orl 300 mg

Carbolith 00236683 BSL ACDEFGV  
 Lithane 00406775 SLP ACDEFGV  
 Apo-Lithium Carbonate 02242838 APX ACDEFGV  
 pms-Lithium Carbonate 02216140 PMS ACDEFGV

Cap Orl 600 mg

Carbolith 02011239 BSL ACDEFGV

SRT Orl 300 mg

Lithmax SR 02266695 AAP ACDEFGV

**N05AX OTHER ANTIPSYCHOTICS**

N05AX08 RISPERIDONE

Liq Orl 1 mg/mL

Risperdal 02236950 JAN ACDEFGV  
 Jamp-Risperidone 02454319 JPC ACDEFGV  
 pms-Risperidone 02279266 PMS ACDEFGV

Pws IM 12.5 mg

Risperdal Consta 02298465 JAN (SA)

Pws IM 25 mg

Risperdal Consta 02255707 JAN (SA)

Pws IM 37.5 mg

Risperdal Consta 02255723 JAN (SA)

Pws IM 50 mg

Risperdal Consta 02255758 JAN (SA)

Tab Orl 0.25 mg

Apo-Risperidone 02282119 APX ACDEFGV  
 Jamp-Risperidone 02359529 JPC ACDEFGV  
 Mar-Risperidone 02371766 MAR ACDEFGV  
 Mint-Risperidone 02359790 MNT ACDEFGV  
 pms-Risperidone 02252007 PMS ACDEFGV  
 Ran-Risperidone 02328305 SUN ACDEFGV  
 Risperidone 02356880 SAS ACDEFGV  
 Sandoz Risperidone 02303655 SDZ ACDEFGV  
 Teva-Risperidone 02282690 TEV ACDEFGV



N05AX08 RISPERIDONE

Tab Orl 0.5 mg

Apo-Risperidone 02282127 APX ACDEFGV  
 Jamp-Risperidone 02359537 JPC ACDEFGV  
 Mar-Risperidone 02371774 MAR ACDEFGV  
 Mint-Risperidone 02359804 MNT ACDEFGV  
 pms-Risperidone 02252015 PMS ACDEFGV  
 Ran-Risperidone 02328313 SUN ACDEFGV  
 Risperidone 02356899 SAS ACDEFGV  
 Sandoz Risperidone 02303663 SDZ ACDEFGV

Tab Orl 1 mg

Apo-Risperidone 02282135 APX ACDEFGV  
 Jamp-Risperidone 02359545 JPC ACDEFGV  
 Mar-Risperidone 02371782 MAR ACDEFGV  
 Mint-Risperidone 02359812 MNT ACDEFGV  
 pms-Risperidone 02252023 PMS ACDEFGV  
 Ran-Risperidone 02328321 SUN ACDEFGV  
 Risperidone 02356902 SAS ACDEFGV  
 Sandoz Risperidone 02279800 SDZ ACDEFGV  
 Teva-Risperidone 02264196 TEV ACDEFGV

Tab Orl 2 mg

Apo-Risperidone 02282143 APX ACDEFGV  
 Jamp-Risperidone 02359553 JPC ACDEFGV  
 Mar-Risperidone 02371790 MAR ACDEFGV  
 Mint-Risperidone 02359820 MNT ACDEFGV  
 pms-Risperidone 02252031 PMS ACDEFGV  
 Ran-Risperidone 02328348 SUN ACDEFGV  
 Risperidone 02356910 SAS ACDEFGV  
 Sandoz Risperidone 02279819 SDZ ACDEFGV  
 Teva-Risperidone 02264218 TEV ACDEFGV

Tab Orl 3 mg

Apo-Risperidone 02282151 APX ACDEFGV  
 Jamp-Risperidone 02359561 JPC ACDEFGV  
 Mar-Risperidone 02371804 MAR ACDEFGV  
 Mint-Risperidone 02359839 MNT ACDEFGV  
 pms-Risperidone 02252058 PMS ACDEFGV  
 Ran-Risperidone 02328364 SUN ACDEFGV  
 Risperidone 02356929 SAS ACDEFGV  
 Sandoz Risperidone 02279827 SDZ ACDEFGV  
 Teva-Risperidone 02264226 TEV ACDEFGV

N05AX08 RISPERIDONE

Tab Orl 4 mg

Apo-Risperidone 02282178 APX ACDEFGV  
 Jamp-Risperidone 02359588 JPC ACDEFGV  
 Mar-Risperidone 02371812 MAR ACDEFGV  
 Mint-Risperidone 02359847 MNT ACDEFGV  
 pms-Risperidone 02252066 PMS ACDEFGV  
 Risperidone 02356937 SAS ACDEFGV  
 Sandoz Risperidone 02279835 SDZ ACDEFGV  
 Taro-Risperidone 02328372 SUN ACDEFGV  
 Teva-Risperidone 02264234 TEV ACDEFGV

N05AX12 ARIPIPRAZOLE

Pws IM 300 mg

Abilify Maintena 02420864 OTS (SA)

Pws IM 400 mg

Abilify Maintena 02420872 OTS (SA)

Tab Orl 2 mg

Abilify 02322374 OTS ACDEFGV  
 Apo-Aripiprazole 02471086 APX ACDEFGV  
 Aripiprazole 02506688 SAS ACDEFGV  
 Auro-Aripiprazole 02460025 ARO ACDEFGV  
 Mint-Aripiprazole 02483556 MNT ACDEFGV  
 pms-Aripiprazole 02466635 PMS ACDEFGV  
 Sandoz Aripiprazole 02473658 SDZ ACDEFGV

Tab Orl 5 mg

Abilify 02322382 OTS ACDEFGV  
 Apo-Aripiprazole 02471094 APX ACDEFGV  
 Aripiprazole 02506718 SAS ACDEFGV  
 Auro-Aripiprazole 02460033 ARO ACDEFGV  
 Mint-Aripiprazole 02483564 MNT ACDEFGV  
 pms-Aripiprazole 02466643 PMS ACDEFGV  
 Sandoz Aripiprazole 02473666 SDZ ACDEFGV

Tab Orl 10 mg

Abilify 02322390 OTS ACDEFGV  
 Apo-Aripiprazole 02471108 APX ACDEFGV  
 Aripiprazole 02506726 SAS ACDEFGV  
 Auro-Aripiprazole 02460041 ARO ACDEFGV  
 Mint-Aripiprazole 02483572 MNT ACDEFGV  
 pms-Aripiprazole 02466651 PMS ACDEFGV  
 Sandoz Aripiprazole 02473674 SDZ ACDEFGV

N05AX12 ARIPIPRAZOLE

Tab Orl 15 mg

Abilify 02322404 OTS ACDEFGV  
 Apo-Aripiprazole 02471116 APX ACDEFGV  
 Aripiprazole 02506734 SAS ACDEFGV  
 Auro-Aripiprazole 02460068 ARO ACDEFGV  
 Mint-Aripiprazole 02483580 MNT ACDEFGV  
 pms-Aripiprazole 02466678 PMS ACDEFGV  
 Sandoz Aripiprazole 02473682 SDZ ACDEFGV

Tab Orl 20 mg

Abilify 02322412 OTS ACDEFGV  
 Apo-Aripiprazole 02471124 APX ACDEFGV  
 Aripiprazole 02506750 SAS ACDEFGV  
 Auro-Aripiprazole 02460076 ARO ACDEFGV  
 Mint-Aripiprazole 02483599 MNT ACDEFGV  
 pms-Aripiprazole 02466686 PMS ACDEFGV  
 Sandoz Aripiprazole 02473690 SDZ ACDEFGV

Tab Orl 30 mg

Abilify 02322455 OTS ACDEFGV  
 Apo-Aripiprazole 02471132 APX ACDEFGV  
 Aripiprazole 02506785 SAS ACDEFGV  
 Auro-Aripiprazole 02460084 ARO ACDEFGV  
 Mint-Aripiprazole 02483602 MNT ACDEFGV  
 pms-Aripiprazole 02466694 PMS ACDEFGV  
 Sandoz Aripiprazole 02473704 SDZ ACDEFGV

N05AX13 PALIPERIDONE  
 PALIPERIDONE PALMITATE

Liq IM 175 mg / 0.875 mL

Invega Trinza 02455943 JAN (SA)

Liq IM 263 mg / 1.315 mL

Invega Trinza 02455986 JAN (SA)

Liq IM 350 mg / 1.75 mL

Invega Trinza 02455994 JAN (SA)

Liq IM 525 mg / 2.625 mL

Invega Trinza 02456001 JAN (SA)

Sus IM 50 mg / 0.5 mL

Invega Sustenna 02354217 JAN (SA)

Sus IM 75 mg / 0.75 mL

Invega Sustenna 02354225 JAN (SA)

Sus IM 100 mg/mL

Invega Sustenna 02354233 JAN (SA)

Sus IM 150 mg / 1.5 mL

Invega Sustenna 02354241 JAN (SA)

N05AX16	BREXPIPIRAZOLE					
Tab	Orl	0.25 mg	Rexulti	02461749	OTS	ACDEFGV
Tab	Orl	0.50 mg	Rexulti	02461757	OTS	ACDEFGV
Tab	Orl	1 mg	Rexulti	02461765	OTS	ACDEFGV
Tab	Orl	2 mg	Rexulti	02461773	OTS	ACDEFGV
Tab	Orl	3 mg	Rexulti	02461781	OTS	ACDEFGV
Tab	Orl	4 mg	Rexulti	02461803	OTS	ACDEFGV

**N05B ANXIOLYTICS**

**N05BA BENZODIAZEPINE DERIVATIVES**

N05BA01	DIAZEPAM					
Liq	Inj	5 mg/mL	Diazepam	00399728	SDZ	ACDEFGV
Tab	Orl	2 mg	Diazepam	00405329	AAP	ACDEFGV
Tab	Orl	5 mg	Valium	00013285	SLP	ACDEFGV
			Diazepam	00362158	AAP	ACDEFGV
Tab	Orl	10 mg	Diazepam	00405337	AAP	ACDEFGV
N05BA02	CHLORDIAZEPOXIDE					
Cap	Orl	5 mg	Chlordiazepoxide	00522724	AAP	ACDEFGV
Cap	Orl	10 mg	Chlordiazepoxide	00522988	AAP	ACDEFGV
Cap	Orl	25 mg	Chlordiazepoxide	00522996	AAP	ACDEFGV
N05BA04	OXAZEPAM					
Tab	Orl	10 mg	Apo-Oxazepam	00402680	APX	ACDEFGV
Tab	Orl	15 mg	Apo-Oxazepam	00402745	APX	ACDEFGV
Tab	Orl	30 mg	Apo-Oxazepam	00402737	APX	ACDEFGV
N05BA05	CLORAZEPATE DIPOTASSIUM					
Cap	Orl	3.75 mg	Clorazepate	00860689	AAP	ACDEFGV
Cap	Orl	7.5 mg	Clorazepate	00860700	AAP	ACDEFGV

N05BA05	CLORAZEPATE DIPOTASSIUM						
Cap	Orl	15 mg	Clorazepate	00860697	AAP	ACDEFGV	
N05BA06	LORAZEPAM						
Liq	Inj	4 mg/mL	Lorazepam	02243278	SDZ	ACDEFVW	
Slit	Orl	0.5 mg	Ativan SL	02041456	PFI	ACDEFGVW	
			Lorazepam Sublingual	02410745	AAP	ACDEFGVW	
Slit	Orl	1 mg	Ativan SL	02041464	PFI	ACDEFGVW	
			Lorazepam Sublingual	02410753	AAP	ACDEFGVW	
Slit	Orl	2 mg	Ativan SL	02041472	PFI	ACDEFGVW	
			Lorazepam Sublingual	02410761	AAP	ACDEFGVW	
Tab	Orl	0.5 mg	Ativan	02041413	PFI	ACDEFGVW	
			Apo-Lorazepam	00655740	APX	ACDEFGVW	
			pms-Lorazepam	00728187	PMS	ACDEFGVW	
			Teva-Lorazepam	00711101	TEV	ACDEFGVW	
Tab	Orl	1 mg	Ativan	02041421	PFI	ACDEFGVW	
			Apo-Lorazepam	00655759	APX	ACDEFGVW	
			pms-Lorazepam	00728195	PMS	ACDEFGVW	
			Teva-Lorazepam	00637742	TEV	ACDEFGVW	
Tab	Orl	2 mg	Ativan	02041448	PFI	ACDEFGVW	
			Apo-Lorazepam	00655767	APX	ACDEFGVW	
			pms-Lorazepam	00728209	PMS	ACDEFGVW	
			Teva-Lorazepam	00637750	TEV	ACDEFGVW	
N05BA08	BROMAZEPAM						
Tab	Orl	3 mg	Apo-Bromazepam	02177161	APX	ACDEFGV	
			Teva-Bromazepam	02230584	TEV	ACDEFGV	
Tab	Orl	6 mg	Apo-Bromazepam	02177188	APX	ACDEFGV	
			Teva-Bromazepam	02230585	TEV	ACDEFGV	
N05BA09	CLOBAZAM						
Tab	Orl	10 mg	Apo-Clobazam	02244638	APX	ACDEFGV	
			Teva-Clobazam	02238334	TEV	ACDEFGV	
N05BA12	ALPRAZOLAM						

N05BA12 ALPRAZOLAM

Tab Orl 0.25 mg

Xanax 00548359 UJC ACDEFGV  
 Apo-Alpraz 00865397 APX ACDEFGV  
 Teva-Alprazolam 01913484 TEV ACDEFGV

Tab Orl 0.5 mg

Xanax 00548367 UJC ACDEFGV  
 Apo-Alpraz 00865400 APX ACDEFGV  
 Teva-Alprazolam 01913492 TEV ACDEFGV

**N05BB DIPHENYLMETHANE DERIVATIVES**

N05BB01 HYDROXYZINE

Cap Orl 10 mg

Hydroxyzine 00646059 AAP ACDEFGVW  
 Novo-Hydroxyzine 00738824 TEV ACDEFGVW

Cap Orl 25 mg

Hydroxyzine 00646024 AAP ACDEFGVW  
 Novo-Hydroxyzine 00738832 TEV ACDEFGVW

Cap Orl 50 mg

Hydroxyzine 00646016 AAP ACDEFGVW  
 Novo-Hydroxyzine 00738840 TEV ACDEFGVW

Syr Orl 2 mg/mL

Atarax 00024694 SLP ACDEFGVW

**N05BE AZASPIRODECANEDIONE DERIVATIVES**

N05BE01 BUSPIRONE

Tab Orl 10 mg

Apo-Buspirone 02211076 APX ACDEFGV  
 Auro-Buspirone 02500213 ARO ACDEFGV  
 Buspirone 02447851 SAS ACDEFGV  
 Jamp Buspirone 02509911 JPC ACDEFGV  
 Mint-Buspirone 02519054 MNT ACDEFGV  
 pms-Buspirone 02230942 PMS ACDEFGV  
 Teva-Buspirone 02231492 TEV ACDEFGV

**N05C HYPNOTICS AND SEDATIVES**

**N05CD BENZODIAZEPINE DERIVATIVES**

N05CD01 FLURAZEPAM

Cap Orl 15 mg

Flurazepam 00521698 AAP ACDEFGV

Cap Orl 30 mg

Flurazepam 00521701 AAP ACDEFGV

N05CD02 NITRAZEPAM

Tab Orl 5 mg

Mogadon 00511528 AAP ACDEFGV

N05CD02	NITRAZEPAM							
Tab	Orl	10 mg		Mogadon	00511536	AAP	ACDEFGV	
N05CD05	TRIAZOLAM							
Tab	Orl	0.25 mg		Triazolam	00808571	AAP	ACDEFGV	
N05CD07	TEMAZEPAM							
Cap	Orl	15 mg		Restoril	00604453	AAP	ACDEFGV	
Cap	Orl	30 mg		Restoril	00604461	AAP	ACDEFGV	
N05CD08	MIDAZOLAM							
Liq	Inj	1 mg/mL		Midazolam	02240285	SDZ	ACDEFGVW	
Liq	Inj	5 mg/mL		Midazolam	02240286	SDZ	ACDEFGVW	

**N05CF BENZODIAZEPINE RELATED DRUGS**

N05CF01	ZOPICLONE							
Tab	Orl	3.75 mg		pms-Zopiclone	02458543	PMS	ACDEFGV	
Tab	Orl	5 mg		Apo-Zopiclone	02245077	APX	ACDEFGV	
				Jamp-Zopiclone	02406969	JPC	ACDEFGV	
				M-Zopiclone	02467941	MRA	ACDEFGV	
				Mar-Zopiclone	02386771	MAR	ACDEFGV	
				Mint-Zopiclone	02391716	MNT	ACDEFGV	
				NRA-Zopiclone	02477378	NRA	ACDEFGV	
				pms-Zopiclone	02243426	PMS	ACDEFGV	
				ratio-Zopiclone	02246534	TEV	ACDEFGV	
				Zopiclone	02344122	SAS	ACDEFGV	
				Zopiclone	02385821	SIV	ACDEFGV	

N05CF01 ZOPICLONE

Tab Orl 7.5 mg

Imovane 01926799 SAV ACDEFGV  
Apo-Zopiclone 02218313 APX ACDEFGV  
Jamp-Zopiclone 02406977 JPC ACDEFGV  
M-Zopiclone 02467968 MRA ACDEFGV  
Mar-Zopiclone 02386798 MAR ACDEFGV  
Mint-Zopiclone 02391724 MNT ACDEFGV  
NRA-Zopiclone 02477386 NRA ACDEFGV  
pms-Zopiclone 02240606 PMS ACDEFGV  
ratio-Zopiclone 02242481 TEV ACDEFGV  
Zopiclone 02282445 SAS ACDEFGV  
Zopiclone 02385848 SIV ACDEFGV

**N06 PSYCHOANALEPTICS**

**N06A ANTIDEPRESSANTS**

**N06AA NON-SELECTIVE MONOAMINE REUPTAKE INHIBITORS**

N06AA01 DESIPRAMINE

Tab Orl 10 mg

Desipramine 02216248 AAP ACDEFGV

Tab Orl 25 mg

Desipramine 02216256 AAP ACDEFGV

Tab Orl 50 mg

Desipramine 02216264 AAP ACDEFGV

Tab Orl 75 mg

Desipramine 02216272 AAP ACDEFGV

Tab Orl 100 mg

Desipramine 02216280 AAP ACDEFGV

N06AA02 IMIPRAMINE

Tab Orl 10 mg

Imipramine 00360201 AAP ACDEFGV

Tab Orl 25 mg

Imipramine 00312797 AAP ACDEFGV

Tab Orl 50 mg

Imipramine 00326852 AAP ACDEFGV

Tab Orl 75 mg

Imipramine 00644579 AAP ACDEFGV

N06AA04 CLOMIPRAMINE

Cap Orl 25 mg

Taro-Clomipramine 02497506 TAR ACDEFGV

Cap Orl 50 mg

Taro-Clomipramine 02497514 TAR ACDEFGV

Tab Orl 10 mg

Anafranil 00330566 APX ACDEFGV



N06AA04	CLOMIPRAMINE					
Tab	Orl	25 mg	Anafranil	00324019	APX	ACDEFGV
Tab	Orl	50 mg	Anafranil	00402591	APX	ACDEFGV
N06AA06	TRIMIPRAMINE					
Tab	Orl	12.5 mg	Trimipramine	00740799	AAP	ACDEFGV
Tab	Orl	25 mg	Trimipramine	00740802	AAP	ACDEFGV
Tab	Orl	50 mg	Trimipramine	00740810	AAP	ACDEFGV
Tab	Orl	75 mg	Trimipramine	02070987	AAP	ACDEFGV
Tab	Orl	100 mg	Trimipramine	00740829	AAP	ACDEFGV
N06AA09	AMITRIPTYLINE					
Tab	Orl	10 mg	Elavil	00335053	AAP	ACDEFGV
			Amitriptyline	00370991	PDL	ACDEFGV
			Apo-Amitriptyline	02403137	APX	ACDEFGV
Tab	Orl	25 mg	Elavil	00335061	AAP	ACDEFGV
			Amitriptyline	00371009	PDL	ACDEFGV
			Apo-Amitriptyline	02403145	APX	ACDEFGV
Tab	Orl	50 mg	Elavil	00335088	AAP	ACDEFGV
			Apo-Amitriptyline	02403153	APX	ACDEFGV
Tab	Orl	75 mg	Elavil	00754129	AAP	ACDEFGV
			Apo-Amitriptyline	02403161	APX	ACDEFGV
N06AA10	NORTRIPTYLINE					
Cap	Orl	10 mg	Aventyl	00015229	AAP	ACDEFGV
Cap	Orl	25 mg	Aventyl	00015237	AAP	ACDEFGV
N06AA12	DOXEPIN					
Cap	Orl	10 mg	Sinequan	00024325	AAP	ACDEFGV
Cap	Orl	25 mg	Sinequan	00024333	AAP	ACDEFGV
Cap	Orl	50 mg	Sinequan	00024341	AAP	ACDEFGV

**N06AB SELECTIVE SEROTONIN REUPTAKE INHIBITORS (SSRI'S)**

**N06AB03 FLUOXETINE**

Cap Orl 10 mg

Prozac 02018985 LIL ACDEFGV  
 Apo-Fluoxetine 02216353 APX ACDEFGV  
 Auro-Fluoxetine 02385627 ARO ACDEFGV  
 Fluoxetine 02393441 AHI ACDEFGV  
 Fluoxetine 02286068 SAS ACDEFGV  
 Fluoxetine 02374447 SIV ACDEFGV  
 Jamp-Fluoxetine 02401894 JPC ACDEFGV  
 M-Fluoxetine 02529432 MRA ACDEFGV  
 NRA-Fluoxetine 02503875 NRA ACDEFGV  
 pms-Fluoxetine 02177579 PMS ACDEFGV  
 Teva-Fluoxetine 02216582 TEV ACDEFGV

Cap Orl 20 mg

Prozac 00636622 LIL ACDEFGV  
 Apo-Fluoxetine 02216361 APX ACDEFGV  
 Auro-Fluoxetine 02385635 ARO ACDEFGV  
 Fluoxetine 02383241 AHI ACDEFGV  
 Fluoxetine 02286076 SAS ACDEFGV  
 Fluoxetine 02374455 SIV ACDEFGV  
 Jamp-Fluoxetine 02386402 JPC ACDEFGV  
 M-Fluoxetine 02529440 MRA ACDEFGV  
 NRA-Fluoxetine 02503883 NRA ACDEFGV  
 pms-Fluoxetine 02177587 PMS ACDEFGV  
 Teva-Fluoxetine 02216590 TEV ACDEFGV

Cap Orl 40 mg

pms-Fluoxetine 02464640 PMS ACDEFGV

Cap Orl 60 mg

pms-Fluoxetine 02464659 PMS ACDEFGV

Liq Orl 20 mg / 5 mL

Apo-Fluoxetine 02231328 APX (SA)  
 Odan-Fluoxetine 02459361 ODN (SA)

**N06AB04 CITALOPRAM**

N06AB04 CITALOPRAM

Tab Orl 10 mg

Citalopram 02430517 JPC ACDEFGV  
 Citalopram 02445719 SAS ACDEFGV  
 Citalopram 02387948 SIV ACDEFGV  
 Citalopram-10 02325047 PDL ACDEFGV  
 Mar-Citalopram 02371871 MAR ACDEFGV  
 Mint-Citalopram 02429691 MNT ACDEFGV  
 Nat-Citalopram 02409003 NAT ACDEFGV  
 pms-Citalopram 02270609 PMS ACDEFGV  
 Teva-Citalopram 02312336 TEV ACDEFGV

Tab Orl 20 mg

Celexa 02239607 VLH ACDEFGV  
 Apo-Citalopram 02246056 APX ACDEFGV  
 Auro-Citalopram 02275562 ARO ACDEFGV  
 CCP-Citalopram 02459914 CCM ACDEFGV  
 Citalopram 02430541 JPC ACDEFGV  
 Citalopram 02353660 SAS ACDEFGV  
 Citalopram 02387956 SIV ACDEFGV  
 Citalopram-20 02257513 PDL ACDEFGV  
 Mar-Citalopram 02371898 MAR ACDEFGV  
 Mint-Citalopram 02429705 MNT ACDEFGV  
 Nat-Citalopram 02409011 NAT ACDEFGV  
 pms-Citalopram 02248010 PMS ACDEFGV  
 Septa-Citalopram 02355272 SPT ACDEFGV  
 Teva-Citalopram 02293218 TEV ACDEFGV

Tab Orl 30 mg

CTP 30 02296152 SNV ACDEFGV

N06AB04 CITALOPRAM

Tab Orl 40 mg

Celexa	02239608	VLH	ACDEFGV
Apo-Citalopram	02246057	APX	ACDEFGV
Auro-Citalopram	02275570	ARO	ACDEFGV
CCP-Citalopram	02459922	CCM	ACDEFGV
Citalopram	02430568	JPC	ACDEFGV
Citalopram	02353679	SAS	ACDEFGV
Citalopram	02387964	SIV	ACDEFGV
Citalopram-40	02257521	PDL	ACDEFGV
Mar-Citalopram	02371901	MAR	ACDEFGV
Mint-Citalopram	02429713	MNT	ACDEFGV
Nat-Citalopram	02409038	NAT	ACDEFGV
pms-Citalopram	02248011	PMS	ACDEFGV
Septa-Citalopram	02355280	SPT	ACDEFGV
Teva-Citalopram	02293226	TEV	ACDEFGV

N06AB05 PAROXETINE

Tab Orl 10 mg

Paxil	02027887	GSK	ACDEFGV
Apo-Paroxetine	02240907	APX	ACDEFGV
Auro-Paroxetine	02383276	ARO	ACDEFGV
Jamp Paroxetine Tablets	02507773	JPC	ACDEFGV
Jamp-Paroxetine	02368862	JPC	ACDEFGV
M-Paroxetine	02467402	MRA	ACDEFGV
Mar-Paroxetine	02411946	MAR	ACDEFGV
Mint-Paroxetine	02421372	MNT	ACDEFGV
NRA-Paroxetine	02479753	NRA	ACDEFGV
Paroxetine	02282844	SAS	ACDEFGV
Paroxetine	02388227	SIV	ACDEFGV
pms-Paroxetine	02247750	PMS	ACDEFGV
Teva-Paroxetine	02248556	TEV	ACDEFGV

N06AB05 PAROXETINE

Tab Orl 20 mg

Paxil	01940481	GSK	ACDEFGV
Apo-Paroxetine	02240908	APX	ACDEFGV
Auro-Paroxetine	02383284	ARO	ACDEFGV
Jamp Paroxetine Tablets	02507781	JPC	ACDEFGV
Jamp-Paroxetine	02368870	JPC	ACDEFGV
M-Paroxetine	02467410	MRA	ACDEFGV
Mar-Paroxetine	02411954	MAR	ACDEFGV
Mint-Paroxetine	02421380	MNT	ACDEFGV
NRA-Paroxetine	02479761	NRA	ACDEFGV
Paroxetine	02248914	PDL	ACDEFGV
Paroxetine	02282852	SAS	ACDEFGV
Paroxetine	02388235	SIV	ACDEFGV
pms-Paroxetine	02247751	PMS	ACDEFGV
Teva-Paroxetine	02248557	TEV	ACDEFGV

Tab Orl 30 mg

Paxil	01940473	GSK	ACDEFGV
Apo-Paroxetine	02240909	APX	ACDEFGV
Auro-Paroxetine	02383292	ARO	ACDEFGV
Jamp Paroxetine Tablets	02507803	JPC	ACDEFGV
Jamp-Paroxetine	02368889	JPC	ACDEFGV
M-Paroxetine	02467429	MRA	ACDEFGV
Mar-Paroxetine	02411962	MAR	ACDEFGV
Mint-Paroxetine	02421399	MNT	ACDEFGV
NRA-Paroxetine	02479788	NRA	ACDEFGV
Paroxetine	02248915	PDL	ACDEFGV
Paroxetine	02282860	SAS	ACDEFGV
Paroxetine	02388243	SIV	ACDEFGV
pms-Paroxetine	02247752	PMS	ACDEFGV

N06AB06 SERTRALINE

N06AB06 SERTRALINE

Cap Orl 25 mg

Zoloft	02132702	UJC	ACDEFGV
Apo-Sertraline	02238280	APX	ACDEFGV
Auro-Sertraline	02390906	ARO	ACDEFGV
M-Sertraline	02530937	MRA	ACDEFGV
Mar-Sertraline	02399415	MAR	ACDEFGV
Mint-Sertraline	02402378	MNT	ACDEFGV
NRA-Sertraline	02488434	NRA	ACDEFGV
pms-Sertraline	02244838	PMS	ACDEFGV
Sertraline	02469626	JPC	ACDEFGV
Sertraline	02353520	SAS	ACDEFGV
Sertraline	02386070	SIV	ACDEFGV
Teva-Sertraline	02240485	TEV	ACDEFGV

Cap Orl 50 mg

Zoloft	01962817	UJC	ACDEFGV
Apo-Sertraline	02238281	APX	ACDEFGV
Auro-Sertraline	02390914	ARO	ACDEFGV
M-Sertraline	02530945	MRA	ACDEFGV
Mar-Sertraline	02399423	MAR	ACDEFGV
Mint-Sertraline	02402394	MNT	ACDEFGV
NRA-Sertraline	02488442	NRA	ACDEFGV
pms-Sertraline	02244839	PMS	ACDEFGV
Sertraline	02469634	JPC	ACDEFGV
Sertraline	02353539	SAS	ACDEFGV
Sertraline	02386089	SIV	ACDEFGV
Teva-Sertraline	02240484	TEV	ACDEFGV

Cap Orl 100 mg

Zoloft	01962779	UJC	ACDEFGV
Apo-Sertraline	02238282	APX	ACDEFGV
Auro-Sertraline	02390922	ARO	ACDEFGV
M-Sertraline	02530953	MRA	ACDEFGV
Mar-Sertraline	02399431	MAR	ACDEFGV
Mint-Sertraline	02402408	MNT	ACDEFGV
NRA-Sertraline	02488450	NRA	ACDEFGV
pms-Sertraline	02244840	PMS	ACDEFGV
Sertraline	02469642	JPC	ACDEFGV
Sertraline	02353547	SAS	ACDEFGV
Sertraline	02386097	SIV	ACDEFGV
Teva-Sertraline	02240481	TEV	ACDEFGV

N06AB08 FLUVOXAMINE

Tab Orl 50 mg

Luvox 01919342 BGP ACDEFGV  
 Act Fluvoxamine 02255529 TEV ACDEFGV  
 Apo-Fluvoxamine 02231329 APX ACDEFGV

Tab Orl 100 mg

Luvox 01919369 BGP ACDEFGV  
 Act Fluvoxamine 02255537 TEV ACDEFGV  
 Apo-Fluvoxamine 02231330 APX ACDEFGV

N06AB10 ESCITALOPRAM

Tab Orl 10 mg

Cipralext 02263238 VLH ACDEFGV  
 Ach-Escitalopram 02434652 AHI ACDEFGV  
 Apo-Escitalopram 02295016 APX ACDEFGV  
 Auro-Escitalopram 02397358 ARO ACDEFGV  
 Escitalopram 02430118 SAS ACDEFGV  
 Escitalopram 02429039 SIV ACDEFGV  
 Jamp-Escitalopram 02429780 JPC ACDEFGV  
 M-Escitalopram 02471418 MRA ACDEFGV  
 Mar-Escitalopram 02423480 MAR ACDEFGV  
 Mint-Escitalopram 02407418 MNT ACDEFGV  
 Mylan-Escitalopram 02309467 MYL ACDEFGV  
 Nat-Escitalopram 02440296 NAT ACDEFGV  
 NRA-Escitalopram 02476851 NRA ACDEFGV  
 pms-Escitalopram 02469243 PMS ACDEFGV  
 Sandoz Escitalopram 02364077 SDZ ACDEFGV  
 Taro-Escitalopram 02385481 SUN ACDEFGV  
 Teva-Escitalopram 02318180 TEV ACDEFGV

Tab Orl 15 mg

Kye-Escitalopram 02512653 KYE ACDEFGV

N06AB10 ESCITALOPRAM

Tab Orl 20 mg

Ciprallex	02263254	VLH	ACDEFGV
Ach-Escitalopram	02434660	AHI	ACDEFGV
Apo-Escitalopram	02295024	APX	ACDEFGV
Auro-Escitalopram	02397374	ARO	ACDEFGV
Escitalopram	02430126	SAS	ACDEFGV
Escitalopram	02429047	SIV	ACDEFGV
Jamp-Escitalopram	02429799	JPC	ACDEFGV
M-Escitalopram	02471426	MRA	ACDEFGV
Mar-Escitalopram	02423502	MAR	ACDEFGV
Mint-Escitalopram	02407434	MNT	ACDEFGV
Mylan-Escitalopram	02309475	MYL	ACDEFGV
Nat-Escitalopram	02440318	NAT	ACDEFGV
NRA-Escitalopram	02476878	NRA	ACDEFGV
pms-Escitalopram	02469251	PMS	ACDEFGV
Sandoz Escitalopram	02364085	SDZ	ACDEFGV
Taro-Escitalopram	02385503	SUN	ACDEFGV
Teva-Escitalopram	02318202	TEV	ACDEFGV

**N06AF MONOAMINE OXIDASE INHIBITORS, NON-SELECTIVE**

N06AF03 PHENELZINE

Tab Orl 15 mg

Nardil 00476552 SLP ACDEFGV

N06AF04 TRANYLCPROMINE

Tab Orl 10 mg

Parnate 01919598 GSK ACDEFGV

**N06AG MONOAMINE OXIDASE TYPE A INHIBITORS**

N06AG02 MOCLOBEMIDE

Tab Orl 100 mg

Moclobemide 02232148 AAP ACDEFGV

Tab Orl 150 mg

Manerix 00899356 BSL ACDEFGV

Moclobemide 02232150 AAP ACDEFGV

Tab Orl 300 mg

Manerix 02166747 BSL ACDEFGV

Moclobemide 02240456 AAP ACDEFGV

**N06AX OTHER ANTIDEPRESSANTS**

N06AX02 TRYPTOPHAN

Cap Orl 500 mg

Tryptan 00718149 BSL ACDEFGV

Apo-Tryptophan 02248540 APX ACDEFGV

Teva-Tryptophan 02240334 TEV ACDEFGV



N06AX02 TRYPTOPHAN

Tab	Orl	250 mg		Tryptan	02239326	BSL	ACDEFGV
Tab	Orl	500 mg		Tryptan	02029456	BSL	ACDEFGV
				Apo-Tryptophan	02248538	APX	ACDEFGV
				Teva-Tryptophan	02240333	TEV	ACDEFGV
Tab	Orl	750 mg		Tryptan	02239327	BSL	ACDEFGV
				Apo-Tryptophan	02458721	APX	ACDEFGV
Tab	Orl	1 000 mg		Tryptan	00654531	BSL	ACDEFGV
				Apo-Tryptophan	02248539	APX	ACDEFGV
				Teva-Tryptophan	02237250	TEV	ACDEFGV

N06AX05 TRAZODONE

Tab	Orl	50 mg		Apo-Trazodone	02147637	APX	ACDEFGV
				Jamp Trazodone	02442809	JPC	ACDEFGV
				pms-Trazodone	01937227	PMS	ACDEFGV
				Teva-Trazodone	02144263	TEV	ACDEFGV
				Trazodone	02348772	SAS	ACDEFGV
Tab	Orl	100 mg		Apo-Trazodone	02147645	APX	ACDEFGV
				Jamp Trazodone	02442817	JPC	ACDEFGV
				pms-Trazodone	01937235	PMS	ACDEFGV
				Teva-Trazodone	02144271	TEV	ACDEFGV
				Trazodone	02348780	SAS	ACDEFGV
Tab	Orl	150 mg		Apo-Trazodone D	02147653	APX	ACDEFGV
				Jamp Trazodone	02442825	JPC	ACDEFGV
				Teva-Trazodone	02144298	TEV	ACDEFGV
				Trazodone	02348799	SAS	ACDEFGV

N06AX11 MIRTAZAPINE

ODT	Orl	15 mg		Remeron RD	02248542	ORG	(SA)
				Auro-Mirtazapine OD	02299801	ARO	(SA)
ODT	Orl	30 mg		Remeron RD	02248543	ORG	(SA)
				Auro-Mirtazapine OD	02299828	ARO	(SA)
ODT	Orl	45 mg		Remeron RD	02248544	ORG	(SA)
				Auro-Mirtazapine OD	02299836	ARO	(SA)

N06AX11 MIRTAZAPINE

Tab Orl 15 mg

Apo-Mirtazapine 02286610 APX ACDEFGV  
 Auro-Mirtazapine 02411695 ARO ACDEFGV  
 Mirtazapine 02496666 SIV ACDEFGV  
 Mylan-Mirtazapine 02256096 MYL ACDEFGV  
 pms-Mirtazapine 02273942 PMS ACDEFGV  
 Sandoz Mirtazapine 02250594 SDZ ACDEFGV

Tab Orl 30 mg

Remeron 02243910 ORG ACDEFGV  
 Apo-Mirtazapine 02286629 APX ACDEFGV  
 Auro-Mirtazapine 02411709 ARO ACDEFGV  
 Mirtazapine 02370689 SAS ACDEFGV  
 Mirtazapine 02496674 SIV ACDEFGV  
 Mylan-Mirtazapine 02256118 MYL ACDEFGV  
 pms-Mirtazapine 02248762 PMS ACDEFGV  
 Sandoz Mirtazapine 02250608 SDZ ACDEFGV  
 Teva-Mirtazapine 02259354 TEV ACDEFGV

Tab Orl 45 mg

Apo-Mirtazapine 02286637 APX ACDEFGV  
 Auro-Mirtazapine 02411717 ARO ACDEFGV  
 Mirtazapine 02496682 SIV ACDEFGV

N06AX12 BUPROPION

ERT Orl 150 mg

Wellbutrin XL 02275090 BSL ACDEFGV  
 Taro-Bupropion XL 02475804 SUN ACDEFGV  
 Teva-Bupropion XL 02439654 TEV ACDEFGV

ERT Orl 150 mg

Zyban 02238441 BSL (SA)

ERT Orl 300 mg

Wellbutrin XL 02275104 BSL ACDEFGV  
 Taro-Bupropion XL 02475812 SUN ACDEFGV  
 Teva-Bupropion XL 02439662 TEV ACDEFGV

SRT Orl 100 mg

Odan Bupropion SR 02275074 ODN ACDEFGV

SRT Orl 150 mg

Odan Bupropion SR 02275082 ODN ACDEFGV

N06AX16 VENLAFAXINE

N06AX16 VENLAFAXINE

SRC Orl 37.5 mg

Effexor XR	02237279	BGP	ACDEFGV
Act Venlafaxine XR	02304317	TEV	ACDEFGV
Apo-Venlafaxine XR	02331683	APX	ACDEFGV
Auro-Venlafaxine XR	02452839	ARO	ACDEFGV
M-Venlafaxine XR	02471280	MRA	ACDEFGV
pms-Venlafaxine XR	02278545	PMS	ACDEFGV
pmsc-Venlafaxine XR	02521466	PMS	ACDEFGV
Sandoz Venlafaxine XR	02310317	SDZ	ACDEFGV
Taro-Venlafaxine XR	02380072	SUN	ACDEFGV
Teva-Venlafaxine XR	02275023	TEV	ACDEFGV
Venlafaxine XR	02516535	JPC	ACDEFGV
Venlafaxine XR	02339242	PDL	ACDEFGV
Venlafaxine XR	02354713	SAS	ACDEFGV
Venlafaxine XR	02385929	SIV	ACDEFGV

SRC Orl 75 mg

Effexor XR	02237280	BGP	ACDEFGV
Act Venlafaxine XR	02304325	TEV	ACDEFGV
Apo-Venlafaxine XR	02331691	APX	ACDEFGV
Auro-Venlafaxine XR	02452847	ARO	ACDEFGV
M-Venlafaxine XR	02471299	MRA	ACDEFGV
pms-Venlafaxine XR	02278553	PMS	ACDEFGV
pmsc-Venlafaxine XR	02521482	PMS	ACDEFGV
Sandoz Venlafaxine XR	02310325	SDZ	ACDEFGV
Taro-Venlafaxine XR	02380080	SUN	ACDEFGV
Teva-Venlafaxine XR	02275031	TEV	ACDEFGV
Venlafaxine XR	02516543	JPC	ACDEFGV
Venlafaxine XR	02339250	PDL	ACDEFGV
Venlafaxine XR	02354721	SAS	ACDEFGV
Venlafaxine XR	02385937	SIV	ACDEFGV

N06AX16 VENLAFAXINE

SRC Orl 150 mg

Effexor XR	02237282	BGP	ACDEFGV
Act Venlafaxine XR	02304333	TEV	ACDEFGV
Apo-Venlafaxine XR	02331705	APX	ACDEFGV
Auro-Venlafaxine XR	02452855	ARO	ACDEFGV
M-Venlafaxine XR	02471302	MRA	ACDEFGV
pms-Venlafaxine XR	02278561	PMS	ACDEFGV
pmsc-Venlafaxine XR	02521474	PMS	ACDEFGV
Sandoz Venlafaxine XR	02310333	SDZ	ACDEFGV
Taro-Venlafaxine XR	02380099	SUN	ACDEFGV
Teva-Venlafaxine XR	02275058	TEV	ACDEFGV
Venlafaxine XR	02516551	JPC	ACDEFGV
Venlafaxine XR	02339269	PDL	ACDEFGV
Venlafaxine XR	02354748	SAS	ACDEFGV
Venlafaxine XR	02385945	SIV	ACDEFGV

N06AX21 DULOXETINE

CDR Orl 30 mg

Cymbalta	02301482	LIL	(SA)
Apo-Duloxetine	02440423	APX	(SA)
Auro-Duloxetine	02436647	ARO	(SA)
Duloxetine	02490889	SAS	(SA)
Duloxetine	02453630	SIV	(SA)
Jamp-Duloxetine	02451913	JPC	(SA)
M-Duloxetine	02473208	MRA	(SA)
Mar-Duloxetine	02446081	MAR	(SA)
Mint-Duloxetine	02438984	MNT	(SA)
NRA-Duloxetine	02482126	NRA	(SA)
pms-Duloxetine	02429446	PMS	(SA)
Sandoz Duloxetine	02439948	SDZ	(SA)
Teva-Duloxetine	02456753	TEV	(SA)

N06AX21 DULOXETINE

CDR Orl 60 mg

Cymbalta 02301490 LIL (SA)  
 Apo-Duloxetine 02440431 APX (SA)  
 Auro-Duloxetine 02436655 ARO (SA)  
 Duloxetine 02490897 SAS (SA)  
 Duloxetine 02453649 SIV (SA)  
 Jamp-Duloxetine 02451921 JPC (SA)  
 M-Duloxetine 02473216 MRA (SA)  
 Mar-Duloxetine 02446103 MAR (SA)  
 Mint-Duloxetine 02438992 MNT (SA)  
 NRA-Duloxetine 02482134 NRA (SA)  
 pms-Duloxetine 02429454 PMS (SA)  
 Sandoz Duloxetine 02439956 SDZ (SA)  
 Teva-Duloxetine 02456761 TEV (SA)

N06AX26 VORTIOXETINE

Tab Orl 5 mg

Trintellix 02432919 VLH ACDEFGV

Tab Orl 10 mg

Trintellix 02432927 VLH ACDEFGV

Tab Orl 20 mg

Trintellix 02432943 VLH ACDEFGV

**N06B PSYCHOSTIMULANTS, AGENTS USED FOR ADHD AND NOOTROPICS**

**N06BA CENTRALLY ACTING SYMPATHOMIMETICS**

N06BA01 AMPHETAMINE

MIXED SALTS AMPHETAMINE

ERC Orl 5 mg

Adderall XR 02248808 TAK ACDEFG  
 Act Amphetamine XR 02439239 TEV ACDEFG  
 Apo-Amphetamine XR 02445492 APX ACDEFG  
 pms-Amphetamines XR 02440369 PMS ACDEFG  
 Sandoz Amphetamine XR 02457288 SDZ ACDEFG

ERC Orl 10 mg

Adderall XR 02248809 TAK ACDEFG  
 Act Amphetamine XR 02439247 TEV ACDEFG  
 Apo-Amphetamine XR 02445506 APX ACDEFG  
 pms-Amphetamines XR 02440377 PMS ACDEFG  
 Sandoz Amphetamine XR 02457296 SDZ ACDEFG

N06BA01	AMPHETAMINE						
	MIXED SALTS AMPHETAMINE						
ERC	Orl	15 mg	Adderall XR	02248810	TAK	ACDEFG	
			Act Amphetamine XR	02439255	TEV	ACDEFG	
			Apo-Amphetamine XR	02445514	APX	ACDEFG	
			pms-Amphetamines XR	02440385	PMS	ACDEFG	
			Sandoz Amphetamine XR	02457318	SDZ	ACDEFG	
ERC	Orl	20 mg	Adderall XR	02248811	TAK	ACDEFG	
			Act Amphetamine XR	02439263	TEV	ACDEFG	
			Apo-Amphetamine XR	02445522	APX	ACDEFG	
			Sandoz Amphetamine XR	02457326	SDZ	ACDEFG	
ERC	Orl	25 mg	Adderall XR	02248812	TAK	ACDEFG	
			Act Amphetamine XR	02439271	TEV	ACDEFG	
			Apo-Amphetamine XR	02445530	APX	ACDEFG	
			pms-Amphetamines XR	02440407	PMS	ACDEFG	
			Sandoz Amphetamine XR	02457334	SDZ	ACDEFG	
ERC	Orl	30 mg	Adderall XR	02248813	TAK	ACDEFG	
			Act Amphetamine XR	02439298	TEV	ACDEFG	
			Apo-Amphetamine XR	02445549	APX	ACDEFG	
			Sandoz Amphetamine XR	02457342	SDZ	ACDEFG	
N06BA02	DEXAMPHETAMINE						
SRC	Orl	10 mg	Dexedrine	01924559	PAL	ACDEFG	
			Act-Dextroamphetamine SR	02448319	TEV	ACDEFG	
SRC	Orl	15 mg	Dexedrine	01924567	PAL	ACDEFG	
			Act-Dextroamphetamine SR	02448327	TEV	ACDEFG	
Tab	Orl	5 mg	Dexedrine	01924516	PAL	ACDEFG	
			Dextroamphetamine	02443236	AAP	ACDEFG	
N06BA04	METHYLPHENIDATE						
ERC	Orl	10 mg	Biphentin	02277166	ELV	(SA)	
ERC	Orl	15 mg	Biphentin	02277131	ELV	(SA)	
ERC	Orl	20 mg	Biphentin	02277158	ELV	(SA)	
ERC	Orl	30 mg	Biphentin	02277174	ELV	(SA)	

N06BA04 METHYLPHENIDATE

ERC	Orl	40 mg	Biphentin	02277182	ELV	(SA)
ERC	Orl	50 mg	Biphentin	02277190	ELV	(SA)
ERC	Orl	60 mg	Biphentin	02277204	ELV	(SA)
ERC	Orl	80 mg	Biphentin	02277212	ELV	(SA)
ERT	Orl	18 mg	Concerta ER	02247732	JAN	ACDEFGV
			Act Methylphenidate ER	02441934	TEV	ACDEFGV
			Apo-Methylphenidate ER	02452731	APX	ACDEFGV
ERT	Orl	27 mg	Concerta ER	02250241	JAN	ACDEFGV
			Act Methylphenidate ER	02441942	TEV	ACDEFGV
			Apo-Methylphenidate ER	02452758	APX	ACDEFGV
ERT	Orl	36 mg	Concerta ER	02247733	JAN	ACDEFGV
			Act Methylphenidate ER	02441950	TEV	ACDEFGV
			Apo-Methylphenidate ER	02452766	APX	ACDEFGV
ERT	Orl	54 mg	Concerta ER	02247734	JAN	ACDEFGV
			Act Methylphenidate ER	02441969	TEV	ACDEFGV
			Apo-Methylphenidate ER	02330377	APX	ACDEFGV
SRT	Orl	20 mg	Apo-Methylphenidate SR	02266687	APX	ACDEFGV
Tab	Orl	5 mg	Apo-Methylphenidate	02273950	APX	ACDEFGV
			pms-Methylphenidate	02234749	PMS	ACDEFGV
Tab	Orl	10 mg	Apo-Methylphenidate	02249324	APX	ACDEFGV
			pms-Methylphenidate	00584991	PMS	ACDEFGV
Tab	Orl	20 mg	Apo-Methylphenidate	02249332	APX	ACDEFGV
			pms-Methylphenidate	00585009	PMS	ACDEFGV

N06BA07 MODAFINIL

N06BA07 MODAFINIL

Tab Orl 100 mg

Alertec 02239665 TEV ACDEFGV  
 Apo-Modafinil 02285398 APX ACDEFGV  
 Auro-Modafinil 02430487 ARO ACDEFGV  
 Jamp Modafinil 02503727 JPC ACDEFGV  
 Mar-Modafinil 02432560 MAR ACDEFGV  
 Modafinil 02530244 SAS ACDEFGV  
 Teva-Modafinil 02420260 TEV ACDEFGV

N06BA09 ATOMOXETINE

Cap Orl 10 mg

Strattera 02262800 LIL ACDEFG  
 Apo-Atomoxetine 02318024 APX ACDEFG  
 Atomoxetine 02467747 SAS ACDEFG  
 Atomoxetine 02445883 SIV ACDEFG  
 Auro-Atomoxetine 02471485 ARO ACDEFG  
 Jamp Atomoxetine 02506807 JPC ACDEFG  
 pms-Atomoxetine 02381028 PMS ACDEFG  
 Sandoz Atomoxetine 02386410 SDZ ACDEFG  
 Teva-Atomoxetine 02314541 TEV ACDEFG

Cap Orl 18 mg

Strattera 02262819 LIL ACDEFG  
 Apo-Atomoxetine 02318032 APX ACDEFG  
 Atomoxetine 02467755 SAS ACDEFG  
 Atomoxetine 02445905 SIV ACDEFG  
 Auro-Atomoxetine 02471493 ARO ACDEFG  
 Jamp Atomoxetine 02506815 JPC ACDEFG  
 pms-Atomoxetine 02381036 PMS ACDEFG  
 Sandoz Atomoxetine 02386429 SDZ ACDEFG  
 Teva-Atomoxetine 02314568 TEV ACDEFG

Cap Orl 25 mg

Strattera 02262827 LIL ACDEFG  
 Apo-Atomoxetine 02318040 APX ACDEFG  
 Atomoxetine 02467763 SAS ACDEFG  
 Atomoxetine 02445913 SIV ACDEFG  
 Auro-Atomoxetine 02471507 ARO ACDEFG  
 Jamp Atomoxetine 02506823 JPC ACDEFG  
 pms-Atomoxetine 02381044 PMS ACDEFG  
 Sandoz Atomoxetine 02386437 SDZ ACDEFG  
 Teva-Atomoxetine 02314576 TEV ACDEFG



N06BA09 ATOMOXETINE

Cap Orl 40 mg

Strattera (Disc/non disp May 29/24) 02262835 LIL ACDEFG  
 Apo-Atomoxetine 02318059 APX ACDEFG  
 Atomoxetine 02467771 SAS ACDEFG  
 Atomoxetine 02445948 SIV ACDEFG  
 Auro-Atomoxetine 02471515 ARO ACDEFG  
 Jamp Atomoxetine 02506831 JPC ACDEFG  
 pms-Atomoxetine 02381052 PMS ACDEFG  
 Sandoz Atomoxetine 02386445 SDZ ACDEFG  
 Teva-Atomoxetine 02314584 TEV ACDEFG

Cap Orl 60 mg

Strattera 02262843 LIL ACDEFG  
 Apo-Atomoxetine 02318067 APX ACDEFG  
 Atomoxetine 02467798 SAS ACDEFG  
 Atomoxetine 02445956 SIV ACDEFG  
 Auro-Atomoxetine 02471523 ARO ACDEFG  
 Jamp Atomoxetine 02506858 JPC ACDEFG  
 pms-Atomoxetine 02381060 PMS ACDEFG  
 Sandoz Atomoxetine 02386453 SDZ ACDEFG  
 Teva-Atomoxetine 02314592 TEV ACDEFG

Cap Orl 80 mg

Strattera 02279347 LIL ACDEFG  
 Apo-Atomoxetine 02318075 APX ACDEFG  
 Atomoxetine 02467801 SAS ACDEFG  
 Auro-Atomoxetine 02471531 ARO ACDEFG  
 Jamp Atomoxetine 02506866 JPC ACDEFG  
 Sandoz Atomoxetine 02386461 SDZ ACDEFG  
 Teva-Atomoxetine 02362511 TEV ACDEFG

Cap Orl 100 mg

Strattera (Disc/non disp Jun 23/24) 02279355 LIL ACDEFG  
 Apo-Atomoxetine 02318083 APX ACDEFG  
 Atomoxetine 02467828 SAS ACDEFG  
 Auro-Atomoxetine 02471558 ARO ACDEFG  
 Jamp Atomoxetine 02506874 JPC ACDEFG  
 Sandoz Atomoxetine 02386488 SDZ ACDEFG

N06BA12 LISDEXAMFETAMINE

Cap Orl 10 mg

Vyvanse 02439603 TAK (SA)

Cap Orl 20 mg

Vyvanse 02347156 TAK (SA)

N06BA12 LISDEXAMFETAMINE

Cap	Orl	30 mg	Vyvanse	02322951	TAK	(SA)
Cap	Orl	40 mg	Vyvanse	02347164	TAK	(SA)
Cap	Orl	50 mg	Vyvanse	02322978	TAK	(SA)
Cap	Orl	60 mg	Vyvanse	02347172	TAK	(SA)
TabC	Orl	10 mg	Vyvanse	02490226	TAK	(SA)
TabC	Orl	20 mg	Vyvanse	02490234	TAK	(SA)
TabC	Orl	30 mg	Vyvanse	02490242	TAK	(SA)
TabC	Orl	40 mg	Vyvanse	02490250	TAK	(SA)
TabC	Orl	50 mg	Vyvanse	02490269	TAK	(SA)
TabC	Orl	60 mg	Vyvanse	02490277	TAK	(SA)

**N06D ANTI-DEMENTIA DRUGS**

**N06DA ANTICHOLINESTERASES**

N06DA02 DONEPEZIL

Tab	Orl	5 mg	Aricept	02232043	PFI	ACDEFV
			Apo-Donepezil	02362260	APX	ACDEFV
			Auro-Donepezil	02400561	ARO	ACDEFV
			Donepezil	02402645	AHI	ACDEFV
			Donepezil	02475278	RIV	ACDEFV
			Donepezil	02426846	SAS	ACDEFV
			Donepezil	02420597	SIV	ACDEFV
			Jamp-Donepezil	02416948	JPC	ACDEFV
			M-Donepezil	02467453	MRA	ACDEFV
			Mar-Donepezil	02402092	MAR	ACDEFV
			Mint-Donepezil	02408600	MNT	ACDEFV
			Nat-Donepezil	02439557	NAT	ACDEFV
			pms-Donepezil	02322331	PMS	ACDEFV
			Sandoz Donepezil	02328666	SDZ	ACDEFV
			Septa-Donepezil	02428482	SPT	ACDEFV
			Taro-Donepezil	02381508	SUN	ACDEFV
			Teva-Donepezil	02340607	TEV	ACDEFV

N06DA02 DONEPEZIL

Tab Orl 10 mg

Aricept	02232044	PFI	ACDEFV
Apo-Donepezil	02362279	APX	ACDEFV
Auro-Donepezil	02400588	ARO	ACDEFV
Donepezil	02402653	AHI	ACDEFV
Donepezil	02475286	RIV	ACDEFV
Donepezil	02426854	SAS	ACDEFV
Donepezil	02420600	SIV	ACDEFV
Jamp-Donepezil	02416956	JPC	ACDEFV
M-Donepezil	02467461	MRA	ACDEFV
Mar-Donepezil	02402106	MAR	ACDEFV
Mint-Donepezil	02408619	MNT	ACDEFV
Nat-Donepezil	02439565	NAT	ACDEFV
pms-Donepezil	02322358	PMS	ACDEFV
Sandoz Donepezil	02328682	SDZ	ACDEFV
Septa-Donepezil	02428490	SPT	ACDEFV
Taro-Donepezil	02381516	SUN	ACDEFV
Teva-Donepezil	02340615	TEV	ACDEFV

N06DA03 RIVASTIGMINE

Cap Orl 1.5 mg

Exelon	02242115	KNI	(SA)
Apo-Rivastigmine	02336715	APX	(SA)
Jamp-Rivastigmine	02485362	JPC	(SA)
Med-Rivastigmine	02401614	GMP	(SA)
Sandoz Rivastigmine	02324563	SDZ	(SA)

Cap Orl 3 mg

Exelon	02242116	KNI	(SA)
Apo-Rivastigmine	02336723	APX	(SA)
Jamp-Rivastigmine	02485370	JPC	(SA)
Med-Rivastigmine	02401622	GMP	(SA)
Sandoz Rivastigmine	02324571	SDZ	(SA)

Cap Orl 4.5 mg

Exelon	02242117	KNI	(SA)
Apo-Rivastigmine	02336731	APX	(SA)
Jamp-Rivastigmine	02485389	JPC	(SA)
Med-Rivastigmine	02401630	GMP	(SA)
Sandoz Rivastigmine	02324598	SDZ	(SA)

**N06DA03 RIVASTIGMINE**

Cap Orl 6 mg

Exelon	02242118	KNI	(SA)
Apo-Rivastigmine	02336758	APX	(SA)
Jamp-Rivastigmine	02485397	JPC	(SA)
Med-Rivastigmine	02401649	GMP	(SA)
Sandoz Rivastigmine	02324601	SDZ	(SA)

Liq Orl 2 mg

Exelon	02245240	KNI	(SA)
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**N06DA04 GALANTAMINE**

ERC Orl 8 mg

Auro-Galantamine ER	02425157	ARO	(SA)
Galantamine ER	02443015	SAS	(SA)
Mylan-Galantamine ER	02339439	MYL	(SA)
pms-Galantamine ER	02398370	PMS	(SA)

ERC Orl 16 mg

Auro-Galantamine ER	02425165	ARO	(SA)
Galantamine ER	02443023	SAS	(SA)
Mylan-Galantamine ER	02339447	MYL	(SA)
pms-Galantamine ER	02398389	PMS	(SA)

ERC Orl 24 mg

Auro-Galantamine ER	02425173	ARO	(SA)
Galantamine ER	02443031	SAS	(SA)
Mylan-Galantamine ER	02339455	MYL	(SA)
pms-Galantamine ER	02398397	PMS	(SA)

**N07 OTHER NERVOUS SYSTEM DRUGS****N07A PARASYMPATHOMIMETICS****N07AA ANTICHOLINESTERASES****N07AA01 NEOSTIGMINE**

Liq Inj 1 mg/mL

Neostigmine Omega	02230592	OMG	V
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Liq Inj 2.5 mg/mL

Neostigmine Omega	02387166	OMG	V
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**N07AA02 PYRIDOSTIGMINE**

SRT Orl 180 mg

Mestinon SR	00869953	BSL	ACDEFGV
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Tab Orl 60 mg

Mestinon	00869961	BSL	ACDEFGV
Jamp Pyridostigmine Bromide	02508362	JPC	ACDEFGV
Riva-Pyridostigmine	02495643	RIV	ACDEFGV

**N07AB CHOLINE ESTERS****N07AB02 BETHANECHOL**

N07AB02 BETHANECHOL

Tab	Orl	10 mg	Duvoid	01947958	PAL	ACDEFGV
Tab	Orl	25 mg	Duvoid	01947931	PAL	ACDEFGV
Tab	Orl	50 mg	Duvoid	01947923	PAL	ACDEFGV

**N07AX OTHER PARASYMPATHOMIMETICS**

N07AX01 PILOCARPINE

Tab	Orl	5 mg	Salagen	02216345	MTP	(SA)
			Accel-Pilocarpine	02496119	MRA	(SA)
			Jamp Pilocarpine	02509571	JPC	(SA)

**N07B DRUGS USED IN ADDICTIVE DISORDERS**

**N07BA DRUGS USED IN NICOTINE DEPENDENCE**

N07BA01 NICOTINE

Gum	Orl	2 mg	Actavis	80015240	ACT	(SA)
			Compliments	80015240	SOB	(SA)
			Exact	80025660	SDM	(SA)
			Life Brand	80025660	SDM	(SA)
			Personnelle	80015240	PJC	(SA)
Loz	Orl	1 mg	Nic-Hit (mini-lozenge)	80061161	NHI	(SA)
Loz	Orl	2 mg	Nic-Hit (mini-lozenge)	80059877	NHI	(SA)
Loz	Orl	4 mg	Nic-Hit (mini-lozenge)	80059869	NHI	(SA)
Pth	Trd	7 mg	Actavis	80044393	ACT	(SA)
			Compliments	80044393	SOB	(SA)
			Equate	02241227	WAL	(SA)
			Exact	80014321	SDM	(SA)
			Life Brand	80014321	SDM	(SA)
			Personnelle	80044393	PJC	(SA)
			Pharmasave	02241227	PSV	(SA)
			Pharmasave	80014321	PSV	(SA)

N07BA01 NICOTINE

Pth Trd 14 mg

Actavis 80044392 ACT (SA)  
 Compliments 80044392 SOB (SA)  
 Equate 02241226 WAL (SA)  
 Exact 80013549 SDM (SA)  
 Life Brand 80013549 SDM (SA)  
 Personnelle 80044392 PJC (SA)  
 Pharmasave 02241226 PSV (SA)  
 Pharmasave 80013549 PSV (SA)

Pth Trd 21 mg

Actavis 80044389 ACT (SA)  
 Compliments 80044389 SOB (SA)  
 Equate 02241228 WAL (SA)  
 Exact 80014250 SDM (SA)  
 Life Brand 80014250 SDM (SA)  
 Personnelle 80044389 PJC (SA)  
 Pharmasave 02241228 PSV (SA)  
 Pharmasave 80014250 PSV (SA)

N07BA03 VARENICLINE

Kit Orl 0.5 mg, 1 mg

Champix Starter Kit 02298309 PFI (SA)  
 Apo-Varenicline 02435675 APX (SA)  
 Teva-Varenicline 02426781 TEV (SA)

Tab Orl 0.5 mg

Champix 02291177 PFI (SA)  
 Apo-Varenicline 02419882 APX (SA)  
 Teva-Varenicline 02426226 TEV (SA)

Tab Orl 1 mg

Champix 02291185 PFI (SA)  
 Apo-Varenicline 02419890 APX (SA)  
 Teva-Varenicline 02426234 TEV (SA)

**N07BB DRUGS USED IN ALCOHOL DEPENDENCE**

N07BB03 ACAMPROSATE

SRT Orl 333 mg

Campral 02293269 MYL ACDEFGV

N07BB04 NALTREXONE

Tab Orl 50 mg

Revia 02213826 TEV ACDEFGV  
 Apo-Naltrexone 02444275 APX ACDEFGV  
 Naltrexone Hydrochloride 02451883 JPC ACDEFGV

**N07BC DRUGS USED IN OPIOID DEPENDENCE****N07BC01 BUPRENORPHINE**

Kit	SC	80 mg	Probuphine (Disc/non disp May 30/24)	02474921	KNI	(SA)
Liq	SC	100 mg / 0.5 mL	Sublocade	02483084	IUK	ACDEFGV
Liq	SC	300 mg / 1.5 mL	Sublocade	02483092	IUK	ACDEFGV

**N07BC02 METHADONE**

Liq	Orl	1 mg/mL	Metadol	02247694	PAL	(SA)
Liq	Orl	10 mg/mL	Metadol	02241377	PAL	(SA)
			Metadol-D	02244290	PAL	ACDEFGV
			Jamp-Methadone	02495783	JPC	ACDEFGV
			Odan-Methadone (cherry flavoured)	02495872	ODN	ACDEFGV
			Odan-Methadone (unflavoured)	02495880	ODN	ACDEFGV

Pws	Orl		Methadone Compounded Oral Solution			
			Opioid Dependence / dépendance aux opiacés	00999734		ACDEFGV
			Pain Management / gestion de la douleur	00999801		(SA)

Tab	Orl	1 mg	Metadol	02247698	PAL	(SA)
Tab	Orl	5 mg	Metadol	02247699	PAL	(SA)
Tab	Orl	10 mg	Metadol	02247700	PAL	(SA)
Tab	Orl	25 mg	Metadol	02247701	PAL	(SA)

**N07BC51 BUPRENORPHINE, COMBINATIONS  
BUPRENORPHINE / NALOXONE**

Slit	Orl	2 mg / 0.5 mg	Suboxone	02295695	IUK	ACDEFGV
			Act Buprenorphine/Naloxone	02453908	TEV	ACDEFGV
			pms-Buprenorphine/Naloxone	02424851	PMS	ACDEFGV
Slit	Orl	8 mg / 2 mg	Suboxone	02295709	IUK	ACDEFGV
			Act Buprenorphine/Naloxone	02453916	TEV	ACDEFGV
			pms-Buprenorphine/Naloxone	02424878	PMS	ACDEFGV

**N07C ANTIVERTIGO PREPARATIONS****N07CA ANTIVERTIGO PREPARATIONS****N07CA01 BETAHISTINE**

N07CA01 BETAHISTINE

Tab Orl 16 mg

Serc 02243878 BGP ACDEFGV  
 Auro-Betahistine 02449153 ARO ACDEFGV  
 Betahistine 02466449 SAS ACDEFGV  
 M-Betahistine 02519690 MRA ACDEFGV  
 pms-Betahistine 02330210 PMS ACDEFGV  
 Teva-Betahistine 02280191 TEV ACDEFGV

Tab Orl 24 mg

Serc 02247998 BGP ACDEFGV  
 Auro-Betahistine 02449161 ARO ACDEFGV  
 Betahistine 02466457 SAS ACDEFGV  
 M-Betahistine 02519704 MRA ACDEFGV  
 pms-Betahistine 02330237 PMS ACDEFGV  
 Teva-Betahistine 02280205 TEV ACDEFGV

N07CA03 FLUNARIZINE

Cap Orl 5 mg

Flunarizine 02246082 AAP ACDEFGV

**N07X OTHER NERVOUS SYSTEM DRUGS**

**N07XX OTHER NERVOUS SYSTEM DRUGS**

N07XX02 RILUZOLE

Tab Orl 50 mg

Rilutek 02242763 SAV ACDEFV  
 Apo-Riluzole 02352583 APX ACDEFV  
 Mylan-Riluzole 02390299 MYL ACDEFV

N07XX05 AMIFAMPRIDINE

Tab Orl 10 mg

Ruzurgi 02503034 MDU (SA)

N07XX06 TETRABENAZINE

Tab Orl 25 mg

Nitoman 02199270 BSL ACDEFGV  
 Apo-Tetrabenazine 02407590 APX ACDEFGV  
 pms-Tetrabenazine 02402424 PMS ACDEFGV

N07XX08 TAFAMIDIS

Cap Orl 20 mg

Vyndaqel 02495732 PFI (SA)

Cap Orl 61 mg

Vyndamax 02517841 PFI (SA)

N07XX12 PATISIRAN

Liq IV 2 mg/mL

Onpattro 02489252 ALN (SA)

N07XX14 EDARAVONE



N07XX14	EDARAVONE								
	Liq	IV	0.3 mg/mL			Radicava	02475472	MBT	(SA)
	Susp	Orl	105 mg / 5 mL			Radicava	02532611	MBT	(SA)
N07XX15	INOTERSEN								
	Liq	SC	284 mg / 1.5 mL			Tegsedi	02481383	AKT	(SA)
N07XX99	SODIUM PHENYL BUTYRATE AND URSODOXICOLTAURINE								
	Pws.	Orl	3 g / 1 g			Albrioza	02527707	ALY	(SA)

**P ANTIPARASITIC PRODUCTS, INSECTICIDES AND REPELLANTS**

**P01 ANTIPROTOZOALS**

**P01A AGENTS AGAINST AMOEBIASIS & OTHER PROTOZOAL DISEASES**

**P01AX OTHER AGENTS AGAINST AMOEBIASIS & OTHER PROTOZOAL DISEASES**

P01AX06	ATOVAQUONE								
	Sus	Orl	750 mg / 5 mL			Mepron	02217422	GSK	ACDEFGV
						GLN-Atovaquone	02528495	GLM	ACDEFGV

**P01B ANTIMALARIALS**

**P01BA AMINOQUINOLINES**

P01BA02	HYDROXYCHLOROQUINE								
	Tab	Orl	200 mg			Plaquenil	02017709	SAV	ACDEFGV
						Apo-Hydroxyquine	02246691	APX	ACDEFGV
						Hydroxychloroquine	02519348	SAS	ACDEFGV
						Jamp-Hydroxychloroquine Sulfate	02491427	JPC	ACDEFGV
						Mint-Hydroxychloroquine	02424991	MNT	ACDEFGV
						NRA-Hydroxychloroquine	02511886	NRA	ACDEFGV

P01BA03	PRIMAQUINE								
	Tab	Orl	15 mg			Primaquine	02017776	SAV	ACDEFGV

**P01C AGENTS AGAINST LEISHMANIASIS AND TRYPANOSOMIASIS**

**P01CX OTHER AGENTS AGAINST LEISHMANIASIS AND TRYPANOSOMIASIS**

P01CX01	PENTAMIDINE ISETIONATE								
	Pws	Inj	300 mg			Pentamidine Isetionate	02183080	PFI	ACDEFGV

**P02 ANTHELMINTICS**

**P02B ANTIREMATODALS**

**P02BA QUINOLINE DERIVATIVES AND RELATED SUBSTANCES**

P02BA01	PRAZIQUANTEL								
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P02BA01 PRAZIQUANTEL

Tab Orl 600 mg

Biltricide 02230897 BAY ACDEFGV

**P02C ANTINEMATODAL AGENTS**

**P02CA BENZIMIDAZOLE AGENTS**

P02CA01 MEBENDAZOLE

Tab Orl 100 mg

Vermox 00556734 JAN ACDEFGV

**P02CC TETRAHYDROPIRIMIDINE DERIVATIVES**

P02CC01 PYRANTEL

Tab Orl 125 mg

Combantrin 01944363 JNJ EFG

**P03 ECTOPARASITICIDES, INCLUDING SCABICIDES, INSECTICIDES & REPELLANTS**

**P03A ECTOPARASITICIDES, INCLUDING SCABICIDES**

**P03AC PYRETHRINS, INCLUDING SYNTHETIC COMPOUNDS**

P03AC04 PERMETHRIN

Crm Top 1%

Kwellada-P Cream Rinse 1% 02231480 MDI EFGV

Nix Cream 00771368 INP EFGV

Crm Top 5%

Nix Dermal 02219905 GCH EFGV

Lot Top 5%

Kwellada-P 02231348 MDI EFGV

P03AC51 PYRETHRUM, COMBINATIONS

PYRETHRINS / PIPERONYL BUTOXIDE

Shp Top 0.33% / 3%

R & C Shampoo and Conditioner 02125447 MDI EFGV

**P03AX OTHER ECTOPARACITICIDES, INCLUDING SCABICIDES**

CROTAMITON

Crm Top 10%

Eurax 00623377 CLC EFGV

ISOPROPYL MYRISTATE

Liq Top 50%

Resultz 02279592 ARZ EFGV

**R RESPIRATORY SYSTEM**

**R01 NASAL PREPARATIONS**

**R01A DECONGESTANTS AND OTHER NASAL PREPARATIONS FOR TOPICAL USE**

**R01AD CORTICOSTEROIDS**

R01AD01 BECLOMETHASONE

Aem Nas 50 mcg

Apo-Beclo methasone AQ 02238796 APX ACDEFGV

Mylan-Beclo AQ 02172712 MYL ACDEFGV

R01AD05	BUDESONIDE							
Aem	Nas	64 mcg						
			Rhinocort Aqua	02231923	JNJ	ACDEFGV		
			Mylan-Budesonide AQ	02241003	MYL	ACDEFGV		
Aem	Nas	100 mcg						
			Mylan-Budesonide AQ	02230648	MYL	ACDEFGV		

R01AD08	FLUTICASONE							
Aem	Nas	50 mcg						
			Apo-Fluticasone	02294745	APX	ABCDEFGV		

R01AD09	MOMETASONE							
Asp	Nas	0.1%						
			Nasonex Aqueous	02238465	ORG	ACDEFGV		
			Apo-Mometasone	02403587	APX	ACDEFGV		
			Mometasone	02519127	SAS	ACDEFGV		
			Sandoz Mometasone	02449811	SDZ	ACDEFGV		
			Teva-Mometasone	02475863	TEV	ACDEFGV		

R01AD11	TRIAMCINOLONE							
Liq	Nas	55 mcg						
			Nasacort AQ	02213834	SNC	ACDEFGV		
			Apo-Triamcinolone AQ	02437635	APX	ACDEFGV		

**R01AX OTHER NASAL PREPARATIONS**

R01AX03	IPRATROPIUM BROMIDE							
Spr	Nas	0.03%						
			pms-Ipratropium	02239627	PMS	ACDEFGV		

**R03 DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES**

**R03A ADRENERGICS, INHALANTS**

**R03AC SELECTIVE BETA2-ADRENOCEPTOR AGONISTS**

R03AC02	SALBUTAMOL							
Aem	Inh	100 mcg						
			Airomir	02232570	BSL	ABCDEFGVW		
			Ventolin	02241497	GSK	ABCDEFGVW		
			Apo-Salvent CFC Free	02245669	APX	ABCDEFGVW		
			Novo-Salbutamol HFA	02326450	TEV	ABCDEFGVW		
			Salbutamol HFA	02419858	SAS	ABCDEFGVW		
Liq	Inh	0.5 mg/mL						
			pms-Salbutamol	02208245	PMS	W (SA)		
Liq	Inh	1 mg/mL						
			pms-Salbutamol	02208229	PMS	W (SA)		
			Teva-Salbutamol Sterinebs	01926934	TEV	W (SA)		
Liq	Inh	2 mg/mL						
			pms-Salbutamol	02208237	PMS	W (SA)		
			Teva-Salbutamol Sterinebs	02173360	TEV	W (SA)		

R03AC02	SALBUTAMOL							
	Liq	Inh	5 mg/mL		Ventolin	02213486	GSK	W (SA)
	Pwr	Inh	200 mcg		Ventolin Diskus	02243115	GSK	ACDEFGVW
R03AC03	TERBUTALINE							
	Pwr	Inh	0.5 mg		Bricanyl Turbuhaler	00786616	AZE	ACDEFGV
R03AC12	SALMETEROL							
	Pwr	Inh	50 mcg		Serevent Diskus	02231129	GSK	(SA)
R03AC13	FORMOTEROL							
	Cap	Inh	12 mcg		Foradil (Disc/non disp Jan 20/24)	02230898	NVR	(SA)
	Pwr	Inh	6 mcg		Oxeze Turbuhaler	02237225	AZE	(SA)
	Pwr	Inh	12 mcg		Oxeze Turbuhaler	02237224	AZE	(SA)
R03AC18	INDACATEROL							
	Cap	Inh	75 mcg		Onbrez Breezhaler	02376938	NVR	(SA)

**R03AK ADRENERGICS AND OTHER DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES**

R03AK06	SALMETEROL AND FLUTICASONE							
	Aem	Inh	25 mcg / 125 mcg		Advair	02245126	GSK	(SA)
	Aem	Inh	25 mcg / 250 mcg		Advair	02245127	GSK	(SA)
	Pwr	Inh	50 mcg / 100 mcg		Advair Diskus	02240835	GSK	(SA)
					pms-Fluticasone Propionate/Salmeterol	02494507	PMS	(SA)
					Wixela Inhub	02495597	MYL	(SA)
	Pwr	Inh	50 mcg / 250 mcg		Advair Diskus	02240836	GSK	(SA)
					pms-Fluticasone Propionate/Salmeterol	02494515	PMS	(SA)
					Wixela Inhub	02495600	MYL	(SA)
	Pwr	Inh	50 mcg / 500 mcg		Advair Diskus	02240837	GSK	(SA)
					pms-Fluticasone Propionate/Salmeterol	02494523	PMS	(SA)
					Wixela Inhub	02495619	MYL	(SA)
R03AK07	FORMOTEROL AND BUDESONIDE							
	Pwr	Inh	6 mcg / 100 mcg		Symbicort Turbuhaler	02245385	AZE	(SA)
	Pwr	Inh	6 mcg / 200 mcg		Symbicort Turbuhaler	02245386	AZE	(SA)

R03AK09	FORMOTEROL AND MOMETASONE					
Aem	Inh	5 mcg / 100 mcg		Zenhale	02361752	ORG (SA)
Aem	Inh	5 mcg / 200 mcg		Zenhale	02361760	ORG (SA)
R03AK10	VILANTEROL AND FLUTICASONE					
Pwr	Inh	25 mcg / 100 mcg		Breo Ellipta	02408872	GSK (SA)
Pwr	Inh	25 mcg / 200 mcg		Breo Ellipta	02444186	GSK (SA)
R03AK14	INDACATEROL AND MOMETASONE					
Cap	Inh	150 mcg / 80 mcg		Aectura Breezhaler	02498685	NVR (SA)
Cap	Inh	150 mcg / 160 mcg		Aectura Breezhaler	02498707	NVR (SA)
Cap	Inh	150 mcg / 320 mcg		Aectura Breezhaler	02498693	NVR (SA)

**R03AL ADRENERGICS IN COMBINATION WITH ANTICHOLINERGICS**

R03AL02	SALBUTAMOL AND IPRATROPIUM BROMIDE					
Liq	Inh	0.5 mg / 2.5 mg / 2.5 mL	Ipratropium Bromide and Salbutamol Sulfate		02483394	JNO (SA)
				Teva-Combo Sterinebs	02272695	TEV (SA)
Liq	Inh	100 mcg / 20 mcg		Combivent RespiMAT	02419106	BOE ACDEFGV
R03AL03	VILANTEROL AND UMECLIDINIUM BROMIDE					
Pwr	Inh	25 mcg / 62.5 mcg		Anoro Ellipta	02418401	GSK (SA)
R03AL04	INDACATEROL AND GLYCOPYRRONIUM BROMIDE					
Cap	Inh	110 mcg / 50 mcg		Ultibro Breezhaler	02418282	CPC (SA)
R03AL05	FORMOTEROL AND ACLIDINIUM BROMIDE					
Pwr	Inh	12 mcg / 400 mcg		Duaklir Genuair	02439530	CPC (SA)
R03AL06	OLODATEROL AND TIOTROPIUM BROMIDE					
Liq	Inh	2.5 mcg / 2.5 mcg		Inspiolto RespiMAT	02441888	BOE (SA)
R03AL08	VILANTEROL, UMECLIDINIUM BROMIDE AND FLUTICASONE					
Pwr	Inh	25 mcg / 62.5 mcg / 100mcg		Trelegy Ellipta	02474522	GSK (SA)
R03AL11	FORMOTEROL GLYCOPYRRONIUM BROMIDE AND BUDESONIDE					
Aem	Inh	5.8 mcg / 8.2 mcg / 182 mcg		Breztri Aerosphere	02518058	AZE (SA)

## R03AL12 INDACATEROL, GLYCOPYRRONIUM BROMIDE AND MOMETASONE

Cap	Inh	150 mcg / 50 mcg / 150mcg	Energair Breezhaler	02501244	VAL	(SA)
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**R03B OTHER DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES, INHALANTS****R03BA GLUCOCORTICOIDS**

## R03BA01 BECLOMETHASONE

Aem	Inh	50 mcg		Qvar	02242029	BSL	ACDEFGV
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Aem	Inh	100 mcg		Qvar	02242030	BSL	ACDEFGV
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## R03BA02 BUDESONIDE

Pwr	Inh	100 mcg	Pulmicort Turbuhaler	00852074	AZE	ACDEFGV
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Pwr	Inh	200 mcg	Pulmicort Turbuhaler	00851752	AZE	ACDEFGV
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Pwr	Inh	400 mcg	Pulmicort Turbuhaler	00851760	AZE	ACDEFGV
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Sus	Inh	0.125 mg/mL	Pulmicort Nebuamp	02229099	AZE	(SA)
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Taro-Budesonide	02494264	TAR	(SA)
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Teva-Budesonide	02465949	TEV	(SA)
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Sus	Inh	0.25 mg/mL	Pulmicort Nebuamp	01978918	AZE	(SA)
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Taro-Budesonide	02494272	TAR	(SA)
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Sus	Inh	0.5 mg/mL	Pulmicort Nebuamp	01978926	AZE	(SA)
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Taro-Budesonide	02494280	TAR	(SA)
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Teva-Budesonide	02465957	TEV	(SA)
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## R03BA05 FLUTICASONE

Aem	Inh	50 mcg	Flovent Metered Dose HFA	02244291	GSK	ACDEFGV
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Aem	Inh	125 mcg	Flovent Metered Dose HFA	02244292	GSK	ACDEFGV
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Apo-Fluticasone HFA	02526557	APX	ACDEFGV
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pms-Fluticasone HFA	02503123	PMS	ACDEFGV
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Aem	Inh	250 mcg	Flovent Metered Dose HFA	02244293	GSK	ACDEFGV
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Apo-Fluticasone HFA	02510987	APX	ACDEFGV
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pms-Fluticasone HFA	02503131	PMS	ACDEFGV
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Pwr	Inh	55 mcg	Aermony Respiclick	02467895	TEV	ACDEFGV
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Pwr	Inh	100 mcg	Flovent Diskus	02237245	GSK	ACDEFGV
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R03BA05	FLUTICASONE							
Pwr	Inh	113 mcg		Aermony Respiclick	02467909	TEV	ACDEFGV	
Pwr	Inh	232 mcg		Aermony Respiclick	02467917	TEV	ACDEFGV	
Pwr	Inh	250 mcg		Flovent Diskus	02237246	GSK	ACDEFGV	
Pwr	Inh	500 mcg		Flovent Diskus	02237247	GSK	ACDEFGV	

R03BA07	MOMETASONE							
Pwr	Inh	100 mcg		Asmanex Twisthaler	02438690	ORG	CDEFG	
Pwr	Inh	200 mcg		Asmanex Twisthaler	02243595	ORG	ACDEFGV	
Pwr	Inh	400 mcg		Asmanex Twisthaler	02243596	ORG	ACDEFGV	

R03BA08	CICLESONIDE							
Aem	Inh	100 mcg		Alvesco	02285606	CPC	ACDEFGV	
Aem	Inh	200 mcg		Alvesco	02285614	CPC	ACDEFGV	

R03BA09	FLUTICASONE FUROATE							
Pwr	Inh	100 mcg		Arnuity Ellipta	02446561	GSK	ACDEFGV	
Pwr	Inh	200 mcg		Arnuity Ellipta	02446588	GSK	ACDEFGV	

**R03BB ANTICHOLINERGICS**

R03BB01	IPRATROPIUM BROMIDE							
Aem	Inh	20 mcg		Atrovent HFA	02247686	BOE	ABCDEFGHIWV	
Liq	Inh	125 mcg/mL		pms-lpratropium	02231135	PMS	(SA)	
Liq	Inh	250 mcg/mL		Apo-lpravent	02126222	APX	W (SA)	
				pms-lpratropium	02231136	PMS	W (SA)	
				pms-lpratropium (1mL nebulas)	02231244	PMS	W (SA)	
				pms-lpratropium (2mL nebulas)	02231245	PMS	W (SA)	
				Teva-lpratropium	02216221	TEV	W (SA)	

R03BB04	TIOTROPIUM BROMIDE							
Cap	Inh	18 mcg		Spiriva	02246793	BOE	(SA)	
Liq	Inh	2.5 mcg		Spiriva Respimat	02435381	BOE	(SA)	

R03BB05 ACLIDINIUM BROMIDE

Pwr Inh 400 mcg

Tudorza Genuair 02409720 ALM (SA)

R03BB06 GLYCOPYRRONIUM BROMIDE

Cap Inh 50 mcg

Seebri Breezhaler 02394936 CPC (SA)

R03BB07 UMECLIDIINIUM BROMIDE

Pwr Inh 62.5 mcg

Incruse Ellipta 02423596 GSK (SA)

R03BX OTHER DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES, INHALANTS

R03BX99 HYPERTONIC SODIUM CHLORIDE

Liq Inh 7%

Hyper-Sal 80029414 KEG BCDEFG

Nebusal 80029758 STR BCDEFG

R03D OTHER SYSTEMIC DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES

R03DA XANTHINES

R03DA04 THEOPHYLLINE

Liq Orl 80 mg / 15 mL

Theolair 01966219 BSL ACDEFGV

SRT Orl 100 mg

AA-Theo LA 00692689 AAP ACDEFGV

SRT Orl 200 mg

AA-Theo LA 00692697 AAP ACDEFGV

SRT Orl 300 mg

AA-Theo LA 00692700 AAP ACDEFGV

SRT Orl 400 mg

Theo ER 02360101 AAP ACDEFGV

SRT Orl 600 mg

Theo ER 02360128 AAP ACDEFGV

R03DC LEUKOTRIENE RECEPTOR ANTAGONISTS

R03DC03 MONTELUKAST

Gran Orl 4 mg

Singulair 02247997 ORG ACDEFGV

Sandoz Montelukast 02358611 SDZ ACDEFGV



R03DC03 MONTELUKAST

Tab Orl 10 mg

Singulair	02238217	ORG	ACDEFGV
Apo-Montelukast	02374609	APX	ACDEFGV
Auro-Montelukast	02401274	ARO	ACDEFGV
Jamp-Montelukast	02391422	JPC	ACDEFGV
M-Montelukast	02488183	MRA	ACDEFGV
Mar-Montelukast	02399997	MAR	ACDEFGV
Mint-Montelukast	02408643	MNT	ACDEFGV
Montelukast	02379333	SAS	ACDEFGV
Montelukast	02382474	SIV	ACDEFGV
Montelukast Sodium	02379236	AHI	ACDEFGV
Nat-Montelukast	02522136	NAT	ACDEFGV
NRA-Montelukast	02489821	NRA	ACDEFGV
pms-Montelukast	02373947	PMS	ACDEFGV
Sandoz Montelukast	02328593	SDZ	ACDEFGV
Taro-Montelukast	02389517	SUN	ACDEFGV
Teva-Montelukast	02355523	TEV	ACDEFGV

TabC Orl 4 mg

Singulair	02243602	ORG	ACDEFGV
Apo-Montelukast	02377608	APX	ACDEFGV
Jamp Montelukast Chewable	02514877	JPC	ACDEFGV
Jamp-Montelukast (Disc/non disp Apr 20/24)	02442353	JPC	ACDEFGV
Mar-Montelukast	02399865	MAR	ACDEFGV
Mint-Montelukast	02408627	MNT	ACDEFGV
Montelukast	02382458	SIV	ACDEFGV
Nat-Montelukast	02522101	NAT	ACDEFGV
pms-Montelukast	02354977	PMS	ACDEFGV
Sandoz Montelukast	02330385	SDZ	ACDEFGV
Teva-Montelukast	02355507	TEV	ACDEFGV

R03DC03 MONTELUKAST

TabC Orl 5 mg

Singulair	02238216	ORG	ACDEFGV
Apo-Montelukast	02377616	APX	ACDEFGV
Jamp Montelukast Chewable	02514885	JPC	ACDEFGV
Jamp-Montelukast (Disc/non disp Apr 20/24)	02442361	JPC	ACDEFGV
Mar-Montelukast	02399873	MAR	ACDEFGV
Mint-Montelukast	02408635	MNT	ACDEFGV
Montelukast	02379325	SAS	ACDEFGV
Montelukast	02382466	SIV	ACDEFGV
Nat-Montelukast	02522128	NAT	ACDEFGV
pms-Montelukast	02354985	PMS	ACDEFGV
Sandoz Montelukast	02330393	SDZ	ACDEFGV
Teva-Montelukast	02355515	TEV	ACDEFGV

**R03DX OTHER SYSTEMIC DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES**

R03DX05 OMALIZUMAB

Liq SC 150 mg/mL

Xolair (prefilled syringe) 02459795 NVR (SA)

Pws SC 150 mg

Xolair (single-use vial) 02260565 NVR (SA)

R03DX09 MEPOLIZUMAB

Liq SC 100 mg/mL

Nucala (autoinjector) 02492989 GSK (SA)

Nucala (prefilled syringe) 02492997 GSK (SA)

Pws SC 100 mg/mL

Nucala (single-use vial) 02449781 GSK (SA)

R03DX10 BENRALIZUMAB

Liq SC 30 mg/mL

Fasenra (autoinjector) 02496135 AZE (SA)

Fasenra (prefilled syringe) 02473232 AZE (SA)

**R05 COUGH AND COLD PREPARATIONS**

**R05C EXPECTORANTS, EXCLUDING COMBINATIONS WITH COUGH SUPPRESSANTS**

**R05CA EXPECTORANTS**

R05CA03 GUAIFENESIN

Syr Orl 100 mg / 5 mL

Balminil 00608920 TEV G

Balminil Expect Sans Sucrose 00609951 TEV G

Robitussin 01931032 GCH G

**R05CB MUCOLYTICS**

R05CB01 ACETYLCYSTEINE

Liq Inh 200 mg/mL

Acetylcysteine 02243098 SDZ ACDEFGV

R05CB13 DORNASE ALFA

Liq Inh 1 mg/mL

Pulmozyme 02046733 HLR (SA)

**R05D COUGH SUPPRESSANTS, EXCLUDING COMBINATIONS WITH EXPECTORANTS**

**R05DA OPIUM ALKALOIDS AND DERIVATIVES**

R05DA04 CODEINE

Liq Inj 30 mg/mL

Codeine Phosphate 00544884 SDZ W

SRT Orl 50 mg

Codeine Contin 02230302 PFR W (SA)

SRT Orl 100 mg

Codeine Contin 02163748 PFR W (SA)

SRT Orl 150 mg

Codeine Contin 02163780 PFR W (SA)

SRT Orl 200 mg

Codeine Contin 02163799 PFR W (SA)

Syr Orl 5 mg/mL

Codeine Phosphate 00050024 ATL ACDEFGVW

Tab Orl 15 mg

Teva-Codeine 00593435 TEV ACDEFGVW

Tab Orl 30 mg

Teva-Codeine 00593451 TEV ACDEFGVW

R05DA09 DEXTROMETHORPHAN

Liq Orl 15 mg/mL

Koffex Sugar Free Clear 01928791 TEV G

Syr Orl 3 mg/mL

Benylin DM 01944738 JNJ G

Koffex DM 01928783 TEV G

**R05F COUGH SUPPRESSANTS AND EXPECTORANTS, COMBINATIONS**

**R05FA OPIUM DERIVATIVES AND EXPECTORANTS**

R05FA02 OPIUM DERIVATIVES AND EXPECTORANTS

DEXTROMETHORPHAN / GUAIFENESIN

Liq Orl 3 mg / 20 mg

Robitussin DM Exp 01931024 GCH G

**R06 ANTIHISTAMINES FOR SYSTEMIC USE**

**R06A ANTIHISTAMINES FOR SYSTEMIC USE**

**R06AA AMINOALKYL ETHERS**

R06AA02 DIPHENHYDRAMINE

Elx Orl 12.5 mg / 5 mL

Benadryl 02019736 JNJ G

Diphenhydramine HCl Elixir USP 02298503 JPC G

R06AA02	DIPHENHYDRAMINE						
Liq	Inj	50 mg/mL	Diphenhydramine HCl	00596612	SDZ	ACDEFGVW	
			Diphenist	02219336	OMG	ACDEFGVW	
Tab	Orl	25 mg	Benadryl	02017849	JNJ	G	
			Diphenhydramine	02257548	JPC	G	
Tab	Orl	50 mg	Diphenhydramine	02257556	JPC	G	
R06AA11	DIMENHYDRINATE						
Liq	Inj	50 mg/mL	Dimenhydrinate Injection USP	00392537	SDZ	ACDEFGVW	
R06AA59	DOXYLAMINE, COMBINATIONS						
	DOXYLAMINE / PYRIDOXINE						
SRT	Orl	10 mg / 10 mg	Diclectin	00609129	DUI	ACDEFGV	
			Apo-Doxylamine/B6	02413248	APX	ACDEFGV	
			pms-Doxylamine-Pyridoxine	02406187	PMS	ACDEFGV	

**R06AB SUBSTITUTED ALKYL AMINES**

R06AB04	CHLORPHENAMINE (CHLORPHENIRAMINE)						
Tab	Orl	4 mg	Novo-Pheniram	00021288	TEV	G	

**R06AD PHENOTHIAZINE DERIVATIVES**

R06AD01	ALIMEMAZINE (TRIMEPRAZINE)						
Tab	Orl	2.5 mg	Panectyl	01926306	SLP	ACDEFGV	
Tab	Orl	5 mg	Panectyl	01926292	SLP	ACDEFGV	

**R06AE PIPERAZINE DERIVATIVES**

R06AE07	CETIRIZINE						
Tab	Orl	10 mg	Reactine	02223554	JNJ	G	
			Apo-Cetirizine	02231603	APX	G	
			Cetirizine Extra Strength	02517566	JPC	G	
			Jamp Cetirizine	02451778	JPC	G	

R06AE07 CETIRIZINE

Tab Orl 20 mg

Reactine 01900978 JNJ (SA)  
 Apo-Cetirizine 02453363 APX (SA)  
 Cetirizine 02515695 SAS (SA)  
 Jamp Cetirizine Tablets 02517353 JPC (SA)  
 M-Cetirizine 02512025 MRA (SA)  
 Mar-Cetirizine 02427141 MAR (SA)  
 pms-Cetirizine 02315963 PMS (SA)  
 Teva-Cetirizine 02528681 TEV (SA)

**R06AX OTHER ANTIHISTAMINES FOR SYSTEMIC USE**

R06AX13 LORATADINE

Tab Orl 10 mg

Claritin 00782696 BAY G  
 Apo-Loratadine 02243880 APX G

R06AX17 KETOTIFEN

Tab Orl 1 mg

Zaditen 00577308 TEV CDEFG

**R07 OTHER RESPIRATORY SYSTEM PRODUCTS**

**R07A OTHER RESPIRATORY SYSTEM PRODUCTS**

**R07AX OTHER RESPIRATORY SYSTEM PRODUCTS**

R07AX02 IVACAFTOR

Tab Orl 150 mg

Kalydeco 02397412 VTX (SA)

R07AX32 IVACAFTOR, TEZACAFTOR AND ELEXACAFTOR

Tab Orl 37.5 mg / 25 mg / 50 mg,  
75mg

Trikafta 02526670 VTX (SA)

Tab Orl 75 mg / 50 mg / 100 mg,  
150mg

Trikafta 02517140 VTX (SA)

**S SENSORY ORGANS**

**S01 OPHTHALMOLOGICALS**

**S01A ANTIINFECTIVES**

**S01AA ANTIBIOTICS**

S01AA12 TOBRAMYCIN

Liq Oph 0.3%

Tobrex 00513962 NVR ACDEFGV  
 Sandoz Tobramycin 02241755 SDZ ACDEFGV

Ont Oph 0.3%

Tobrex 00614254 NVR ACDEFGV

S01AA17 ERYTHROMYCIN

S01AA17 ERYTHROMYCIN  
 Ont Oph 0.5% Erythromycin 02326663 SGQ ACDEFGV  
 pdp-Erythromycin 01912755 PDP ACDEFGV

S01AA30 COMBINATIONS OF DIFFERENT ANTIBIOTICS  
 POLYMYXIN B SULFATE / BACITRACIN ZINC  
 Ont Oph 10 000 IU / 500 IU Polysporin 02239157 JNJ G  
 POLYMYXIN B SULFATE / TRIMETHOPRIM SULFATE  
 Liq Oph 10000 U/mL Polytrim (Disc/non disp Oct 19/23) 02011956 ALL ACDEFGV  
 Sandoz Polytrimethoprim 02239234 SDZ ACDEFGV

**S01AD ANTIVIRALS**

S01AD02 TRIFLURIDINE  
 Liq Oph 1% Viroptic 00687456 BSL ACDEFGV

**S01AE FLUOROQUINOLONES**

S01AE01 OFLOXACIN  
 Liq Oph 0.3% Ocuflox 02143291 ABV (SA)

S01AE03 CIPROFLOXACIN  
 Liq Oph 0.3% Ciloxan 01945270 NVR (SA)  
 Sandoz Ciprofloxacin 02387131 SDZ (SA)

Ont Oph 0.3% Ciloxan 02200864 NVR (SA)

S01AE06 GATIFLOXACIN  
 Liq Oph 0.3% Zymar 02257270 ABV ACDEFGV  
 Apo-Gatifloxacin 02327260 APX ACDEFGV

S01AE07 MOXIFLOXACIN  
 Liq Oph 0.5% Vigamox 02252260 NVR ACDEFGV  
 Apo-Moxifloxacin 02406373 APX ACDEFGV  
 Jamp-Moxifloxacin 02472120 JPC ACDEFGV  
 Moxifloxacin 02529076 SAS ACDEFGV  
 pms-Moxifloxacin 02432218 PMS ACDEFGV  
 Sandoz Moxifloxacin 02411520 SDZ ACDEFGV

**S01B ANTIINFLAMMATORY AGENTS**

**S01BA CORTICOSTEROIDS, PLAIN**

S01BA01 DEXAMETHASONE  
 Dps Oph 0.1% Maxidex 00042560 NVR ACDEFGV

S01BA01	DEXAMETHASONE							
	Ont	Oph 0.1%		Maxidex	00042579	NVR	ACDEFGV	
S01BA04	PREDNISOLONE							
	Sus	Oph 1%		Pred Forte	00301175	ABV	ACDEFGV	
				Sandoz Prednisolone	01916203	SDZ	ACDEFGV	
				Teva-Prednisolone	00700401	TEV	ACDEFGV	
S01BA07	FLUOROMETHOLONE							
	Dps	Oph 0.1%		FML	00247855	ABV	ACDEFGV	
				Sandoz Fluorometholone	00432814	SDZ	ACDEFGV	
	Sus	Oph 0.1%		Flarex	00756784	NVR	ACDEFGV	
<b>S01BC</b>	<b>ANTIINFLAMMATORY AGENTS, NON STEROIDS</b>							
S01BC03	DICLOFENAC							
	Liq	Oph 0.1%		Voltaren	01940414	NVR	ACDEFGV	
				Apo-Diclofenac	02441020	APX	ACDEFGV	
				Diclofenac	02475065	PST	ACDEFGV	
				Mint-Diclofenac	02475197	MNT	ACDEFGV	
				Sandoz Diclofenac Ophtha	02454807	SDZ	ACDEFGV	
S01BC05	KETOROLAC							
	Liq	Oph 0.45%		Acuvail	02369362	ABV	ACDEFGV	
	Liq	Oph 0.5%		Acular	01968300	ABV	ACDEFGV	
				Ketorolac	02245821	AAP	ACDEFGV	
<b>S01C</b>	<b>ANTIINFLAMMATORY AGENTS &amp; ANTIINFECTIVES IN COMBINATION</b>							
<b>S01CA</b>	<b>CORTICOSTEROIDS AND ANTIINFECTIVES IN COMBINATION</b>							
S01CA01	DEXAMETHASONE AND ANTIINFECTIVES							
		DEXAMETHASONE / NEOMYCIN / POLYMYXIN B						
	Ont	Oph 1 mg / 3.5 mg / 6 000 IU		Maxitrol	00358177	NVR	ACDEFGV	
	Sus	Oph 1 mg / 3.5 mg / 6 000 IU		Maxitrol	00042676	NVR	ACDEFGV	
		DEXAMETHASONE / TOBRAMYCIN						
	Ont	Oph 0.1% / 0.3%		Tobradex	00778915	NVR	ACDEFGV	
	Sus	Oph 0.1% / 0.3%		Tobradex	00778907	NVR	ACDEFGV	
<b>S01E</b>	<b>ANTIGLAUCOMA PREPARATIONS AND MIOTICS</b>							

**S01EA SYMPATHOMIMETICS IN GLAUCOMA THERAPY**

S01EA03 APRACLONIDINE

Liq Oph 0.5%

Iopidine 02076306 NVR ACDEFGV

S01EA05 BRIMONIDINE

Liq Oph 0.15%

Alphagan P 02248151 ABV ACDEFGV

Brimonidine P 02301334 AAP ACDEFGV

Liq Oph 0.2%

Alphagan 02236876 ABV ACDEFGV

Brimonidine Tartrate 02515377 HIK ACDEFGV

Jamp-Brimonidine 02449226 JPC ACDEFGV

Med-Brimonidine 02507811 GMP ACDEFGV

pms-Brimonidine 02246284 PMS ACDEFGV

Sandoz Brimonidine 02305429 SDZ ACDEFGV

**S01EB PARASYMPATHOMIMETICS**

S01EB01 PILOCARPINE

Dps Oph 2%

Isopto Carpine 00000868 NVR ACDEFGV

**S01EC CARBONIC ANHYDRASE INHIBITORS**

S01EC01 ACETAZOLAMIDE

Tab Orl 250 mg

Acetazolamide 00545015 AAP ACDEFGV

S01EC03 DORZOLAMIDE

Liq Oph 2%

Trusopt 02216205 ELV ACDEFGV

Jamp-Dorzolamide 02453347 JPC ACDEFGV

Med-Dorzolamide 02457210 GMP ACDEFGV

Sandoz Dorzolamide 02316307 SDZ ACDEFGV

S01EC04 BRINZOLAMIDE

Liq Oph 1%

Azopt 02238873 NVR ACDEFGV

S01EC05 METHAZOLAMIDE

Tab Orl 50 mg

Methazolamide 02245882 AAP ACDEFGV

S01EC54 BRINZOLAMIDE, COMBINATIONS

BRINZOLAMIDE / BRIMONIDINE

Liq Oph 1% / 0.2%

Simbrinza 02435411 NVR ACDEFGV

**S01ED BETA BLOCKING AGENTS**

S01ED01 TIMOLOL

Dps Oph 0.25%

Sandoz Timolol Maleate 02166712 SDZ ACDEFGV



## S01ED01 TIMOLOL

Dps Oph 0.5%

Timoptic Oph 00451207 ELV ACDEFGV

Apo-Timop 00755834 APX ACDEFGV

Jamp-Timolol 02447800 JPC ACDEFGV

Sandoz Timolol Maleate 02166720 SDZ ACDEFGV

Timo-Stulln (Temporary Benefit) 09858120 PST ACDEFGV

Liq Oph 0.25%

Timolol Maleate-EX 02242275 SDZ ACDEFGV

Liq Oph 0.5%

Timoptic-XE Oph 02171899 ELV ACDEFGV

Timolol Maleate-EX 02242276 SDZ ACDEFGV

## S01ED02 BETAXOLOL

Sus Oph 0.25%

Betoptic S 01908448 NVR ACDEFGV

## S01ED51 TIMOLOL COMBINATIONS

## TIMOLOL / BRIMONIDINE

Liq Oph 0.5% / 0.2%

Combigan 02248347 ABV ACDEFGV

Apo-Brimonidine-Timop 02375311 APX ACDEFGV

Jamp Brimonidine/Timolol 02531704 JPC ACDEFGV

## TIMOLOL / BRINZOLAMIDE

Sus Oph 0.5% / 1%

Azarga 02331624 NVR ACDEFGV

## TIMOLOL / DORZOLAMIDE

Liq Oph 0.5% / 2%

Cosopt 02240113 ELV ACDEFGV

Cosopt PF 02258692 ELV ACDEFGV

Apo-Dorzo-Timop 02299615 APX ACDEFGV

Dorzolamide and Timolol 02489635 HIK ACDEFGV

Dorzolamide-Timolol 02522020 JPC ACDEFGV

Jamp-Dorzolamide-Timolol 02457539 JPC ACDEFGV

Med-Dorzolamide-Timolol 02437686 GMP ACDEFGV

Riva-Dorzolamide/Timolol 02441659 RIV ACDEFGV

Sandoz Dorzolamide/Timolol 02344351 SDZ ACDEFGV

## TIMOLOL / LATANOPROST

S01ED51 TIMOLOL COMBINATIONS  
TIMOLOL / LATANOPROST

Liq	Oph	0.5% / 0.005%	Xalacom	02246619	BGP	ACDEFGV
			Act Latanoprost/Timolol	02436256	TEV	ACDEFGV
			GD-Latanoprost/Timolol	02373068	UJC	ACDEFGV
			Jamp-Latanoprost-Timolol	02453770	JPC	ACDEFGV
			Latanoprost and Timolol Ophthalmic (Disc/non disp Aug 8/24)	02489368	HIK	ACDEFGV
			M-Latanoprost-Timolol	02514516	MRA	ACDEFGV
			Med-Latanoprost-Timolol	02454505	GMP	ACDEFGV

TIMOLOL / TRAVOPROST

Liq	Oph	0.5% / 0.004%	Duo Trav PQ	02278251	NVR	ACDEFGV
			Apo-Travoprost-Timop	02415305	APX	ACDEFGV

**S01EE PROSTAGLANDIN ANALOGUES**

S01EE01 LATANOPROST

Liq	Oph	0.005%	Xalatan	02231493	UJC	ACDEFGV
			Apo-Latanoprost	02296527	APX	ACDEFGV
			GD-Latanoprost	02373041	UJC	ACDEFGV
			Jamp-Latanoprost	02453355	JPC	ACDEFGV
			Latanoprost Ophthalmic Solution	02489570	HIK	ACDEFGV
			M-Latanoprost	02513285	MRA	ACDEFGV
			Med-Latanoprost	02426935	GMP	ACDEFGV
			pms-Latanoprost	02317125	PMS	ACDEFGV
			Riva-Latanopost	02341085	RIV	ACDEFGV
			Sandoz Latanoprost	02367335	SDZ	ACDEFGV
			Teva-Latanoprost	02254786	TEV	ACDEFGV

S01EE03 BIMATOPROST

Liq	Oph	0.01%	Lumigan RC	02324997	ABV	ACDEFGV
Liq	Oph	0.03%	Vistitan	02429063	SDZ	ACDEFGV

S01EE04 TRAVOPROST

Liq	Oph	0.003%	Izba	02457997	NVR	ACDEFGV
Liq	Oph	0.004%	Travatan Z	02318008	NVR	ACDEFGV
			Apo-Travoprost Z	02415739	APX	ACDEFGV
			Sandoz Travoprost	02413167	SDZ	ACDEFGV

S01EE06 LATANOPROSTENE BUNOD

S01EE06 LATANOPROSTENE BUNOD

Liq Oph 0.024%

Vyzulta 02484218 BSH ACDEFGV

**S01F MYDRIATICS AND CYCLOPLEGICS**

**S01FA ANTICHOLINERGICS**

S01FA01 ATROPINE

Dps Oph 1%

Isopto Atropine 00035017 ALC ACDEFGVW

Atropine 02023695 PST ACDEFGVW

S01FA04 CYCLOPENTOLATE

Liq Oph 1%

Cyclogyl 00252506 ALC ACDEFGV

S01FA06 TROPICAMIDE

Liq Oph 0.5%

Mydriacyl 00000981 ALC ACDEFGV

Liq Oph 1%

Mydriacyl 00001007 ALC ACDEFGV

**S01G DECONGESTANTS AND ANTIALLERGICS**

**S01GX OTHER ANTIALLERGICS**

S01GX01 CROMOGLICIC ACID

Liq Oph 2%

Cromolyn Ophthalmic Solution 02009277 PDP ACDEFGV

S01GX08 KETOTIFEN

Liq Oph 0.025%

Zaditor 02242324 LTH ACDEFGV

S01GX09 OLOPATADINE

Liq Oph 0.1%

Patanol 02233143 NVR ACDEFGV

Apo-Olopatadine 02305054 APX ACDEFGV

Jamp-Olopatadine 02458411 JPC ACDEFGV

Sandoz Olopatadine 02358913 SDZ ACDEFGV

Liq Oph 0.2%

Pataday 02362171 NVR ACDEFGV

Apo-Olopatadine 02402823 APX ACDEFGV

Mint-Olopatadine 02508605 MNT ACDEFGV

Sandoz Olopatadine 02420171 SDZ ACDEFGV

**S01L OCULAR VASCULAR DISORDER AGENTS**

**S01LA ANTINEOVASCULARISATION AGENTS**

S01LA04 RANIBIZUMAB

Liq IVL 10 mg/mL

Byoovis 02525852 BIG (SA)

S01LA05 AFLIBERCEPT

S01LA05	AFLIBERCEPT								
Liq	IVL	40 mg/mL			Eylea	02415992	BAY	(SA)	
S01LA06	BROLUCIZUMAB								
Liq	IVL	6 mg / 0.05 mL			Beovu	02496976	NVR	(SA)	
S01LA09	FARICIMAB								
Liq	IVL	6 mg / 0.05 mL			Vabysmo	02527618	HLR	(SA)	
<b>S01X</b>	<b>OTHER OPHTHALMOLOGICALS</b>								
<b>S01XA</b>	<b>OTHER OPHTHALMOLOGICALS</b>								
S01XA03	SODIUM CHLORIDE, HYPERTONIC								
Dps	Oph	5%			Muro 128	00750824	BSH	AEFGV	
					Odan-Sodium Chloride	80046737	ODN	AEFGV	
Ont	Oph	5%			Muro 128	00750816	BSH	AEFGV	
					Odan-Sodium Chloride	80046696	ODN	AEFGV	
S01XA18	CICLOSPORIN								
Eml	Oph	0.1%			Verkazia	02484137	SNN	(SA)	
S01XA21	MERCAPTAMINE (CYSTEAMINE)								
Liq	Oph	0.37%			Cystadrops	02485605	RRD	(SA)	
<b>S02</b>	<b>OTOLOGICALS</b>								
<b>S02C</b>	<b>CORTICOSTEROIDS AND ANTIINFECTIVES IN COMBINATION</b>								
<b>S02CA</b>	<b>CORTICOSTEROIDS AND ANTIINFECTIVES IN COMBINATION</b>								
S02CA02	FLUMETASONE AND ANTIINFECTIVES								
	FLUMETASONE / CLIOQUINOL								
Dps	Ot	0.2% / 1%			Locacorten-Vioform	00074454	PAL	ACDEFGV	
S02CA06	DEXAMETHASONE AND ANTIINFECTIVES								
	DEXAMETHASONE / CIPROFLOXACIN								
Sus	Ot	0.1% / 0.3%			Ciprodex	02252716	NVR	ACDEFGV	
					Sandoz Ciprofloxacin/Dexamethasone	02506882	SDZ	ACDEFGV	
					Taro-Ciprofloxacin/Dexamethasone	02481901	TAR	ACDEFGV	
<b>S03</b>	<b>OPHTHALMOLOGICAL AND OTOLOGICAL PREPARATIONS</b>								
<b>S03C</b>	<b>CORTICOSTEROIDS AND ANTIINFECTIVES IN COMBINATION</b>								
<b>S03CA</b>	<b>CORTICOSTEROIDS AND ANTIINFECTIVES IN COMBINATION</b>								
S03CA01	DEXAMETHASONE AND ANTIINFECTIVES								
	DEXAMETHASONE / FRAMYCETIN / GRAMICIDIN								

S03CA01	DEXAMETHASONE AND ANTIINFECTIVES						
	DEXAMETHASONE / FRAMYCETIN / GRAMICIDIN						
Dps	Oph	0.5 mg / 5 mg / 0.05 mg		Sofracort E/E	02224623	SAV	ACDEFGV
<b>V</b>	<b>VARIOUS</b>						
<b>V01</b>	<b>ALLERGENS</b>						
<b>V01A</b>	<b>ALLERGENS</b>						
<b>V01AA</b>	<b>ALLERGEN EXTRACTS</b>						
V01AA02	GRASS POLLEN						
Kit	SC	105, 250, 700, 2 150 PNU		Pollinex-R	00464988	PAL	(SA)
Slit	Orl	100 IR		Oralair	02381885	STA	(SA)
Slit	Orl	300 IR		Oralair	02381893	PAL	(SA)
V01AA20	VARIOUS ALLERGEN EXTRACTS						
Liq	Inj			Allergy Sera	00999938	HJM	EF-18G
<b>V03</b>	<b>ALL OTHER THERAPEUTIC PRODUCTS</b>						
<b>V03A</b>	<b>ALL OTHER THERAPEUTIC PRODUCTS</b>						
<b>V03AB</b>	<b>ANTIDOTES</b>						
V03AB06	THIOSULFATE						
	SODIUM THIOSULFATE						
Liq	Inj	250 mg/mL		Seacalphyx	02386666	SFD	ACDEFGVW
<b>V03AC</b>	<b>IRON CHELATING AGENTS</b>						
V03AC01	DEFEROXAMINE						
Pws	Inj	500 mg		Desferal	01981242	NVR	ACDEFGV
				Deferoxamine Mesilate	02241600	PFI	ACDEFGV
Pws	Inj	2 g		Deferoxamine Mesilate	02247022	PFI	ACDEFGV
V03AC02	DEFERIPRONE						
Liq	Orl	100 mg/mL		Ferriprox	02436523	APO	(SA)
Tab	Orl	1 000 mg		Ferriprox	02436558	CCC	(SA)
V03AC03	DEFERASIROX						

V03AC03 DEFERASIROX

Tab	Orl	90 mg	Jadenu	02452219	NVR	(SA)
			Apo-Deferasirox (Type J)	02485265	APX	(SA)
			pms-Deferasirox (Type J)	02528290	PMS	(SA)
			Sandoz Deferasirox (Type J)	02489899	SDZ	(SA)
			Taro-Deferasirox (Type J)	02507315	TAR	(SA)
Tab	Orl	180 mg	Jadenu	02452227	NVR	(SA)
			Apo-Deferasirox (Type J)	02485273	APX	(SA)
			pms-Deferasirox (Type J)	02528304	PMS	(SA)
			Sandoz Deferasirox (Type J)	02489902	SDZ	(SA)
			Taro-Deferasirox (Type J)	02507323	TAR	(SA)
Tab	Orl	360 mg	Jadenu	02452235	NVR	(SA)
			Apo-Deferasirox (Type J)	02485281	APX	(SA)
			pms-Deferasirox (Type J)	02528312	PMS	(SA)
			Sandoz Deferasirox (Type J)	02489910	SDZ	(SA)
			Taro-Deferasirox (Type J)	02507331	TAR	(SA)

**V03AE FOR TREATMENT OF HYPERKALEMIA AND HYPERPHOSPHATEMIA**

V03AE01 POLYSTYRENE SULFONATE

CALCIUM POLYSTYRENE SULFONATE

Pws	Orl	999 mg/g	Resonium Calcium	02017741	SAV	ACDEFGV
			Jamp Calcium Polystyrene Sulfonate	02502631	JPC	ACDEFGV

SODIUM POLYSTYRENE SULONATE

Pws	Orl	1 g/g	Kayexalate	02026961	SAV	ACDEFGV
			Jamp Sodium Polystyrene Sulfonate	02497557	JPC	ACDEFGV
			Odan-Sodium Polystyrene Sulfonate	02473941	ODN	ACDEFGV
			Solystat	00755338	PDP	ACDEFGV
Sus	Orl	250 mg/mL	Odan-Sodium Polystyrene Sulfonate	02473968	ODN	ACDEFGV
			Solystat	00769541	PDP	ACDEFGV

V03AE02 SEVELAMER

Pws.	Orl	0.8 g	Renvela	02485559	SAV	(SA)
Pws.	Orl	2.4 g	Renvela	02485567	SAV	(SA)
Tab	Orl	800 mg	Renagel	02244310	SAV	ACDEFGV
			Accel-Sevelamer	02461501	ACC	ACDEFGV

V03AE03 LANTHANUM CARBONATE

TabC	Orl	250 mg	Fosrenol (Disc/non disp Apr 30/24)	02287145	TAK	(SA)
TabC	Orl	500 mg	Fosrenol	02287153	TAK	(SA)
TabC	Orl	750 mg	Fosrenol	02287161	TAK	(SA)
TabC	Orl	1000 mg	Fosrenol	02287188	TAK	(SA)

V03AE05 SUCROFERRIC OXYHYDROXIDE

TabC	Orl	500 mg	Velphoro	02471574	VFM	(SA)
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**V03AF DETOXIFYING AGENTS FOR ANTINEOPLASTIC TREATMENT**

V03AF01 MESNA

MESNA

Pws	Inj	100 mg/mL	Uromitexan	02241411	BAX	ACDEFGV
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V03AF03 CALCIUM FOLINATE

LEUCOVORIN CALCIUM

Tab	Orl	5 mg	Lederle Leucovorin	02170493	PFI	ACDEFGV
			Mint-Leucovorin	02496828	MNT	ACDEFGV
			Riva Leucovorin	02493357	RIV	ACDEFGV

**V03AG DRUGS FOR TREATMENT OF HYPERCALCEMIA**

V03AG99 DRUGS FOR TREATMENT OF HYPERCALCEMIA

SODIUM ACID PHOSPHATE / SODIUM BICARBONATE / POTASSIUM

Evt	Orl	500 mg / 469 mg / 123 mg	Jamp-Sodium Phosphate	80047562	JPC	ACDEFGV
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**V03AH FOR TREATMENT OF HYPOGLYCEMIA**

V03AH01 DIAZOXIDE

Cap	Orl	100 mg	Proglycem	00503347	FRS	ACDEFGV
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**V04 DIAGNOSTIC AGENTS**

**V04C OTHER DIAGNOSTIC AGENTS**

**V04CJ TESTS FOR THYREOIDEA FUNCTION**

V04CJ01 THYROTROPIN

Pws	IM	0.9 mg	Thyrogen	02246016	GZM	(SA)
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**APPENDIX I-A / ANNEXE I-A**

**ABBREVIATIONS OF DOSAGE FORMS / ABRÉVIATIONS DES FORMES POSOLOGIQUES**

FORM	CODE	FORME
Metered-Dose Aerosol	Aem/Aém.	Aérosol-dose mesurée
Aerosol (with propellants)	Aer/Aér.	Aérosol (avec agents de propulsion)
Aerosol (without propellants)	Asp	Aérosol (sans agents de propulsion)
Capsule	Cap/Caps	Capsule
Chewable Tablets	TabC/Co.C.	Comprimés à croquer
Controlled Delivery Capsules	CDC/Caps.L.C.	Capsules à libération contrôlée
Cleanser	Clr/Net	Nettoyant
Cream	Crm/Cr.	Crème
Cartridge	Ctg/Cart	Cartouche
Douche	Dch	Douche
Delayed Action (Injectables)	Dla	Soluté injectable-retard
Delayed Release Capsule	CDR/Caps.L.R.	Capsule à libération retardée
Drop	Dps/Gttes	Gouttes
Dressing	Dre	Pansement
Enteric Coated Capsule	ECC/Caps.Ent.	Capsule entérique
Each	Ech/Ch	Chacun
Enteric Coated Granule	Ecg	Granule entérique
Enteric Coated Tablet	ECT/Co.Ent	Comprimés entérique
Elixir	Elx	Élixir
Emulsion	Em/Émuls	Émulsion
Enema	Enm/Lav.	Lavement
Extended Release	ER	À libération prolongée
Extended Release Capsules	ERC/Caps.L.P.	Capsules à libération prolongée
Extended Release Tablets	ERT/Co.L.P.	Comprimés à libération prolongée
Effervescent Granule	Evg/Gev	Granule effervescente
Effervescent Powder	Ecp/Pev	Poudre effervescente
Effervescent Tablet	Evt/Co.Eff.	Comprimé effervescent
Film Coated	FC	Pelliculés
Gas	Gas	Gaz
Gel	Gel	Gelée
Granules	Gran	Granules
Immediate release	IR	Libération immédiate



**APPENDIX I-A / ANNEXE I-A**

**ABBREVIATIONS OF DOSAGE FORMS / ABRÉVIATIONS DES FORMES POSOLOGIQUES**

FORM	CODE	FORME
Inhaler	Inh	Inhalateur
Instrument	Ins	Pièce à insérer
Insulin	Ins	Insuline
Intra-Articular	IA	Intra-articulaire
Kit	Kit/Tro	Trousse
Liniment	Lin	Liniment
Liquid	Liq	Liquide
Lente Suspension	Lla/Susp.	Suspension
Lotion	Lot	Lotion
Lozenge	Loz/Pas	Pastille
Miscellaneous	Misc	Divers
Mist, Aerosol	Mst/Baer	Bruine en aérosol
Mouthwash	MWH/R.-B.	rince-bouche
Nebules	Neb	Nébules
Orally Disintegrating Film	ODF	Film à désintégration orale
Orally Disintegrating Tablet	ODT/Co.D.O.	Comprimés à désintégration orale
Oral liquid	O/L	Liquide Oral
Ointment	Ont	Onguent, pomade
Pad	Pad/Gaze	Compresse
Package	Pck	Paquet
Paste	Pst	Pâte
Patch	Pth	Timbre cutané
Preservative Free	PF	Sans agent de conservation
Powder	Pwr/Pd.	Poudre
Powder for Solution / Powder for Suspension	Pws/Pds.	Poudre pour solution / Poudre pour suspension
Rapid Dissolving	RD	Dissolution rapide
Rapid Disintegrating	RPD	Désintégration rapide
Shampoo	Shp	Shampooing
Semi-Lente Suspension	SLA	Suspension semi-lente
Slow release	SR	Libération lente
Sublingual Tablet	SlT/Co.S.L.	Comprimé sublingual

**APPENDIX I-A / ANNEXE I-A**

**ABBREVIATIONS OF DOSAGE FORMS / ABRÉVIATIONS DES FORMES POSOLOGIQUES**

FORM	CODE	FORME
Spray	Spr/Vap	Vaporisateur
Sustained-Released Capsule	SRC/Caps.L.L.	Capsule à libération lente
Packet	Packet/Sachets	Sachet/Paquet,
Sustained-Release Disc	Srd	Disque à action soutenue
Sustained-Release Syrup	SRS	Sirop à action soutenue
Sustained-Release Tablet	SRT/Co.L.L.	Comprimé à libération lente
Suppository	Sup/Supp.	Suppositoire
Suspension	Susp/Susp	Suspension
Syrup	Syr/Sir.	Sirop
Tablet	Tab/Co.	Comprimé
Ultra-Lente Suspension	Ula	Suspension ultra-lente
Wafer	Waf	Gaufrette

**APPENDIX I-B/ ANNEXE I-B**

**ABBREVIATIONS OF ROUTES / ABRÉVIATIONS DES VOIES D'ADMINISTRATION**

ROUTE	CODE	VOIE
Buccal	Buc	Buccale, orale
Dental	Den	Dentaire
Inhalation	Inh	Inhalation
Injectable	Inj	Injectable
Instillation	ISL	Instillation
Instrument(s)	Ins	Instrument(s)
Intervertebral	IND	Intervertébrale
Intra Articular	IA	Intra-articulaire
Intrabursal	IBU	Intrabursique
Intracardiac	ICD	Intracardiaque
Intracavity	ICV	Intra-cavitaire
Intradermal	ID	Intradermique
Intrafollicular	INF	Intra-folliculaire
Intraintestinal	ITT	Intraintestinale
Intramuscular	IM	Intramusculaire
Intraocular	IO	Intraoculaire
Intraperitoneal	IP	Intrapéritonéale
Intrapleural	IPL	Intrapleurale
Intrapulmonary	IPU	Intrapulmonaire
Intrathecal	INT	Intra-thécale
Intravenous	IV	Intraveineuse
Intraventricular	IVR	intraventriculaire
Intravesicular	ITV	Intravésicale
Intravitreal	IVL	Intravitréenne
Irrigation	IR	Irrigation
Miscellaneous	Mis	Divers
Nasal	Nas	Nasale
Nil	NIL	Néant
Ophthalmic	Oph	Ophtalmique
Oral	Orl	Orale
Otic	Ot	Otique

APPENDIX I-B/ ANNEXE I-B

ABBREVIATIONS OF ROUTES / ABRÉVIATIONS DES VOIES D'ADMINISTRATION

ROUTE	CODE	VOIE
Retrobulbar	RB	Rétrobulbaire
Rectal	Rt	Rectale
Subcutaneous	SC	Sous-cutané
Sublingual	Slg	Sublinguale
Topical	Top	Topique
Transdermal	Trd	Transdermique
Vaginal	Vag	Vaginale

**APPENDIX I-C/ ANNEXE I-C**

**ABBREVIATIONS OF UNITS / ABRÉVIATIONS DES UNITÉS DE MESURE**

UNIT	CODE	UNITÉS
Ampoule	Amp	Ampoule
Billion	B	Milliard
Bottle	Bottl	Flacon, bouteille
Box	Box	Boîte
Capsule	Cap	Capsule
Cubic Centimetre	CC	Centimètre cube
Centimetre	cm	Centimètre
Disk	Disk	Disque
Fluid Ounce	Fl oz	Once liquid
Gallon	Gal	Gallon
Gram	g	Gramme
Grain	Gr	Grain
Kilogram	kg	Kilogramme
Kit	Kit/Tro	Trousse
Litre	L	Litre
Pound	lb	Livre
Lozenge	Loz/Pas	Pastille
Million	M	Million
Microgram	mcg	Microgramme
Milli-equivalent	mEq	Milli-équivalent
Milligram	mg	Milligramme
Drop	dps/gttes	Goutte
Millitre	mL	Millilitre
Millimole	Mmol	Millimole
Nil	Nil	Néant
Ounce	oz	Once
Package	Pcg	Paquet, emballage
Syringe	SYR	Seringue
Tablet	Tab/Co.	Comprimé
Tablespoon	Tbs	Cuillerée à soupe
Trace	Trace	Trace

APPENDIX I-C/ ANNEXE I-C

ABBREVIATIONS OF UNITS / ABRÉVIATIONS DES UNITÉS DE MESURE

UNIT	CODE	UNITÉS
Teaspoon	Tsp	Cuillerée à thé
Tube	Tube	Tube
International Unit	IU	Unité international
Vial	Vial	Fiole
By Weight	w/w	En poids

**APPENDIX I-D / ANNEXE I-D**

**ABBREVIATIONS OF MANUFACTURER'S NAMES/ABRÉVIATIONS DES NOMS DE FABRICANTS**

AAP	AA Pharma Inc.	FRE	Fresenius Medical Care Canada
ABB	Abbott Laboratories, Ltd.	FRS	Merck Canada Inc.
ABV	Abbvie Corporation	GAC	Galderma Canada Inc.
ACC	Accel Pharma	GCH	GlaxoSmithKline Consumer Healthcare Inc.
ACT	Actelion Pharmaceuticals Canada Inc.	GIL	Gilead Sciences Inc.
ADZ	Advanz Pharma Canada Inc.	GLM	Glenmark Pharmaceuticals Canada Inc.
AGA	Amgen Canada Inc.	GMD	GenMed, a division of Pfizer Canada Inc.
AHC	Athena Canada Inc.	GMP	Generic Medical Partners
AHI	Accord Healthcare Inc.	GSK	GlaxoSmithKline
AKT	Akcea Therapeutics Inc.	GZM	Genzyme- A Division of Sanofi-Aventis
ALC	Alcon Canada Inc.	HIK	Hikma Canada Ltd.
ALL	Allergan Inc.	HJM	Medavie Blue Cross
ALM	Almirall Canada Ltd.	HLR	Hoffmann-La Roche Ltd/Ltee.
ALN	Alnylam Netherlands B.V.	HLS	HLS Therapeutics Inc.
ALX	Alexion Pharma	HLZ	Hill Dermaceuticals Inc.
ALY	Amylyx Canada Inc.	HOS	Hospira Healthcare Corporation
AMT	Amicus Therapeutics UK Ltd.	HRZ	Horizon Pharma Ireland Ltd.
APN	Aspen Pharmacare Canada Inc.	INP	Insight Pharmaceuticals Corp.
APO	ApoPharma Inc.	IPS	Ipsen Biopharmaceuticals
APX	Apotex Inc.	IUK	Indivior UK Limited
ARN	Accelera Pharma Canada Inc.	JAM	Jamieson Laboratories Ltd.
ARO	Auro Pharma Inc.	JAN	Janssen Inc.
ARZ	Aralez Pharmaceuticals Canada Inc.	JCB	Jacobus Pharmaceutical Company Inc.
ASL	Astellas Pharma Canada Inc.	JNJ	Johnson & Johnson Consumer Group
ATL	Laboratoire Atlas Inc.	JNO	Juno Pharmaceuticals Corp
ATS	Altius Healthcare Inc.	JPC	Jamp Pharma Corporation
ATV	Actavis Pharma Company	KEG	Kego Corporation
AVI	Avir Pharma Inc.	KLO	Kaleo Inc.
AXC	Aptalis	KNI	Knight Therapeutics Inc.
AZE	AstraZeneca Canada Inc.	KVR	KVR Pharmaceuticals Inc.
BAX	Baxter Corporation	KYE	Kye Pharmaceuticals Inc.
BAY	Bayer Inc., HealthCare Division	LBI	Leadiant Biosciences Inc.
BGN	BeiGene (Canada) ULC	LBK	Lundbeck Inc.
BGP	BGP Pharma Inc.	LDN	Leadiant Biosciences Inc.
BIG	Biogen Idec Canada, Inc.	LEO	Leo Pharma Inc.
BMR	Biomarin Pharmaceuticals Canada	LIL	Eli Lilly Canada Inc.
BOE	Boehringer Ingelheim (Canada) Ltd.	LIN	Linepharma International Inc.
BOX	Biocodex SA	LTH	Labtician Thea
BRI	Bristol-Myers Squibb Canada Inc.	LUP	Lupin Pharma Canada Ltd.
BSH	Bausch & Lomb Canada Inc.	MAR	Marcan Pharmaceuticals Inc
BSL	Bausch Health Canada Inc.	MBT	Mitsubishi Tanabe Pharma Corporation
BVT	Swedish Orphan Biovitrum AB	MCK	Mckesson Canada Corp.
CBP	Cubist Pharmaceuticals Inc.	MDI	Medtech Products Inc.
CCC	Chiesi Canada Corp	MDK	MendeliKABS Inc.
CCM	CellChem Pharmaceuticals Inc.	MDU	Medunik Canada
CEL	Celgene	MDX	Medexus Inc.
CHC	Pfizer Canada Inc., Consumer Healthcare	MJO	Mead Johnson Canada
CHU	Church and Dwight Canada Corp.	MNT	Mint Pharmaceuticals Inc.
CIP	Cipher Pharmaceuticals Inc.	MRA	Mantra Pharma
CLC	Columbia Laboratories Canada Inc.	MRZ	Merz Pharmaceuticals Canada Ltd.
CPC	Covis Pharma Canada Ltd.	MSD	MSD Inc.
CTL	Celltrion Healthcare Co., Ltd.	MTP	Methapharm Inc.
DPT	Dermtek Pharmaceuticals Ltd	MYL	Mylan Pharmaceuticals ULC
DUI	Duchesnay	NAT	Natco Pharma (Canada) Inc.
EDO	Endo Ventures Ltd.	NHI	Nic-Hit International Inc.
EIS	Eisai Limited	NNO	Novo Nordisk Canada Inc.
ELV	Elvium Life Science- A Purdue Company	NRA	Nora Phama Inc.
EMD	EMD Serono Canada Inc.	NUT	Nutricorp International
ERF	Erfa Canada Inc.	NVR	Novartis Pharmaceuticals Canada Inc.
ETH	Ethypharm Inc.	ODN	Odan Laboratories Ltd.
EXZ	Exzell Pharma Inc.	OMG	Omega Laboratories Limited
FEI	Ferring Inc.	ORG	Organon Canada Inc.
FKB	Fresenius Kabi Canada Ltd.	ORI	Orimed Pharma Corporation

**APPENDIX I-D / ANNEXE I-D**

**ABBREVIATIONS OF MANUFACTURER'S NAMES/ABRÉVIATIONS DES NOMS DE FABRICANTS**

OTS	Otsuka Canada Pharmaceuticals In	WLS	Wellspring Pharmaceutical Canada Corp
PAL	Paladin Labs Inc.	WMD	Waymade Canada Inc.
PCI	Phebra Canada Inc.	WNC	Warner Chilcott Canada Co.
PDL	Pro Doc Laboratories Ltd	WNP	WN Pharmaceuticals Ltd.
PDP	PendoPharm, Division of Pharmascience	XPI	Xediton Pharmaceuticals Inc.
PFB	Pierre Fabre Dermo-Cosmetique		
PFI	Pfizer Canada Inc.		
PFR	Purdue Pharma		
PJC	Pharmacie Jean Coutu.		
PMS	Pharmascience Inc.		
PRZ	Pharmaris Canada Inc.		
PST	Pharma Stulln Inc.		
PSV	Pharmasave		
RAN	Ranbaxy Pharmaceuticals Canada Inc		
RCH	Dr. Reddy's Laboratories Inc.		
RIV	Riva Laboratories Ltee		
ROG	Rougier Pharma Inc, Div of Ratiopharm		
RRD	Recordati Rare Diseases Canada Inc		
SAS	Sanis Health Inc.		
SAV	Sanofi-Aventis Canada Inc.		
SAX	Salix Pharmaceuticals Inc.		
SDM	Shoppers Drug Mart		
SDZ	Sandoz Canada Incorporated		
SEV	Servier Canada Inc.		
SFD	Seaford Pharmaceuticals Inc.		
SGC	Seagen Canada Inc.		
SGQ	Sterigen Inc.		
SHI	Shire Canada Inc.		
SIV	Sivem Pharmaceuticals		
SLP	Searchlight Pharma Inc.		
SNC	Sanofi Consumer Health Inc.		
SNE	Smith & Nephew, Inc.		
SNN	Santen Incorporated		
SNV	Sunovion Pharmaceuticals Canada Inc		
SOB	Sobey's Pharmacy		
SPT	Septa Pharmaceuticals Inc.		
STA	Stallergenes Canada Inc.		
STD	Strides Pharma Canada Inc.		
STR	Sterimax Inc.		
SUN	Sun Pharma Canada Inc.		
TAI	Taiho Pharma Canada Inc.		
TAK	Takeda Canada Inc.		
TAR	Taro Pharmaceuticals Inc.		
TEV	Teva Canada Limited		
TLI	Labs Laboratoire Trianon		
TMP	Teva Canada Innovation		
TOL	Tolmar International Ltd.		
TPH	TaroPharma, Divison of Taro Pharmaceuticals		
UCB	UCB Canada Inc.		
UJC	Upjohn Canada ULC		
UGX	Ultragenyx Canada Inc.		
UTC	United Therapeutics Corporation		
VFM	Vifor Fresenius Medical Care Renal Pharma Ltd.		
VIV	ViiV Healthcare ULC		
VLH	Lundbeck Canada Inc.		
VRT	Verity Pharmaceuticals		
VTH	Vita Health Company (1985) Ltd		
VTX	Vertex Pharmaceuticals (Canada) Inc.		
VVS	Vivus Inc.		
WAL	Walmart Pharmacy		
WAM	Wampole Brands		



## APPENDIX II

### Extemporaneous Preparations (Compounds)

An extemporaneous preparation (compound) is a drug or mixture of drugs prepared or compounded in a pharmacy according to the order of a prescriber.

#### Eligible Benefits

To be eligible as a benefit, a compound must meet one of the following criteria:

1. Contains one or more regular\* benefit drugs
2. Contains one or more special authorization drugs for which approval has been granted
3. Contains a combination of regular\* benefit drugs and special authorization drugs for which approval has been granted
4. Is a compound that has been approved through special authorization

\*Regular benefits include drugs listed on the NB Drug Plans Formulary that do not require special authorization, and the drugs and ingredients used in compounds that are listed below.

#### Non Benefits

A compound is not an eligible benefit if any of the following apply:

1. An alternative is commercially available
2. Contains a drug or product on the exclusion list
3. Made using a proprietary recipe with an undisclosed ingredient list
4. Contains a non-benefit form of a drug (e.g. using powder vs. tablets) unless special authorization approval has been granted
5. Custom-compounded bioidentical hormones

Note: Any drug or product manipulated in accordance with its direction of use (e.g. mixing, reconstituting, prefilling syringes, filling infusion pump reservoirs) is not considered an extemporaneous preparation.

#### Product Shortages

When there is a shortage or no supply of a commercially available product and the healthcare professional has determined a medical need for this product, the product may be compounded during the period of shortage or no supply only. (Health Products and Food Branch Inspectorate Policy on Manufacturing and Compounding Drug Products in Canada, January 26, 2009)

#### Regular Benefit Compounds

<b>Product / Ingredient</b>	<b>PIN</b>	<b>Plans</b>
Anthrakin powder in compounds for topical application	00901113	ACDEFGV
Disulfiram powder	00999087	ACDEFG
Hydrochlorothiazide powders and suspensions for oral use	00999106	ACDEFGV
Hydrocortisone powder for topical applications >0.5%	00990841	ACDEFGV
LCD (Coal Tar Solution) in compounds for topical applications	00358495	ACDEFGV
Meclizine powder	00903076	ACDEFGV
Methoxsalen powder	00903588	ACDEFGVW
Prednisone powders and suspension for oral use	00999108	ACDEFGV
Salicylic Acid in compounds for topical applications	00900788	ACDEFGV
Saturated Solution Potassium Iodide	00999105	ACDEFGV
Spirolactone powders and suspensions for oral use	00999107	ACDEFGV
Sulphur in compounds for topical applications	00900826	ACDEFGV

Note: The PIN can be used to submit claims for any strength of the extemporaneous preparation.

#### Pharmacy Claims

Information on NB Drug Plans Claim Submissions is available [here](#).

- Claims for compounds are to be submitted electronically using the eligible benefit DIN/PIN of at least one of the ingredients contained in the preparation.
- If a preparation contains both a regular benefit drug(s) and a special authorization drug(s), it must be billed using the DIN of the special authorization drug for which prior approval has been granted.
- Claims must be identified by entering the appropriate CPhA version 3 code.

- Manual claims from beneficiaries (pay and submit) will only be accepted for regular benefit preparations. If the preparation does not contain a regular benefit drug, the claim cannot be processed unless special authorization has been granted.
- If a participating provider does not submit an electronic claim for payment and provides a receipt to a beneficiary for a manual (pay and submit) claim, the participating provider must not charge an amount that is greater than the amount that would be paid if the claim was submitted electronically.

#### **Pharmacy Provider Audits**

- Payments made for compounds are subject to audit and recovery.
- Compound Review Verification letters requesting documentation, may be sent to providers to confirm the ingredients contained in the compound and the acquisition cost of each ingredient.
- Although a claim with an eligible benefit DIN/PIN may be accepted electronically, if it contains a drug considered a non-benefit it is subject to recovery.

## APPENDIX III

### New Brunswick Drug Plans Special Authorization Criteria

#### **ABATACEPT (ORENCIA) 250 mg / 15 mL vial**

##### **Polyarticular Juvenile Idiopathic Arthritis**

For the treatment of children (age 6-17) with moderately to severely active polyarticular juvenile idiopathic arthritis (pJIA) who are intolerant to, or who have not had an adequate response from etanercept.

##### Claim Notes:

- Must be prescribed by a rheumatologist.
- Abatacept will not be reimbursed in combination with anti-TNF agents.
- Combined use of more than one biologic drug will not be reimbursed.
- Intravenous infusion: initial IV infusion dose is administered at 0, 2, and 4 weeks then every 4 weeks thereafter.
- Initial treatment is limited to a maximum of 16 weeks. Retreatment is permitted for children who demonstrated an adequate initial treatment response and who are experiencing a disease flare.

#### **ABATACEPT (ORENCIA) 250 mg / 15 mL vial and 125 mg/mL prefilled syringe**

##### **Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

##### Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

##### Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Intravenous infusion: 500mg for patients less than 60 kg, 750mg for patients 60-100 kg and 1000mg for patients greater than 100 kg, given at 0, 2, and 4 weeks then every 4 weeks thereafter.
- Subcutaneous injection: a single IV loading dose of up to 1,000mg may be given, followed by 125mg subcutaneous injection within a day, then once-weekly 125mg subcutaneous injections.
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of continued response is required.

#### **ABIRATERONE (ZYTIGA and generic brands) 250 mg tablet and 500 mg film-coated tablet**

1. In combination with androgen deprivation therapy (ADT) for the treatment of patients with metastatic castration-sensitive prostate cancer (mCSPC) who have had no prior ADT, or are within 6 months of beginning ADT, in the metastatic setting.
2. For the treatment of patients with metastatic castration-resistant prostate cancer.

##### Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

##### Clinical Notes:

1. Patients must have a good performance status.

2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**ABOBOTULINUMTOXINA (DYSPORT THERAPEUTIC)  
300 unit/vial and 500 unit/vial**

1. For the treatment of cervical dystonia (spasmodic torticollis) in adults.
2. For the treatment of upper and lower limb focal spasticity in adults.
3. For the treatment of lower limb spasticity in pediatric patients 2 years of age and older.

**ACALABRUTINIB (CALQUENCE)  
100 mg capsule**

1. As monotherapy for adult patients with previously untreated chronic lymphocytic leukemia (CLL) / small lymphocytic lymphoma (SLL) for whom fludarabine-based treatment is inappropriate due to high-risk cytogenetic markers (i.e., del17p, TP53 mutation, or unmutated IGHV).
2. As monotherapy for adult patients with relapsed or refractory CLL / SLL who have received at least one prior therapy.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will not be considered for patients who experience disease progression on a Bruton's tyrosine kinase (BTK) inhibitor or idelalisib.
- Approval period: 1 year.

**ADALIMUMAB**

**Abrilada 20 mg / 0.4 mL prefilled syringe, 40 mg / 0.8 mL autoinjector and prefilled syringe**  
**Amgevita 20 mg / 0.4 mL prefilled syringe, 40 mg / 0.8 mL autoinjector and prefilled syringe**  
**Hadlima 40 mg / 0.8 mL autoinjector and prefilled syringe**  
**Hulio 20 mg / 0.4 mL prefilled syringe, 40 mg / 0.8 mL autoinjector and prefilled syringe**  
**Hyrimoz 20 mg / 0.4 mL prefilled syringe, 40 mg / 0.8 mL autoinjector and prefilled syringe**  
**Idacio 40 mg / 0.8 mL autoinjector**  
**Simlandi 40 mg / 0.4 mL autoinjector and prefilled syringe, 80 mg / 0.8 mL prefilled syringe**  
**Yuflyma 40 mg / 0.4 mL autoinjector**

**Ankylosing Spondylitis**

For the treatment of patients with active ankylosing spondylitis who are refractory, intolerant or have contraindications to conventional therapy.

Claim Notes:

- Must be prescribed by a rheumatologist or internist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 40 mg every two weeks.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.

**Crohn's Disease**

For the treatment of patients with moderately to severely active Crohn's disease who are refractory, intolerant or have contraindications to conventional therapy.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 160 mg followed by 80 mg two weeks later, then 40 mg every two weeks.
- Initial approval period: 12 weeks.
- Renewal approval period: Long term. Confirmation of response is required.

### **Hidradenitis Suppurativa**

For the treatment of patients with active moderate to severe hidradenitis suppurativa (HS) who are refractory, intolerant or have contraindications to conventional therapy.

#### Claim Notes:

- Must be prescribed by a dermatologist or physician with experience in the treatment of HS.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 160 mg followed by 80 mg two weeks later, then 40 mg every week beginning four weeks after the initial dose.
- Initial approval period: 12 weeks.
- Renewal approval period: Long term. Confirmation of response is required.

### **Plaque Psoriasis**

For the treatment of patients with moderate to severe plaque psoriasis who are refractory, intolerant or have contraindications to conventional therapy.

#### Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 80 mg followed by 40 mg in one week, then 40 mg every two weeks thereafter.
- Initial approval period: 16 weeks.
- Renewal approval period: Long term. Confirmation of response is required.

### **Polyarticular Juvenile Idiopathic Arthritis**

For the treatment of patients with moderately to severely active polyarticular juvenile idiopathic arthritis who are refractory, intolerant or have contraindications to conventional therapy.

#### Claim Notes:

- Must be prescribed by, or in consultation with, a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 40 mg every two weeks.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.

### **Psoriatic Arthritis**

For the treatment of patients with active psoriatic arthritis who are refractory, intolerant or have contraindications to conventional therapy.

#### Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 40 mg every two weeks.
- Initial approval period: 16 weeks.
- Renewal approval period: Long term. Confirmation of response is required.

### **Rheumatoid Arthritis**

For the treatment of patients with moderately to severely active rheumatoid arthritis who are refractory, intolerant or have contraindications to conventional therapy.

#### Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 40 mg every two weeks.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.

### **Ulcerative Colitis**

For the treatment of patients with moderately to severely active ulcerative colitis who are refractory, intolerant or have contraindications to conventional therapy.

#### Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 160 mg followed by 80 mg two weeks later, then 40 mg every two weeks.
- Initial approval period: 8 weeks.
- Renewal approval period: Long term. Confirmation of response is required.

### **Uveitis**

For the treatment of patients with non-infectious uveitis who are refractory, intolerant or have contraindications to conventional therapy.

#### Claim Notes:

- Must be prescribed by, or in consultation with an ophthalmologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 80 mg followed by 40 mg in one week, then 40 mg every two weeks thereafter.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.

### **AFATINIB (GIOTRIF)**

#### **20 mg, 30 mg and 40 mg film-coated tablets**

For the first-line treatment of patients with EGFR mutation-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer.

#### Renewal Criteria:

- Written confirmation that the patient is responding to treatment.

#### Clinical Note:

- Patients must have a good performance status.

#### Claim Notes:

- Approvals will be for a maximum of 40 mg daily.
- Approval period: 1 year.

### **AFLIBERCEPT (EYLEA)**

#### **40 mg/mL solution for intravitreal injection**

#### **Diabetic macular edema**

For the treatment of patients with diabetic macular edema who meet all of the following criteria:

- Clinically significant center-involving macular edema for whom laser photocoagulation is also indicated
- Central retinal thickness greater than or equal to 250 micrometers

#### Claim Notes:

- An initial claim of up to two vials (1 vial per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 vial per eye every 30 days.
- Approval Period: 1 year. Confirmation of continued response is required.

#### **Neovascular (wet) age-related macular degeneration**

For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD).

#### Discontinuation Criteria:

- Reduction in Best Corrected Visual Acuity (BCVA) in the treated eye of 15 letters or more on 2 consecutive visits, attributed to AMD in the absence of other pathology, or
- Reduction in BCVA in the treated eye of 30 letters or more compared to either baseline and/or best recorded level, or
- There is evidence of deterioration of the lesion morphology despite optimum treatment over 3 consecutive visits.

#### Clinical Note:

- BCVA must be provided with initial request and with subsequent renewal requests.

#### Claim Notes:

- An initial claim of up to two vials (1 vial per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 vial per eye every 30 days.
- Approval Period: 1 year.

#### **Retinal vein occlusion (RVO)**

For the treatment of macular edema secondary to central retinal vein occlusion (CRVO) or branch retinal vein occlusion (BRVO).

Claim Notes:

- An initial claim of up to two vials (1 vial per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 vial per eye every 30 days.
- Approval Period: 1 year. Confirmation of continued response is required.

**ALECTINIB (ALECENSARO)**

**150 mg capsule**

For the treatment of patients with anaplastic lymphoma kinase (ALK)-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer when used:

- as first-line therapy, or
- following disease progression on, or intolerance to, crizotinib.

Renewal Criteria

- Written confirmation that the patient is responding to treatment.

Clinical Note:

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

Claim Notes:

- Requests for alectinib will not be considered for patients who experience disease progression on any ALK inhibitor other than crizotinib.
- No further ALK inhibitor will be reimbursed following disease progression on alectinib.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**ALEMTUZUMAB (LEMTRADA)**

**12 mg / 1.2 mL single-use vial**

For the treatment of adult patients with highly active relapsing-remitting multiple sclerosis (RRMS) who meet all the following criteria:

- Confirmed diagnosis based on McDonald criteria.
- Experienced one or more disabling relapses or new MRI activity in the past year.
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5).
- Refractory or intolerant to at least two disease modifying therapies.

Clinical Notes:

1. Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.
2. A relapse is defined as the appearance of new or worsening neurological symptoms in the absence of fever or infection, lasting at least 24 hours yet preceded by stability for at least one month and accompanied by new objective neurological findings observed through evaluation by a neurologist.

Claim Notes:

- Must be prescribed by a neurologist.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Maximum approval quantity and period: 8 vials in 2 years (5 vials approved in year 1 and 3 vials approved in year 2).
- For more information regarding re-treatment, please contact the NB Drug Plans.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**ALGLUCOSIDASE ALFA (MYOZYME)**

**50 mg vial**

For the treatment of infantile-onset Pompe disease, as demonstrated by onset of symptoms and confirmed cardiomyopathy within the first 12 months of life.

Monitoring of therapy

The monitoring of markers of disease severity and response to treatment must include at least:

1. Weight, length and head circumference.
2. Need for ventilatory assistance, including supplementary oxygen, CPAP, BiPAP, or endotracheal intubation and ventilation.
3. Left ventricular mass index (LVMI) as determined by echocardiography (not ECG alone).
4. Periodic consultation with cardiology.
5. Periodic consultation with respiratory.

#### Withdrawal of therapy

1. Patients to be considered for reimbursement of drug costs for alglucosidase alfa treatment must be willing to participate in the long-term evaluation of the efficacy of treatment by periodic medical assessment. Failure to comply with recommended medical assessment and investigations may result in withdrawal of financial support of drug therapy.
2. The development of the need for continuing invasive ventilatory support after the initiation of ERT should be considered a treatment failure. Funding for ERT should not be continued for infants who fail to achieve ventilator-free status, or who deteriorate further, within 6 months after the initiation of ventilatory support.
3. Deterioration of cardiac function, as shown by failure of LV hypertrophy (as indicated by LV mass index) to regress by more than Z=1 unit, or persistent clinical or echocardiographic findings of cardiac systolic or diastolic failure without evidence of improvement, in spite of 24 weeks of ERT, should be considered a treatment failure and funding for ERT should be discontinued.

#### **ALIROCUMAB (PRALUENT) 75 mg/mL and 150 mg/mL prefilled pen**

For the treatment of heterozygous familial hypercholesterolemia (HeFH) in adult patients who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following criteria are met:

- Definite or probable diagnosis of HeFH using the Simon Broome or Dutch Lipid Network criteria or genetic testing; and
- Patient is unable to reach LDL-C target (less than 2.0 mmol/L or at least a 50% reduction in LDL-C from untreated baseline) despite confirmed adherence to at least 3 months of continuous treatment with:
  - high-dose statin (e.g. atorvastatin 80 mg, rosuvastatin 40 mg) in combination with ezetimibe; or
  - ezetimibe alone, if high dose statin is not possible due to rhabdomyolysis, contraindication or intolerance.

#### Initial Renewal Criteria:

- A reduction in LDL-C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L.

#### Subsequent Renewal Criteria:

- The patient continues to maintain a reduction in LDL-C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L.

#### Clinical Notes:

1. LDL-C levels must be provided.
2. Intolerance to high dose statin will be considered if patient has developed documented myopathy or abnormal biomarkers (i.e. creatinine kinase greater than 5 times the upper limit of normal) after trial of at least two statins and
  - for each statin, dose reduction was attempted rather than statin discontinuation, and intolerance was reversible upon statin discontinuation, but reoccurred with statin re-challenge where clinically appropriate; and
  - at least one statin was initiated at the lowest daily starting dose; and
  - other known causes of intolerance have been ruled out.
3. For patients who cannot take ezetimibe due to an intolerance or contraindication, details must be provided.

#### Claim Notes:

- Approvals will be for a maximum of 300mg every 4 weeks.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

#### **ALTEPLASE (CATHFLO) 2 mg vial**

For the treatment of central venous catheter occlusion in home hemodialysis patients.

#### **AMBRISENTAN (VOLIBRIS and generic brands) 5 mg and 10 mg tablets**

For the treatment of patients with Group 1 pulmonary arterial hypertension (PAH) with World Health Organization (WHO) functional class III or IV.

#### Clinical Note:

- The diagnosis of PAH should be confirmed by right heart catheterization.

#### Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of PAH.
- Combined use of more than one endothelin receptor antagonist will not be reimbursed.



- The maximum dose of ambrisentan that will be reimbursed is 10 mg daily.
- Approval period: Long term.

**AMIFAMPRIDINE (RUZURGI)  
10 mg tablet**

For the treatment of Lambert-Eaton myasthenic syndrome (LEMS) in patients 6 years of age or older.

Initial Renewal Criteria:

- An improvement of at least 30% on the Triple Timed Up and Go (3TUG) test compared to baseline measurement.

Subsequent Renewal Criteria:

- The patient continues to maintain an improvement of at least 30% on the 3TUG test compared to baseline measurement.

Clinical Note:

- The 3TUG test score must be provided with initial and renewal requests.

Claim Notes:

- Must be prescribed by a neurologist.
- Approvals will be up to a maximum daily dose of 40 mg for patients weighing less than 45 kg and 100 mg for patients weighing 45 kg or more.
- Initial approval period: 3 months.
- Renewal approval period: 1 year.

**AMLODIPINE (pdp-AMLODIPINE)  
1 mg/mL oral solution**

For use in patients who require administration through a feeding tube or in pediatric patients when oral tablets or capsules are not an option.

Claim Note:

- Approval Period: 1 year.

**APALUTAMIDE (ERLEADA)  
60 mg tablet**

**Non-Metastatic Castration-Resistant Prostate Cancer**

In combination with androgen deprivation therapy (ADT) for the treatment of patients with castration-resistant prostate cancer (CRPC) who meet all of the following criteria:

- No detectable distant metastases by either CT, MRI or technetium-99m bone scan
- Prostate-specific antigen (PSA) doubling time of less than or equal to 10 months during continuous ADT (i.e., high risk of developing metastases)

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of radiographic disease progression.

Clinical Notes:

1. Castration-resistance must be demonstrated during continuous ADT and is defined as a minimum of three rises in PSA, measured at least one week apart, with the last PSA greater than 2 mcg/L.
2. Castrate levels of testosterone must be maintained throughout treatment with apalutamide.
3. Patients must have a good performance status and no risk factors for seizures.
4. Treatment should be discontinued upon radiographic disease progression or unacceptable toxicity.

Claim Notes:

- Requests for apalutamide will not be considered for patients who experience disease progression on enzalutamide or darolutamide.
- Approval period: 1 year.

**Metastatic Castration-Sensitive Prostate Cancer**

In combination with androgen deprivation therapy (ADT) for the treatment of patients with metastatic castration-sensitive prostate cancer (mCSPC) who have had no prior ADT, or are within 6 months of beginning ADT, in the metastatic setting.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status and no risk factors for seizures.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests for apalutamide will not be considered for patients who experience disease progression on enzalutamide.
- Approval period: 1 year.

**APOMORPHINE (KYNMOBI)**  
**10 mg, 15 mg, 20 mg, 25 mg, and 30 mg orally disintegrating film**

For the acute, intermittent treatment of “off” episodes in patients with Parkinson’s Disease (PD) who are receiving optimized PD treatment (i.e. levodopa and derivatives and dopaminergic agonists or MAO-B inhibitors or amantadine derivatives).

Clinical Note:

- Treatment with Kynmobi should be discontinued unless an improvement of at least 3.25 points is achieved in the Movement Disorders Society Unified Parkinson’s Disease Rating Scale Part III (MDS-UPDRS III) score measured within 30 to 60 minutes after a titrated dose of Kynmobi is administered. This assessment should occur not more than one year after Kynmobi has been titrated to a stable and tolerated dose.

Claim Notes:

- The patient must be under the care of a physician experienced in the diagnosis and treatment of PD.
- Approvals will be for a maximum of 90 mg per day not exceeding five films per day.
- Approval period: 1 year.

**APOMORPHINE (MOVAPO)**  
**30 mg / 3 mL prefilled pen**

For the acute, intermittent treatment of hypomobility “off” episodes in patients with advanced Parkinson’s Disease (PD) who are receiving optimized PD treatment (i.e. levodopa and derivatives and dopaminergic agonists).

Clinical Note:

- “Off” episodes are defined as “end of dose wearing off” and unpredictable “on/off” episodes.

Claim Notes:

- The patient must be under the care of a physician experienced in the diagnosis and treatment of PD.
- Approval period: 1 year.

**APREPITANT (EMEND)**  
**80 mg and 125 mg capsules**  
**Tri-Pack 2x80 mg capsules + 125 mg capsule**

In combination with a 5-HT<sub>3</sub> antagonist and dexamethasone for the prevention of acute and delayed nausea and vomiting in patients receiving:

- highly emetogenic chemotherapy, or
- moderately emetogenic chemotherapy who have had inadequate symptom control using a 5-HT<sub>3</sub> antagonist and dexamethasone in a previous cycle.

Claim Note:

- Prescriptions written by hematologists, oncologists, oncology clinical associates, or general practitioners in oncology who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.

**ARIPIRAZOLE (ABILIFY MAINTENA)**  
**300 mg and 400 mg vials**

For the treatment of patients who are:

- not adherent to an oral antipsychotic, or
- currently receiving a long-acting injectable antipsychotic and require an alternative long-acting injectable antipsychotic.

Claim Notes:

- Requests will not be considered for the treatment of psychotic symptoms related to dementia.
- Approval period: Long term.

**ASCIMINIB (SCEMBLIX)  
20 mg and 40 mg tablets**

For the treatment of adult patients with Philadelphia chromosome-positive chronic myeloid leukemia (Ph+ CML) in chronic phase who have resistance or intolerance to at least two tyrosine kinase inhibitors and no evidence of T315i or V299L mutations.

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Clinical Notes:**

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Claim Notes:**

- Requests will not be considered for patients with CML in accelerated or blast phase.
- Approval period : 1 year.

**ASENAPINE (SAPHRIS)  
5 mg and 10 mg sublingual tablets**

For the acute treatment of bipolar I disorder as either:

- Monotherapy, after inadequate response to a trial of lithium or divalproex sodium, and there is a history of inadequate response or intolerance to at least one less expensive antipsychotic agent; or
- Co-therapy with lithium or divalproex sodium, and there is a history of inadequate response or intolerance to at least one less expensive antipsychotic agent.

**Claim Note:**

- Approval period: Long term.

**ASFOTASE ALFA (STRENSIQ)  
18 mg / 0.45 mL, 28 mg / 0.7 mL, 40 mg / 1 mL and 80 mg / 0.8 mL single-use vials**

For the treatment of patients with perinatal, infantile, or juvenile-onset hypophosphatasia (HPP).

**Clinical Note:**

- Eligibility for the treatment of HPP is determined by the Canadian HPP Clinical Expert Committee. Please contact the NB Drug Plans at 1-800-332-3691 for the request form.

**Claim Note:**

- Must be prescribed by a metabolic specialist with expertise in the diagnosis and management of HPP.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**AXITINIB (INLYTA)  
1 mg and 5 mg tablets**

For the treatment of patients with advanced or metastatic renal cell carcinoma when used as:

- first-line therapy in combination with pembrolizumab; or
- second-line therapy following disease progression on a vascular endothelial growth factor receptor tyrosine kinase inhibitor (i.e., sunitinib or pazopanib); or
- third-line therapy following disease progression on first-line nivolumab and ipilimumab combination therapy and a second-line vascular endothelial growth factor receptor tyrosine kinase inhibitor (i.e., sunitinib or pazopanib).

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Clinical Notes:**

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Claim Notes:**

- Requests for axitinib will not be considered for patients who experience disease progression on everolimus, cabozantinib or single-agent nivolumab.
- Approval period: 1 year.

**AZACITIDINE (ONUREG)  
200 mg and 300 mg tablets**

As maintenance therapy for the treatment of adult patients with newly diagnosed acute myeloid leukemia (de novo or secondary to prior MDS or CMML) who meet all of the following criteria:

- Intermediate or poor risk cytogenetics
- Complete remission or complete remission with incomplete blood count recovery following induction therapy, with or without consolidation treatment, within the previous 4 months
- Not eligible for hematopoietic stem cell transplantation

Renewal Criteria:

- Written confirmation that the patient continues to be in complete remission or complete remission with incomplete blood count recovery.

Clinical Note:

- Treatment should be discontinued upon disease relapse (i.e., appearance of greater than 5% blasts in the bone marrow or peripheral blood), unacceptable toxicity or the patient becomes eligible for allogeneic bone marrow or stem cell transplantation.

Claim Notes:

- Requests will not be considered for patients who experience disease progression on hypomethylating agents.
- Approvals will be for a maximum of 300 mg daily for 14 days every 28-day cycle.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**AZITHROMYCIN (generic brands)  
600 mg tablet**

For the prevention of disseminated Mycobacterium Avium Complex (MAC) in HIV positive patients who are severely immunocompromised with CD4 levels  $<0.1 \times 10^9/L$ .

**AZTREONAM (CAYSTON)  
75 mg powder for inhalation**

For the treatment of chronic pulmonary *Pseudomonas aeruginosa* infections, when used as a cyclic treatment, in patients with moderate to severe cystic fibrosis and deteriorating clinical condition despite treatment with inhaled tobramycin.

Clinical Note:

- Cyclic treatment measured in 28-day cycles is defined as 28 days of treatment, followed by 28 days without treatment.

Claim Notes:

- Combined use of aztreonam either concurrently or for antibiotic cycling during off-treatment periods, with other inhaled antibiotics (e.g, tobramycin, levofloxacin) will not be reimbursed.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.

**BARICITINIB (OLUMIANT)  
2 mg tablet**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.

5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 2 mg daily.
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of response is required.

**BENRALIZUMAB (FASENRA)  
30 mg/mL autoinjector and prefilled syringe**

For the adjunctive treatment of severe eosinophilic asthma in adult patients who are inadequately controlled with high dose inhaled corticosteroids (ICS) and one or more additional asthma controller(s) (e.g. long-acting beta-agonist), and meets one of the following criteria:

- blood eosinophil count of  $\geq 0.3 \times 10^9/L$  within the past 12 months and has experienced two or more clinically significant asthma exacerbations in the past 12 months, or
- blood eosinophil count of  $\geq 0.15 \times 10^9/L$  and is receiving maintenance treatment with oral corticosteroids (OCS).

Initial Discontinuation Criteria:

- Baseline asthma control questionnaire score has not improved at 12 months since the initiation of treatment, or
- No decrease in the daily maintenance OCS dose in the first 12 months of treatment, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months

Subsequent Discontinuation Criteria:

- Asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
- Reduction in the daily maintenance OCS dose achieved after the first 12 months of treatment is not maintained subsequently, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months

Clinical Notes:

1. A baseline and annual assessment of asthma symptom control using a validated asthma control questionnaire must be provided.
2. High-dose ICS is defined as greater than or equal to 500 mcg of fluticasone propionate or equivalent daily dose.
3. A significant clinical exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least 3 days or the patient visited an emergency department or was hospitalized.

Claim Notes:

- Must be prescribed by a respirologist, clinical immunologist, allergist or internist experienced in the treatment of severe eosinophilic asthma.
- Combined use of benralizumab with other biologics used to treat asthma will not be reimbursed.
- Approvals will be for a maximum of 30 mg every four weeks for 12 weeks, then every eight weeks thereafter.
- Approval period: 1 year.

**BICTEGRAVIR, EMTRICITABINE AND TENOFOVIR ALAFENAMIDE (BIKTARVY)  
50 mg / 200 mg / 25 mg tablet**

For the treatment of adult patients with HIV-1 infection with no known substitution associated with resistance to the individual components of Biktarvy.

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**BIMEKIZUMAB (BIMZELX)  
160 mg/mL autoinjector and prefilled syringe**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals, or nails
- Refractory, intolerant or unable to access phototherapy

- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for 320 mg given every 4 weeks for 16 weeks then 320 mg every 8 weeks thereafter.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**BINIMETINIB (MEKTOVI)**

**15 mg film-coated tablet**

For the treatment of patients with BRAF V600 mutation-positive locally advanced unresectable or metastatic melanoma when used in combination with encorafenib.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Binimetinib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy.
- Approval period: 6 months.

**BOSENTAN (TRACLEER and generic brands)**

**62.5 mg and 125 mg tablets**

For the treatment of patients with Group 1 pulmonary arterial hypertension (PAH) with World Health Organization (WHO) functional class II, III or IV.

Clinical Note:

- The diagnosis of PAH should be confirmed by right heart catheterization.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of PAH.
- Combined use of more than one endothelin receptor antagonist will not be reimbursed.
- The maximum dose of bosentan that will be reimbursed is 125 mg twice daily.
- Approval period: Long term.

**BOSUTINIB (BOSULIF)**

**100 mg and 500 mg tablets**

For the treatment of adult patients with chronic, accelerated, or blast phase Philadelphia chromosome-positive chronic myelogenous leukemia (Ph+ CML) who have resistance or intolerance to prior tyrosine kinase inhibitor therapy.

Clinical Note:

- Patients must have a good performance status.

Claim Notes:

- Approval period: 1 year.

**BRIGATINIB (ALUNBRIG)**  
**30 mg, 90 mg, 180 mg tablets**

For the treatment of adult patients with anaplastic lymphoma kinase (ALK)-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer who have not been previously treated with an ALK inhibitor.

**Renewal Criteria**

- Written confirmation that the patient is responding to treatment.

**Clinical Note:**

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

**Claim Notes:**

- No further ALK inhibitor will be reimbursed following disease progression on brigatinib.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**BRIVARACETAM (BRIVLERA)**  
**10 mg, 25 mg, 50 mg, 75 mg and 100 mg tablets**

For the adjunctive treatment of refractory partial-onset seizures (POS) in patients who are currently receiving two or more antiepileptic drugs, and who have had an inadequate response or intolerance to at least three other antiepileptic drugs.

**Claim Notes:**

- The patient must be under the care of a physician experienced in the treatment of epilepsy.

**BRODALUMAB (SILIQ)**  
**210 mg / 1.5 mL prefilled syringe**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals, or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

**Clinical Notes:**

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

**Claim Notes:**

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for 210mg at week 0, 1, and 2, then 210mg every two weeks thereafter.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**BROLUCIZUMAB (BEOVU)**  
**6 mg / 0.05 mL prefilled syringe**

**Diabetic Macular Edema**

For the treatment of patients with diabetic macular edema who meet all of the following criteria:

- Clinically significant center-involving macular edema for whom laser photocoagulation is also indicated
- Central retinal thickness greater than or equal to 250 micrometers

**Claim Notes:**

- An initial claim of up to two prefilled syringes (1 per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 prefilled syringe per eye every 6 weeks for 30 weeks, followed by 1 prefilled syringe per eye every 8 weeks thereafter.
- Approval Period: 1 year. Confirmation of continued response is required.

### **Neovascular (wet) age-related macular degeneration**

For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD).

#### Discontinuation Criteria:

- Reduction in Best Corrected Visual Acuity (BCVA) in the treated eye of 15 letters or more on 2 consecutive visits, attributed to AMD in the absence of other pathology, or
- Reduction in BCVA in the treated eye of 30 letters or more compared to either baseline and/or best recorded level, or
- There is evidence of deterioration of the lesion morphology despite optimum treatment over 3 consecutive visits.

#### Clinical Note:

- BCVA must be provided with initial request and with subsequent renewal requests.

#### Claim Notes:

- An initial claim of up to two prefilled syringes (1 per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 prefilled syringe per eye every 4 weeks for 12 weeks, followed by 1 prefilled syringe per eye every 8 weeks thereafter.
- Approval Period: 1 year.

### **BUDESONIDE (PULMICORT NEBUAMP and generic brands) 0.125 mg/mL, 0.25 mg/mL and 0.5 mg/mL suspension for inhalation**

1. For patients who have tried using a budesonide inhaler and
  - cannot follow instructions, or cannot hold the device long enough to actuate it due to cognitive or physical limitations; or
  - have difficulty generating adequate inspiratory effort to achieve therapeutic benefit.

#### Claim Note:

- Approval period: Long term.

2. For patients who require budesonide for sinonasal irrigation when it is prescribed by, or in consultation with, a specialist (e.g., ENT, allergists, immunologists).

#### Claim Notes:

- Initial approval period: 1 year.
- Renewal approval period: Long term.

### **BUPRENORPHINE (PROBUPHINE) 80 mg subdermal implant**

For the treatment of patients with opioid use disorder who have been stabilized on a dose of no more than 8 mg of sublingual buprenorphine for the preceding 90 days.

#### Clinical Notes:

1. Implants are inserted subdermally for up to six months for 4 cycles. Dosing beyond 4 cycles (fifth implantation) is not recommended at this time.
2. Insertion of the subdermal implants should be performed by a healthcare provider who has completed the training program.

#### Claim Notes:

- Approvals will be for 4 implant kits. Requests for additional implants will not be considered.
- Approval period: 2 years.

### **BUPROPION (ZYBAN) 150 mg tablet**

For smoking cessation in adults 18 years of age and older.

#### Clinical Notes:

1. The patient should be participating in a form of smoking cessation counselling.
2. For information on quitting smoking or to obtain the special authorization request form, visit our website [Smoking Cessation Therapies](#).

#### Claim Notes:

- A maximum of 12 weeks of standard therapy (168 tablets) will be reimbursed annually without special authorization.



- Patients who have a high probability of quitting with additional therapy may be approved under special authorization for another 168 tablets.
- Requests for nicotine replacement therapy (patches/gum/lozenge) for use in combination with a non-nicotine prescription smoking cessation drug (varenicline or bupropion) will not be considered.
- Requests for special authorization should be submitted on the Smoking Cessation Therapy Special Authorization Request Form.

**BUROSUMAB (CRYSVITA)**  
**10 mg/mL, 20 mg/mL, 30mg/mL single-use vials**

For the treatment of patients with X-linked hypophosphatemia (XLH) who meet the following criteria:

- Initiated in a pediatric patient who is at least one year of age and in whom epiphyseal closure has not yet occurred
- Fasting hypophosphatemia
- Normal renal function (defined as a serum creatinine below the age-adjusted upper limit of normal)
- Radiographic evidence of rickets with a rickets severity score (RSS) of two or greater
- Confirmed phosphate-regulating endopeptidase homolog, X-linked (PHEX) gene variant in either the patient or in a directly related family member with appropriate X-linked inheritance

**Discontinuation Criteria:**

In pediatric patients under 18 years of age in whom epiphyseal closure has not yet occurred and who met the above criteria, treatment should be discontinued if:

- there is no demonstrated improvement in the 12-month RSS total score from baseline RSS total score; or
- the patient's RSS total score achieved after the first 12 months of therapy has not been maintained subsequently.

In adolescent patients who are 13 to 17 years of age in whom epiphyseal closure has occurred and who met the above criteria and initiated treatment as a pediatric patient, treatment should be discontinued if any of the following occur:

- Hyperparathyroidism; or
- Nephrocalcinosis; or
- Evidence of fracture or pseudo-fracture based on radiographic assessment.

In adult patients who met the above criteria and initiated treatment as a pediatric patient, treatment should be discontinued if any of the following occur:

- Hyperparathyroidism; or
- Nephrocalcinosis; or
- Evidence of fracture or pseudo-fracture based on radiographic assessment.

**Clinical Note:**

- A baseline and annual assessment of the RSS score must be provided for pediatric patients in whom epiphyseal closure has not occurred.

**Claim Notes:**

- Requests will not be considered for treatment-naïve adults.
- Must be prescribed by a physician working in a multidisciplinary team of health care providers who are experienced in the diagnosis and management of XLH.
- Approvals for children (1-17 years of age) will be up to a maximum of 90mg every 2 weeks.
- Approvals for adults (18 years of age and older) will be up to a maximum of 90mg every 4 weeks.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**CABOTEGRAVIR (VOCABRIA)**  
**30 mg tablet**  
**CABOTEGRAVIR and RILPIRIVINE (CABENUVA)**  
**600 mg / 3 mL and 900 mg / 3 mL dosing kit**  
**400 mg / 2 mL and 600 mg / 2 mL dosing kit**

For the treatment of adult patients with HIV-1 infection who are virologically stable and suppressed (HIV-1 RNA less than 50 copies per mL).

**Claim Notes:**

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**CABOZANTINIB (CABOMETYX)  
20 mg, 40 mg, and 60 mg tablets**

**Advanced Hepatocellular Carcinoma**

For the second-line treatment of adult patients with unresectable hepatocellular carcinoma who meet all of the following criteria:

- Disease progression on sorafenib or lenvatinib
- Child-Pugh class status of A
- ECOG performance status of 0 or 1

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and continues to experience clinical benefit.

**Clinical Note:**

- Treatment should continue until the patient no longer experiences clinical benefit or experiences unacceptable toxicity.

**Claim Notes:**

- Requests for cabozantinib will not be considered for patients who experience disease progression on regorafenib or atezolizumab in combination with bevacizumab.
- Approval period: 6 months.

**Metastatic Renal Cell Carcinoma**

For the treatment of patients with advanced or metastatic renal cell carcinoma who have received at least one prior vascular endothelial growth factor receptor (VEGFR) tyrosine kinase inhibitor (TKI) therapy when used as:

- second-line therapy following disease progression on sunitinib, pazopanib or pembrolizumab in combination with axitinib; or
- third-line therapy following disease progression on immunotherapy and VEGFR TKI (i.e., sunitinib or pazopanib), used in any sequence.

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of clinically meaningful disease progression.

**Clinical Note:**

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

**Claim Notes:**

- Requests for cabozantinib will not be considered for patients who experience disease progression on everolimus or axitinib monotherapy.
- Approval period: 1 year.

**CANAGLIFLOZIN (INVOKANA)  
100 mg and 300 mg tablets**

For the treatment of type 2 diabetes mellitus when added to metformin and a sulfonylurea for patients with inadequate glycemic control on metformin and a sulfonylurea.

**Clinical Note:**

- For patients who cannot take metformin and/or a sulfonylurea due to contraindications or intolerances, details must be provided.

**CANAKINUMAB (ILARIS)  
150 mg/mL solution for injection**

For the treatment of active systemic juvenile idiopathic arthritis, in patients 2 years of age or older, who have an inadequate response or intolerance to systemic corticosteroids (with or without methotrexate) and tocilizumab.

**Clinical Note:**

- Intolerance is defined as a serious adverse effect as described in the product monograph. The nature of the intolerance(s) must be clearly documented.

**Claim Notes:**

- Must be prescribed by, or in consultation with, a rheumatologist, who is familiar with the use of biologic DMARDs in children.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for 4 mg/kg for patients weighing more than 9 kg, to a maximum of 300mg, administered every four weeks.

- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**CEFTOLOZANE AND TAZOBACTAM (ZERBAXA)**  
**1 g / 0.5 g vial**

For the treatment of patients with multidrug-resistant *Pseudomonas aeruginosa* when alternative agents are not an option.

Claim Notes:

- Must be prescribed by, or in consultation with, an infectious disease specialist or medical microbiologist.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**CERITINIB (ZYKADIA)**  
**150 mg Capsule**

As monotherapy treatment for patients with anaplastic lymphoma kinase (ALK)-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer who experience disease progression on, or intolerance to, crizotinib.

Renewal Criteria:

- Written confirmation that the patient is responding to treatment.

Clinical Note:

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

Claim Notes:

- Requests for ceritinib will not be considered for patients who experience disease progression on any ALK inhibitor other than crizotinib.
- No further ALK inhibitor will be reimbursed following disease progression on ceritinib.
- Approval: 1 year.

**CERLIPONASE ALFA (BRINEURA)**  
**150 mg / 5 mL solution for intracerebroventricular infusion**

For the treatment of neuronal ceroid lipofuscinosis type 2 (CLN2) disease, if all of the following criteria are met:

- Confirmed diagnosis of CLN2 disease based on tripeptidyl peptidase 1 (TPP1) enzyme activity and CLN2 genotype analysis
- Score of greater than or equal to 1 in each of the motor and language domains of the CLN2 Clinical Rating Scale
- Aggregate motor-language score of greater than or equal to 3 on the CLN2 Clinical Rating Scale

Discontinuation criteria:

- Reduction of greater than or equal to 2 points in the aggregate motor-language score of the CLN2 Clinical Rating Scale that is maintained over any two consecutive 24-week assessments; or
- Aggregate motor-language score of 0 on the CLN2 Clinical Rating Scale at two consecutive 24-week assessments.

Clinical Note:

- Documentation of the most recent motor and language domain scores of the CLN2 Clinical Rating Scale must be provided with all requests.

Claim Notes:

- Must be prescribed by, or in consultation with, a specialist with experience in the treatment of CLN2 disease.
- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**CERTOLIZUMAB PEGOL (CIMZIA)**  
**200 mg/mL autoinjector and prefilled syringe**

**Ankylosing Spondylitis**

- For the treatment of patients with moderate to severe ankylosing spondylitis (e.g., Bath AS Disease Activity Index (BASDAI) score greater than or equal to 4 on 10-point scale) who:
  - have axial symptoms and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each or in whom NSAIDs are contraindicated, or

- have peripheral symptoms and who have failed to respond, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.
- Requests for renewal must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease of at least 2 points on the BASDAI scale, compared with the pre-treatment score, or
  - patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as HAQ or “ability to return to work”).

Clinical Note:

- Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease do not require a trial of NSAIDs.

Claim Notes:

- Must be prescribed by a rheumatologist or internist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 400mg at weeks 0, 2, and 4, then 200mg every two weeks (or 400mg every four weeks).
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**Psoriatic Arthritis**

- For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each.
- For the treatment of patients with predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each; and
  - methotrexate (oral or parenteral) at a dose of greater than or equal to 20mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 8 weeks; and
  - leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 400mg at weeks 0, 2, and 4, then 200mg every two weeks (or 400mg every four weeks)
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 400mg at weeks 0, 2, and 4, then 200mg every two weeks (or 400mg every four weeks)
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**CETIRIZINE (REACTINE and generic brands)  
20 mg film-coated tablet**

For the treatment of patients with moderate to severe chronic urticaria who have had hives, angioedema, or both for at least six weeks.

Claim Note:

- Approval period: Long term.

**CIPROFLOXACIN (CILOXAN and generic brand)  
0.3% ophthalmic solution  
0.3% ophthalmic ointment**

- For the treatment of ophthalmic infections caused by susceptible bacteria.
- For the prevention of ophthalmic infections associated with non-elective eye surgery.

Claim Note:

- Prescriptions written by ophthalmologists and prescribing optometrists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.

**CIPROFLOXACIN (CIPRO and generic brands)  
250 mg, 500 mg and 750 mg tablets**

1. For the treatment of patients with any of the following:
  - Acute exacerbations of chronic obstructive pulmonary disease who are at risk of Pseudomonas infection
  - Bacterial prostatitis
  - Cystic fibrosis-related pulmonary infections
  - Febrile neutropenia
  - Gram-negative infections (e.g., osteomyelitis, joint infections) which are resistant to other oral antibacterials
  - Infections with Pseudomonas aeruginosa (susceptible strains).
  - Severe bacterial gastroenteritis when other antibacterials (e.g., macrolides, sulfamethoxazole/trimethoprim) are ineffective, not tolerated, or contraindicated
  - Severe ("malignant") otitis externa
  - Urinary tract infections or acute uncomplicated pyelonephritis when caused by resistant bacteria or when other antibacterials are ineffective, not tolerated or are contraindicated
2. For chemoprophylaxis of close contacts of a patient with invasive meningococcal disease.
3. For the prevention of endophthalmitis in patients who have had cataract surgery with unplanned vitrectomy.

Claim Notes:

- Prescriptions written by infectious disease specialists, internal medicine specialists, hematologists, medical microbiologists, oncologists, oncology clinical associates, or general practitioners in oncology, respirologists or urologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Ciprofloxacin 250 mg, 500 mg, and 750 mg tablets are regular benefits for beneficiaries of Plan B.

**CIPROFLOXACIN (CIPRO)  
500 mg / 5 mL oral suspension**

For use in patients when oral tablets are not an option and who otherwise meet special authorization criteria for ciprofloxacin tablets.

Claim Note:

- Prescriptions written by infectious disease specialists, internal medicine specialists, hematologists, medical microbiologists, oncologists, oncology clinical associates, or general practitioners in oncology, respirologists or urologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.

**CLADRIBINE (MAVENCLAD)**  
**10 mg tablet**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet all the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Has experienced one or more disabling relapses or new MRI activity in the past year
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)
- Refractory or intolerant to at least one disease modifying therapy (e.g., interferon, glatiramer, dimethyl fumarate, teriflunomide, ocrelizumab)

Clinical Notes:

1. Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.
2. A relapse is defined as the appearance of new or worsening neurological symptoms in the absence of fever or infection, lasting at least 24 hours yet preceded by stability for at least one month and accompanied by new objective neurological findings observed through evaluation by a neurologist.

Claim Notes:

- Must be prescribed by a neurologist.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Approvals will be for 1.75mg/kg to a maximum of 200mg per treatment year.
- Approval period: 2 years.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**COBIMETINIB (COTELLIC)**  
**20 mg tablet**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used in combination with vemurafenib.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Cobimetinib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy.
- Approval period: 6 months.

**CODEINE (CODEINE CONTIN)**  
**50 mg, 100 mg, 150 mg, and 200 mg controlled release tablets**

For the treatment of mild to moderate cancer-related or chronic non-cancer pain.

**CRIZOTINIB (XALKORI)**  
**200 mg and 250 mg capsules**

1. For the treatment of patients with anaplastic lymphoma kinase (ALK)-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer when used as:
  - first-line therapy, or
  - second-line therapy following chemotherapy.
2. As monotherapy for the first-line treatment of patients with ROS1-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer (NSCLC).

Renewal Criteria:

- Written confirmation that the patient is responding to treatment.

Clinical Note:

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

Claim Notes:

- Requests for crizotinib will not be considered for patients who experience disease progression on an ALK inhibitor.
- Approval period: 1 year.

**CYCLOSPORINE (VERKAZIA)  
0.1% ophthalmic emulsion**

For the treatment of pediatric patients between the age of 4 and 18 years of age with severe vernal keratoconjunctivitis (VKC) who meet the following criteria:

- Grade 3 (severe) or 4 (very severe) on the Bonini scale, or
- Grade 4 (marked) or 5 (severe) on the modified Oxford scale.

Discontinuation Criteria:

- Treatment should be discontinued if no improvement in signs and symptoms of VKC is observed, or
- Treatment should be discontinued if signs and symptoms of VKC have resolved.

Clinical Note:

- Documentation of the severity of signs and symptoms of VKC at treatment initiation and renewal must be provided.

Claim Notes:

- The patient must be under the care of a physician experienced in the diagnosis and treatment of VKC.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**CYSTEAMINE (CYSTADROPS)  
0.37% ophthalmic solution**

For the treatment of corneal cystine crystal deposits (CCCDs) in patients 2 years of age and older with cystinosis.

Clinical Note:

- Diagnosis of cystinosis confirmed by cystinosis (lysosomal cystine transporter) gene mutation or elevated white blood cell cystine levels. Documentation must be provided.

Claim Note:

- Must be prescribed by an ophthalmologist experienced in the treatment of CCCDs

**CYSTEAMINE (PROCYSBI)  
25 mg and 75 mg delayed-release capsule**

For the treatment of infantile nephropathic cystinosis with documented cystinosis (lysosomal cystine transporter) gene mutation or elevated white blood cell cystine levels.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician with experience in the diagnosis and management of cystinosis.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**DABIGATRAN ETEXILATE (PRADAXA and generic brand)  
110 mg and 150 mg capsules**

For the prevention of stroke and systemic embolism in patients with atrial fibrillation.

Claim Note:

- Approval period: Long term

**DABRAFENIB (TAFINLAR)  
50 mg and 75 mg capsules**

**Adjuvant Melanoma**

In combination with trametinib for the adjuvant treatment of patients with cutaneous melanoma who meet all of the following criteria:

- Stage IIIA (limited to lymph node metastases of greater than 1 mm) to stage IIID disease (AJCC 8<sup>th</sup> edition)
- BRAF V600-mutation positive
- Completely resected disease including in-transit metastases

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should continue until disease recurrence, unacceptable toxicity, or up to a maximum of 12 months.

Claim Notes:

- Requests will be considered for patients with regional lymph nodes with micrometastases after sentinel lymph node biopsy.
- Requests will not be considered for patients who received adjuvant immunotherapy for greater than three months. Patients may switch to BRAF targeted therapy within the first three months of initiating immunotherapy to complete a total of 12 months of adjuvant treatment.
- Approval period: Up to 12 months.

**Metastatic Melanoma**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used alone or in combination with trametinib.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Dabrafenib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy.
- Approval period: 6 months.

**DALTEPARIN (FRAGMIN)**

**10,000 IU/mL ampoule**

**12,500 IU/mL prefilled syringe**

**25,000 IU/mL multidose vial and prefilled syringe**

1. For the treatment of venous thromboembolism (VTE) and/or pulmonary embolism (PE) for a maximum of 30 days.
2. For the extended treatment of recurrent symptomatic venous thromboembolism (VTE) that has occurred while patients are on therapeutic doses of warfarin.
3. For the prophylaxis of venous thromboembolism (VTE) up to 35 days following elective hip replacement or hip fracture surgery.
4. For the prophylaxis of VTE up to 14 days following elective knee replacement surgery.
5. For the prophylaxis of venous thromboembolism (VTE) post abdominal or pelvic surgery for management of a malignant tumour for up to 28 days.
6. For the treatment and secondary prevention of symptomatic venous thromboembolism (VTE) or pulmonary embolism (PE) for a period of up to 6 months in patients with cancer for whom warfarin therapy is not an option.

Claim Note:

- An annual quantity of 35 days of therapy is available without special authorization.

**DAPAGLIFLOZIN AND METFORMIN (XIGDUO)**

**5 mg / 850 mg, 5 mg / 1000 mg film-coated tablets**

For the treatment of type 2 diabetes mellitus in patients who are already stabilized on therapy with dapagliflozin and metformin, to replace the individual components of dapagliflozin and metformin.

**DAPTOMYCIN (CUBICIN RF)**

**500 mg / 10mL single-use vial**

For the treatment of patients with resistant gram-positive infections, including methicillin-resistant *Staphylococcus aureus* (MRSA) who failed to respond, or have a contraindication or intolerance to vancomycin, or for whom IV vancomycin is not appropriate.

Clinical Note:

- Daptomycin is inhibited by pulmonary surfactant and should not be used to treat respiratory tract infections.



Claim Note:

- Must be prescribed by, or in consultation with, an infectious disease specialist or medical microbiologist.

**DARBEPOETIN ALFA (ARANESP)**

**10 mcg / 0.4 mL, 20 mcg / 0.5 mL, 30 mcg / 0.3 mL, 40 mcg / 0.4 mL, 50 mcg / 0.5 mL, 60 mcg / 0.3 mL, 80 mcg / 0.4 mL, 100 mcg / 0.5 mL, 130 mcg / 0.65 mL, 150 mcg / 0.3 mL, 200 mcg / 0.4 mL, 300 mcg / 0.6 mL and 500 mcg/mL SingleJect® prefilled syringes**

- For the treatment of anemia associated with chronic renal failure.

Claim Note:

- Patients on dialysis (end-stage renal disease) receive darbepoetin through the dialysis units.

- For the treatment of transfusion dependent patients with hematologic malignancies whose transfusion requirements are  $\geq 2$  units of packed red blood cells per month over 3 months.

Clinical Note:

- Approval of further 12 week cycles is dependent on evidence of satisfactory clinical response or reduced treatment requirement to less than 2 units of PRBC monthly.

Claim Note:

- Initial approval for 12 weeks.

**DARIFENACIN (ENABLEX and generic brand)**

**7.5 mg and 15 mg extended-release tablets**

For the treatment of overactive bladder (OAB) with symptoms of urgency, urgency incontinence, and urinary frequency in patients who have an intolerance or insufficient response to an adequate trial of a regular benefit OAB drug (e.g. immediate-release oxybutynin, solifenacin or tolterodine).

Clinical Notes:

1. Requests for the treatment of stress incontinence will not be considered.
2. Not to be used in combination with other pharmacological treatments of OAB.

**DAROLUTAMIDE (NUBEQA)**

**300 mg film-coated tablet**

In combination with androgen deprivation therapy (ADT) for the treatment of patients with non-metastatic castration-resistant prostate cancer who have a prostate-specific antigen (PSA) doubling time of less than or equal to 10 months during continuous ADT (i.e., high risk of developing metastases).

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of radiographic disease progression.

Clinical Notes:

1. Castration-resistance must be demonstrated during continuous ADT and is defined as a minimum of three rises in PSA, measured at least one week apart, with the last PSA greater than 2 mcg/L.
2. Castrate levels of testosterone must be maintained throughout treatment with darolutamide.
3. Patients must have a good performance status.
4. Treatment should be discontinued upon radiographic disease progression or unacceptable toxicity.

Claim Notes:

- Requests for darolutamide will not be considered for patients who experience disease progression on apalutamide or enzalutamide.
- Approval period: 1 year.

**DARUNAVIR AND COBICISTAT (PREZCOBIX)**

**800 mg / 150 mg film-coated tablet**

For treatment of HIV-1 infection in treatment-naïve and treatment-experienced patients without darunavir resistance-associated mutations.

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**DASATINIB (SPRYCEL and generic brands)  
20 mg, 50 mg, 70 mg, 80 mg, 100 mg and 140 mg tablets**

1. For the treatment of adult patients with Philadelphia chromosome-positive chronic myeloid leukemia (Ph+ CML) in chronic, accelerated, or blast phase.
2. For the treatment of patients with Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ ALL).

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Claim Note:**

- Approval period: 1 year.

**DECITABINE / CEDAZURIDINE (INQOVI)  
35 mg / 100 mg tablet**

For the treatment of patients with myelodysplastic syndromes (MDS), including previously treated and untreated, who meet all of the following criteria:

- De novo or secondary MDS including all French-American-British subtypes (i.e., refractory anemia, refractory anemia with ringed sideroblasts, refractory anemia with excess blasts, refractory anemia with excess blasts in transformation, and chronic myelomonocytic leukemia)
- Intermediate-1, intermediate-2, or high-risk MDS, according to the International Prognostic Scoring System
- Have not experienced disease progression on a hypomethylating agent

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Clinical Notes:**

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Claim Note:**

- Approval period: 1 year.

**DEFERASIROX (JADENU and generic brands)  
90 mg, 180 mg and 360 mg film-coated tablet**

For the treatment of chronic iron overload.

**DEFERIPRONE (FERRIPROX)  
1000 mg tablet and 100 mg/mL oral solution**

For the treatment of patients with transfusional iron overload due to thalassemia syndromes when current chelation therapy is inadequate.

**Claim Note:**

- Combined use of more than one iron chelating therapy will not be reimbursed.

**DENOSUMAB (PROLIA)  
60 mg/mL prefilled syringe**

For the treatment of osteoporosis in patients who have:

- a high fracture risk, and
- a contraindication, severe gastrointestinal intolerance, or are refractory to bisphosphonates.

**Clinical Notes:**

1. Refractory is defined as a fragility fracture or evidence of a decline in bone mineral density below pre-treatment baseline levels, despite adherence for one year to osteoporosis therapy.
2. High fracture risk is defined as:
  - Moderate 10-year fracture risk (10% to 20%) as defined by the Canadian Association of Radiologists and Osteoporosis Canada (CAROC) tool or the World Health Organization's Fracture Risk Assessment (FRAX) tool with a prior fragility fracture; or
  - High 10-year fracture risk ( $\geq 20\%$ ) as defined by the CAROC or FRAX tool.

Claim Notes:

- Initial approval period: 1 year.
- Renewal approval period: Long term.

**DENOSUMAB (XGEVA)**  
**120 mg / 1.7 mL single use vial**

For the prevention of skeletal-related events (SREs) in patients with castrate-resistant prostate cancer (CRPC) with one or more documented bone metastases and an ECOG performance status of 0-2\*.

Clinical Note:

- \*Patients who are asymptomatic and those who are symptomatic and in bed less than 50% of the time.

**DESMOPRESSIN (generic brands)**  
**0.1 mg and 0.2 mg tablets**  
**DESMOPRESSIN (DDAVP MELT)**  
**60 mcg and 120 mcg orally disintegrating tablets**

- For the management of diabetes insipidus.
- For the treatment of patients 18 years and older with nocturnal enuresis.

Claim Note:

- Desmopressin oral formulations are a regular benefit for Plans CDEF-18G.

**DESMOPRESSIN (generic brand)**  
**10 mcg metered dose nasal spray**

For the treatment of patients with diabetes insipidus.

Clinical Note:

- The nasal formulations are no longer indicated for nocturnal enuresis due to the risk of hyponatremia.

**DIENOGEST (VISANNE and generic brands)**  
**2 mg tablet**

For the management of pelvic pain associated with endometriosis in patients for whom one or more less costly hormonal options are either ineffective or cannot be used.

Clinical Note:

- Continuous combined oral contraceptives and medroxyprogesterone are examples of less costly hormonal options.

**DIMETHYL FUMARATE (TECFIDERA and generic brands)**  
**120 mg and 240 mg delayed-release capsules**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Experienced one or more disabling relapses or new MRI activity in the past two years
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Prescriptions written by neurologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Combined use with other disease modifying therapies to treat RRMS will not be reimbursed.
- Approval Period: 2 years.

**DIPYRIDAMOLE AND ACETYLSALIC ACID (generic brand)**  
**200 mg / 25 mg capsule**

For the secondary prevention of ischemic stroke/TIA in patients who have experienced a recurrent thrombotic event (stroke, symptoms of TIA) while taking ASA.

**DORNASE ALFA (PULMOZYME)  
1 mg/mL solution**

For the treatment of patients with cystic fibrosis with clinical evidence of lung disease (e.g., frequent pulmonary exacerbations, FEV<sub>1</sub> less than 90% predicted, difficulty clearing secretions).

Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACBDEFGV
- Approval period: Long term.

**DOLUTEGRAVIR AND RILPIVIRINE (JULUCA)  
50 mg / 25 mg tablet**

As a complete regimen to replace the current antiretroviral regimen for the treatment of HIV-1 infection in adult patients who are virologically stable and suppressed (i.e. HIV-1 RNA less than 50 copies per mL).

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**DORAVIRINE (PIFELTRO)  
100 mg tablet**

For use in combination with other antiretrovirals in adult patients with HIV-1 infection, who have no known mutations associated with resistance to doravirine.

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**DULOXETINE (CYMBALTA and generic brands)  
30 mg and 60 mg delayed release capsules**

**Chronic Pain**

For the treatment of patients with chronic pain.

Claim Note:

- The maximum dose reimbursed is 60 mg daily.

**Major Depressive Disorder**

For the treatment of major depressive disorder in patients 18 years and older, who have failed treatment with at least one less costly antidepressant.

Claim Note:

- The maximum dose reimbursed is 60mg daily.

**DUPILUMAB (DUPIXENT)  
200 mg / 1.14 mL prefilled syringe and prefilled pen  
300 mg / 2mL prefilled syringe and prefilled pen**

**Asthma**

1. For the adjunctive treatment of severe asthma with a type 2 or eosinophilic phenotype in patients aged 6 to 11 years of age who are inadequately controlled with medium-to high-dose inhaled corticosteroids (ICS) and one or more additional asthma controller(s) (e.g., long-acting beta-agonist) or high-dose ICS alone and meet the following criteria:

- blood eosinophil count  $\geq 0.15 \times 10^9/L$  within the past 12 months; and
- uncontrolled asthma with at least one clinically significant asthma exacerbation in the past 12 months.

Initial Discontinuation Criteria:

- Baseline asthma control questionnaire score has not improved at 12 months since initiation of treatment, or
- The number of clinically significant asthma exacerbations has increased within the previous 12 months.

**Subsequent Discontinuation Criteria:**

- Asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
- The number of clinically significant asthma exacerbations has increased within the previous 12 months.

**Clinical Notes:**

1. A baseline and annual assessment of asthma symptom control using a validated asthma control questionnaire must be provided.
2. Medium dose ICS is defined as between 200 mcg and 400 mcg of fluticasone propionate or equivalent daily dose and high-dose ICS is defined as greater than 400 mcg of fluticasone propionate or equivalent daily dose.
3. A significant clinical exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least 3 days or the patient visited an emergency department or was hospitalized.

**Claim Notes:**

- Must be prescribed by a pediatric respirologist or allergist experienced in the treatment of severe asthma.
  - Combined use of dupilumab with other biologics used to treat asthma will not be reimbursed.
  - Approvals will be for a maximum of 200 mg every two weeks or 300 mg every four weeks.
  - Approval period: 1 year.
2. For the adjunctive treatment of severe asthma with a type 2 or eosinophilic phenotype or oral corticosteroid (OCS) dependent severe asthma in patients 12 years of age and older who are inadequately controlled with high-dose inhaled corticosteroids (ICS) and one or more additional asthma controller(s) (e.g., long-acting beta-agonist) and meets one of the following criteria:
    - blood eosinophil count  $\geq 0.15 \times 10^9/L$  within the past 12 months, or
    - have OCS dependent asthma.

**Initial Discontinuation Criteria:**

- Baseline asthma control questionnaire score has not improved at 12 months since initiation of treatment, or
- No decrease in the daily maintenance OCS dose in the first 12 months of treatment, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months.

**Subsequent Discontinuation Criteria:**

- Asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
- Reduction in the daily maintenance OCS dose achieved after the first 12 months of treatment is not maintained subsequently, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months.

**Clinical Notes:**

1. A baseline and annual assessment of asthma symptom control using a validated asthma control questionnaire must be provided.
2. A baseline and annual number of clinically significant asthma exacerbations must be provided.
3. High-dose ICS is defined as greater than or equal to 500 mcg of fluticasone propionate or equivalent daily dose.
4. A significant clinical exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least 3 days or the patient visited an emergency department or was hospitalized.

**Claim Notes:**

- Must be prescribed by a respirologist, clinical immunologist, allergist or internist experienced in the treatment of severe asthma.
- Combined use of dupilumab with other biologics used to treat asthma will not be reimbursed.
- Approvals will be for a maximum of 600 mg at week 0, then 300 mg every two weeks thereafter.
- Approval period: 1 year.

**Atopic Dermatitis**

For the treatment of moderate to severe atopic dermatitis (AD) in patients 12 years of age and older who meet all of the following criteria:

- Refractory or have contraindications to an adequate trial of topical prescription therapies combined with phototherapy (where available)
- Refractory, intolerant or have contraindications to an adequate trial of methotrexate, cyclosporine, mycophenolic acid, or azathioprine
- Baseline Physician Global Assessment score of 3 or greater and Eczema Area and Severity Score of 7.1 or greater.

Renewal criteria

- Requests for renewal must provide proof of beneficial clinical effect defined as a 75% or greater improvement from baseline in the Eczema Area and Severity Index (EASI-75) score six months after treatment initiation.
- Proof of maintenance of EASI-75 response from baseline must be provided for subsequent authorizations.

Clinical Note:

- Not to be used in combination with phototherapy or immunomodulatory drugs (e.g., biologics or janus kinase inhibitors) for moderate to severe AD.

Claim Notes:

- Must be prescribed by a dermatologist, pediatrician or clinical immunologist with experience in the treatment of moderate to severe AD.
- Approvals will be for a maximum of 600 mg at week 0, then 300 mg every two weeks thereafter.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**ECULIZUMAB (SOLIRIS)**  
**30 mg / 30 mL single-use vial**

For the treatment of paroxysmal nocturnal hemoglobinuria (PNH).

Clinical Notes:

1. A Request for Coverage including the completed consent and specific special authorization forms must be submitted and the patient must:
  - a) Satisfy the Clinical Criteria for eculizumab (initial or continued coverage, as appropriate);
  - b) Not meet any of the criteria specified in Contraindications to Coverage or Discontinuance of Coverage.
2. Please contact the NB Drug Plans at 1-800-332-3691 for a packet containing the Clinical Criteria and required forms.

Claim Note:

- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**EDARAVONE (RADICAVA)**  
**0.3 mg/mL solution for injection**  
**105 mg / 5 mL oral solution**

For the treatment of patients with probable or definite amyotrophic lateral sclerosis (ALS) who meet all the following criteria:

- ALS Functional Rating Scale – Revised (ALSFRS-R) score of at least two points on each item
- Forced vital capacity (FVC) greater than or equal to 80% of predicted
- ALS symptoms for two years or less
- Permanent non-invasive or invasive ventilation is not required

Discontinuation Criteria:

- The patient is non-ambulatory (ALSFRS-R score less than or equal to 1 for item 8) and unable to cut food and feed themselves without assistance, irrespective of whether a gastrostomy tube is in place (ALSFRS-R score less than 1 for item 5a or 5b); or
- The patient requires permanent non-invasive or invasive ventilation.

Clinical Note:

- ALSFRS-R scores and FVC must be provided.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician with experience in the diagnosis and management of ALS.
- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**ELOSULFASE ALFA (VIMIZIM)**  
**5 mg / 5 mL single-use vial**

For the treatment of patients with mucopolysaccharidosis type IVA (MPS IVA).

Clinical Note:

- Please contact the NB Drug Plans at 1-800-332-3691 for the complete criteria.

Claim Note:

- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**EMPAGLIFLOZIN (JARDIANCE)  
10 mg and 25 mg tablets**

1. For the treatment of type 2 diabetes mellitus when added to metformin and a sulfonylurea for patients with inadequate glycemic control on metformin and a sulfonylurea.
2. As an adjunct to diet, exercise, and standard care therapy to reduce the incidence of cardiovascular death in patients with type 2 diabetes mellitus who have:
  - inadequate glycemic control despite an adequate trial of metformin, or a contraindication or intolerance to metformin; and
  - established cardiovascular disease.

Clinical Notes:

1. For patients who cannot take metformin and/or a sulfonylurea due to contraindications or intolerances, details must be provided.
2. Established cardiovascular disease is defined as one of the following (details must be provided):
  - History of myocardial infarction (MI).
  - Multi-vessel coronary artery disease in two or more major coronary arteries (irrespective of revascularization status).
  - Single-vessel coronary artery disease with significant stenosis and a positive non-invasive stress test.
  - Unstable angina with either coronary multi-vessel or single-vessel disease.
  - History of ischemic or hemorrhagic stroke.
  - Occlusive peripheral artery disease.

**EMPAGLIFLOZIN AND METFORMIN (SYNJARDY)  
5 mg / 500 mg, 5 mg / 850 mg and 5 mg / 1000 mg,  
12.5 mg / 500 mg, 12.5 mg / 850 mg, 12.5 mg / 1000 mg tablet**

For the treatment of type 2 diabetes mellitus in patients who are already stabilized on therapy with empagliflozin and metformin, to replace the individual components of empagliflozin and metformin.

**EMTRICITABINE, RILPIVIRINE AND TENOFOVIR ALAFENAMIDE (ODEFSEY)  
200 mg / 25 mg / 25 mg tablet**

For the treatment of adult patients with HIV-1 infection who meet the following criteria:

- No known mutations associated with resistance to tenofovir, emtricitabine or non-nucleoside reverse transcriptase inhibitor (NNRTI) class.
- Viral load less than or equal to 100,000 copies/mL

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**EMTRICITABINE, TENOFOVIR ALAFENAMIDE, ELVITEGRAVIR AND COBICISTAT (GENVOYA)  
200 mg / 10 mg / 150 mg / 150 mg tablet**

For the treatment of HIV-1 infection in patients 12 years of age and older (weighing at least 35kg) with no known mutations associated with resistance to the individual components of Genvoya.

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval Period: Long term.

**EMTRICITABINE, TENOFOVIR DISOPROXIL, ELVITEGRAVIR AND COBICISTAT (STRIBILD)  
200 mg / 300 mg / 150 mg / 150 mg tablet**

As a complete regimen for antiretroviral treatment naïve HIV-1 infected patients in whom efavirenz is not indicated.

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**ENCORAFENIB (BRAFTOVI)**  
**75 mg capsule**

**Metastatic Colorectal Cancer**

In combination with panitumumab for the treatment of patients with metastatic colorectal cancer who meet all of the following criteria:

- Presence of BRAF V600E mutation
- Disease progression following at least one prior therapy in the metastatic setting
- No previous treatment with an EGFR inhibitor

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Encorafenib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Approval period: 6 months.

**Metastatic Melanoma**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used in combination with binimetinib.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Encorafenib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy.
- Approval period: 6 months.

**ENTRECTINIB (ROZLYTREK)**  
**100 mg and 200 mg capsule**

**Non-Small Cell Lung Cancer**

As monotherapy for the first-line treatment of patients with ROS1-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer.

Renewal Criteria:

- Written confirmation that the patient is responding to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**Solid Tumors with NTRK gene fusion**

As monotherapy for the treatment of adult patients with unresectable locally advanced or metastatic extracranial solid tumors who meet all of the following criteria:

- Tumors have a NTRK gene fusion without a known acquired resistance mutation
- No other satisfactory treatment options
- Not a candidate for surgery and/or radiation due to risk of substantial morbidity

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of radiographic disease progression.



Clinical Notes:

1. Patients must have a good performance status.
2. If central nervous system metastases are present, patients must be asymptomatic.
3. Treatment should be discontinued upon radiographic disease progression or unacceptable toxicity.

Claim Notes:

- Requests will not be considered for patients who experience disease progression on a NTRK inhibitor.
- Approval period: 6 months.

**ENZALUTAMIDE (XTANDI)  
40 mg capsule**

**Metastatic Castration-Resistant Prostate Cancer**

For the treatment of patients with metastatic castration-resistant prostate cancer.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status and no risk factors for seizures.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests for enzalutamide will not be considered for patients who experience disease progression on apalutamide or darolutamide.
- Approval period: 1 year.

**Metastatic Castration-Sensitive Prostate Cancer**

In combination with androgen deprivation therapy (ADT) for the treatment of patients with metastatic castration-sensitive prostate cancer (mCSPC) who have had no prior ADT, or are within 6 months of beginning ADT, in the metastatic setting.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status and no risk factors for seizures.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests for enzalutamide will not be considered for patients who experience disease progression on apalutamide.
- Approval period: 1 year.

**Non-Metastatic Castration-Resistant Prostate Cancer**

In combination with androgen deprivation therapy (ADT) for the treatment of patients with non-metastatic castration-resistant prostate cancer who have a prostate-specific antigen (PSA) doubling time of less than or equal to 10 months during continuous ADT (i.e., high risk of developing metastases).

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of radiographic disease progression.

Clinical Notes:

1. Castration-resistance must be demonstrated during continuous ADT and is defined as a minimum of three rises in PSA, measured at least one week apart, with the last PSA greater than 2 mcg/L.
2. Castrate levels of testosterone must be maintained throughout treatment with enzalutamide.
3. Patients must have a good performance status and no risk factors for seizures.
4. Treatment should be discontinued upon radiographic disease progression or unacceptable toxicity.

Claim Notes:

- Requests for enzalutamide will not be considered for patients who experience disease progression on apalutamide or darolutamide.
- Approval period: 1 year.

**EPLERENONE (INSPIRA and generic brand)  
25 mg and 50 mg tablets**

For the treatment of patients with New York Heart Association (NYHA) class II chronic heart failure with left ventricular systolic dysfunction (with ejection fraction less than or equal to 40%), as an adjunct to standard care therapy.

Clinical Note:

- Patients must be on optimal therapy with an angiotensin-converting–enzyme (ACE) inhibitor or angiotensin-receptor blocker (ARB), and a beta-blocker (unless contraindicated) at the recommended dose or maximal tolerated dose.

**EPOETIN ALFA (EPREX)**

**1,000 IU / 0.5 mL, 2,000 IU / 0.5 mL, 3,000 IU / 0.3 mL, 4,000 IU / 0.4 mL, 5,000 IU / 0.5 mL, 6,000 IU / 0.6 mL, 8,000 IU / 0.8 mL, 10,000 IU/mL, 20,000 IU/mL, 30,000 IU / 0.75 mL and 40,000 IU/mL prefilled syringes**

1. Treatment of anemia associated with chronic renal failure.

Claim Note:

- Patients on dialysis (end-stage renal disease) receive epoetin through the dialysis units
2. Treatment of transfusion dependent anemia related to therapy with zidovudine in HIV-infected patients.
  3. Treatment of transfusion dependent patients with hematologic malignancies whose transfusion requirements are  $\geq 2$  units of packed red blood cells per month over 3 months.

Clinical Note:

- Approval of further 12 week cycles is dependent on evidence of satisfactory clinical response or reduced treatment requirement to less than 2 units of PRBC monthly.

Claim Note:

- Initial approval for 12 weeks.

**EPOPROSTENOL (CARIPUL and FLOLAN)  
0.5 mg and 1.5 mg vials**

For the treatment of patients with Group 1 pulmonary arterial hypertension (PAH) with World Health Organization (WHO) functional class III or IV.

Clinical Note:

- The diagnosis of PAH should be confirmed by right heart catheterization.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of PAH.
- Approval period: Long term.

**ESLICARBAZEPINE (APTIOM)**

**200 mg, 400 mg, 600 mg, 800 mg tablets**

For the adjunctive treatment of refractory partial-onset seizures in patients who are currently receiving two or more antiepileptic drugs and have had an inadequate response or intolerance to at least three other antiepileptic drugs.

Claim Notes:

- The patient must be under the care of a physician experienced in the treatment of epilepsy.

**ETANERCEPT**

**Brenzys 50 mg/mL autoinjector and prefilled syringe**

**Erelzi 25 mg / 0.5 mL prefilled syringe, 50 mg/mL autoinjector and prefilled syringe**

**Ankylosing Spondylitis**

- For the treatment of patients with moderate to severe ankylosing spondylitis (e.g., Bath AS Disease Activity Index (BASDAI) score greater than or equal to 4 on 10-point scale) who:
  - have axial symptoms and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each or in whom NSAIDs are contraindicated, or
  - have peripheral symptoms and who have failed to respond, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.
- Requests for renewal must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease of at least 2 points on the BASDAI scale, compared with the pre-treatment score, or

- patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as HAQ or “ability to return to work”).

**Clinical Note:**

- Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease do not require a trial of NSAIDs.

**Claim Notes:**

- Must be prescribed by a rheumatologist or internist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of etanercept will be approved for the biosimilar versions only.
- Approvals will be for a maximum of 50 mg per week.
- Initial approval period: 6 months.
- Renewal approval period: Long term.

**Plaque Psoriasis**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

**Clinical Notes:**

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

**Claim Notes:**

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of etanercept will be approved for the biosimilar versions only.
- Approvals will be for a maximum of 50 mg twice weekly for 12 weeks, then once weekly thereafter.
- Initial approval period: 16 weeks.
- Renewal approval period: Long term. Confirmation of response is required

**Polyarticular Juvenile Idiopathic Arthritis**

For the treatment of children (age 4-17) with moderately to severely active polyarticular juvenile idiopathic arthritis (pJIA) who have had inadequate response to one or more disease modifying antirheumatic drugs (DMARDs).

**Claim Notes:**

- Must be prescribed by, or in consultation with, a rheumatologist, who is familiar with the use of biologic DMARDs in children.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of etanercept will be approved for the biosimilar version only.
- Approvals will be for a maximum of 0.8mg/kg, up to 50mg per week.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.

**Psoriatic Arthritis**

- For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each.
- For the treatment of patients with predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each; and
  - methotrexate (oral or parenteral) at a dose of greater than or equal to 20mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 8 weeks; and
  - leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months.

**Clinical Notes:**

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.

3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of etanercept will be approved for the biosimilar versions only.
- Approvals will be for a maximum of 50mg once a week.
- Initial approval period: 16 weeks.
- Renewal approval period: Long term. Confirmation of response is required.

**Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of etanercept will be approved for the biosimilar versions only.
- Approvals will be for a maximum of 50mg per week.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.

**ETRAVIRINE (INTELENCE)  
100 mg and 200 mg tablets**

For the treatment of HIV-1 infection in patients who are antiretroviral experienced and have virologic failure due to HIV-1 strains resistant to multiple antiretroviral agents, including other non-nucleoside reverse transcriptase inhibitors.

**EVEROLIMUS (AFINITOR and generic brands)  
2.5 mg, 5 mg and 10 mg tablets**

**Advanced Breast Cancer**

For the treatment of hormone-receptor positive, HER2 negative advanced breast cancer in postmenopausal patients, after recurrence or progression following a non-steroidal aromatase inhibitor, when used in combination with exemestane.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests for everolimus will not be considered for patients who experience disease progression on CDK4/6 inhibitor therapy.
- Approval period: 1 year.

### **Metastatic Renal Cell Carcinoma**

For the treatment of patients with advanced or metastatic renal cell carcinoma following disease progression on tyrosine kinase inhibitor therapy.

#### Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

#### Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

#### Claim Notes:

- Requests for everolimus will not be considered for patients who experience disease progression on axitinib, cabozantinib or nivolumab monotherapy.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

### **Neuroendocrine Tumours**

1. For the treatment of patients with progressive, unresectable, locally advanced or metastatic, well or moderately differentiated pancreatic neuroendocrine tumours (pNET).
2. For the treatment of patients with unresectable, locally advanced or metastatic, well-differentiated, non-functional neuroendocrine tumours (NETs) of gastrointestinal or lung origin (GIL) with documented radiological disease progression within six months.

#### Renewal Criteria

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

#### Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

#### Claim Notes:

- Requests for everolimus will not be considered for patients who experience disease progression on sunitinib for pNET.
- Approval period: 1 year.

### **EVOLOCUMAB (REPATHA)**

**140 mg/mL autoinjector**

**120 mg/mL automated mini-doser with prefilled cartridge**

For the treatment of heterozygous familial hypercholesterolemia (HeFH) in adult patients who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following criteria are met:

- Definite or probable diagnosis of HeFH using the Simon Broome or Dutch Lipid Network criteria or genetic testing; and
- Patient is unable to reach LDL-C target (less than 2.0 mmol/L or at least a 50% reduction in LDL-C from untreated baseline) despite confirmed adherence to at least 3 months of continuous treatment with:
  - high-dose statin (e.g. atorvastatin 80 mg, rosuvastatin 40 mg) in combination with ezetimibe; or
  - ezetimibe alone, if high dose statin is not possible due to rhabdomyolysis, contraindication or intolerance

#### Initial Renewal Criteria:

- A reduction in LDL-C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L.

#### Subsequent Renewal Criteria:

- The patient continues to maintain a reduction in LDL- C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L.

#### Clinical Notes:

1. LDL-C levels must be provided.
2. Intolerance to high dose statin will be considered if patient has developed documented myopathy or abnormal biomarkers (i.e. creatinine kinase greater than 5 times the upper limit of normal) after trial of at least two statins and
  - for each statin, dose reduction was attempted rather than statin discontinuation, and intolerance was reversible upon statin discontinuation, but reoccurred with statin re-challenge where clinically appropriate; and
  - at least one statin was initiated at the lowest daily starting dose; and
  - other known causes of intolerance have been ruled out.

3. For patients who cannot take ezetimibe due to an intolerance or contraindication, details must be provided.

Claim Notes:

- Approvals will be for a maximum of 140mg every 2 weeks or 420mg monthly.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**FARICIMAB (VABYSMO)  
6 mg / 0.05 mL solution for intravitreal injection**

**Diabetic macular edema**

For the treatment of patients with diabetic macular edema who meet all of the following criteria:

- Clinically significant center-involving macular edema for whom laser photocoagulation is also indicated.
- Central retinal thickness greater than or equal to 250 micrometers.

Claim Notes:

- An initial claim of up to two vials (1 vial per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 vial per eye every 4 weeks.
- Approval Period: 1 year. Confirmation of continued response is required.

**Neovascular (wet) age-related macular degeneration**

For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD).

Discontinuation Criteria:

- Reduction in Best Corrected Visual Acuity (BCVA) in the treated eye of 15 letters or more on 2 consecutive visits, attributed to AMD in the absence of other pathology, or
- Reduction in BCVA in the treated eye of 30 letters or more compared to either baseline and/or best recorded level, or
- There is evidence of deterioration of the lesion morphology despite optimum treatment over 3 consecutive visits.

Clinical Note:

- BCVA must be provided with initial request and with subsequent renewal requests.

Claim Notes:

- An initial claim of up to two vials (1 vial per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 vial per eye every 4 weeks for 16 weeks, followed by 1 vial per eye every 8 weeks thereafter.
- Approval Period: 1 year.

**FEBUXOSTAT (generic brands)  
80 mg tablet**

For the treatment of symptomatic gout in patients who are refractory, intolerant or have a contraindication to allopurinol.

**FEDRATINIB (INREBIC)  
100 mg capsule**

For the treatment of splenomegaly and/or disease-related symptoms in adult patients with:

- intermediate-2 or high-risk primary myelofibrosis, post-polycythemia vera myelofibrosis, or post-essential thrombocythemia myelofibrosis; and
- a contraindication or intolerance to ruxolitinib.

Renewal Criteria:

- Confirmation that the patient has responded to treatment as evidenced by a reduction in spleen size or symptom improvement.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued in patients who have progressive increase in spleen size, return of constitutional symptoms or development of serious adverse events.

Claim Notes:

- Requests will not be considered for patients who experience disease progression following treatment with ruxolitinib.
- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**FENTANYL (generic brands)**

**12 mcg/hr, 25 mcg/hr, 50 mcg/hr, 75 mcg/hr and 100 mcg/hr transdermal patch**

For the treatment of cancer-related or chronic non-cancer pain in adult patients who were previously receiving at least 60 mg per day of oral morphine equivalents and who:

- had an inadequate response, intolerance, or contraindication to oral opioids; or
- are unable to take oral therapy.

**FESOTERODINE (TOVIAZ and generic brand)**

**4 mg and 8 mg extended-release tablets**

For the treatment of overactive bladder (OAB) with symptoms of urgency, urgency incontinence, and urinary frequency in patients who have an intolerance or insufficient response to an adequate trial of a regular benefit OAB drug (e.g. immediate-release oxybutynin, solifenacin or tolterodine).

Clinical Notes:

1. Requests for the treatment of stress incontinence will not be considered.
2. Not to be used in combination with other pharmacological treatments of OAB.

**FIDAXOMICIN (DIFICID)**

**200 mg film-coated tablet**

For the treatment of patients with Clostridium difficile infection (CDI), where the patient has:

- a second or subsequent recurrence following treatment with oral vancomycin; or
- treatment failure with oral vancomycin for the current CDI episode; or
- an intolerance or contraindication to oral vancomycin.

Re-treatment criteria:

- Re-treatment with fidaxomicin will only be considered for an early relapse occurring within 8 weeks of the start of the most recent fidaxomicin course.

Clinical Notes:

1. Treatment failure is defined as 14 days of vancomycin therapy without acceptable clinical improvement.
2. Intolerance is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Should be prescribed by, or in consultation with, an infectious disease specialist or gastroenterologist.
- Requests will be approved for 200 mg twice a day for 10 days.

**FILGRASTIM**

**Grastofil 300 mcg / 0.5 mL and 480 mcg / 0.8 mL prefilled syringes**

**Nivestym 300 mcg / 0.5 mL and 480 mcg / 0.8 mL prefilled syringes, 300 mcg/mL and 480 mcg / 1.6 mL vial**

**Chemotherapy Support**

For the prevention of febrile neutropenia in patients receiving myelosuppressive chemotherapy with curative intent who:

- are at high risk of febrile neutropenia due to chemotherapy regimen, co-morbidities or pre-existing severe neutropenia; or
- have had an episode of febrile neutropenia, neutropenic sepsis or profound neutropenia in a previous cycle of chemotherapy; or
- have had a dose reduction, or treatment delay greater than one week due to neutropenia.

Clinical Note:

- Patients with non-curative cancer receiving chemotherapy with palliative intent are not eligible for coverage of filgrastim for prevention of febrile neutropenia.

**Non-Malignant Indications**

- To increase neutrophil count and reduce the incidence and duration of infection in patients with congenital, idiopathic or cyclic neutropenia.
- For the prevention and treatment of neutropenia in patients with HIV infection.

**Stem Cell Transplantation Support**

- For mobilization of peripheral blood progenitor cells for the purpose of stem cell transplantation.
- To enhance engraftment following stem cell transplantation.

Claim Notes:

- All requests for coverage of filgrastim will be approved for the biosimilar versions only.

**FINGOLIMOD (GILENYA and generic brands)  
0.5 mg capsule**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Has experienced one or more disabling relapses or new MRI activity in the past two years
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Combined use with other disease modifying therapies to treat RRMS will not be reimbursed.
- Approval period: 2 years.

**FLUCONAZOLE (DIFLUCAN)  
50 mg / 5 mL powder for oral suspension**

For the treatment of patients who have:

- oropharyngeal candidiasis which failed to respond to nystatin, or
- systemic infections and oral fluconazole tablets are not an option.

**FLUDARABINE (FLUDARA)  
10 mg film-coated tablet**

- For the first-line treatment of patients with chronic lymphocytic leukemia (CLL)/ small lymphocytic lymphoma (SLL) when used in combination with rituximab (with or without cyclophosphamide).
- For the treatment of patients with CLL / SLL who have failed to respond to, or have relapsed during or after previous therapy with an alkylating agent.

**FLUOXETINE (Generic brands)  
20 mg / 5 mL oral solution**

For use in patients for whom oral capsules are not an option.

**FLUTICASONE FUROATE, UMECLIDINIUM AND VILANTEROL (TRELEGY ELLIPTA)  
100 mcg / 62.5 mcg / 25 mcg dry powder for inhalation**

For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry, in patients who experience inadequate control while being treated with a long-acting beta-2 agonist/long-acting anticholinergic (LABA/LAAC).

Clinical Notes:

1. COPD is defined by spirometry as a post-bronchodilator FEV<sub>1</sub>/FVC ratio of less than 0.70. Spirometry reports from any point in time will be accepted. If spirometry cannot be obtained, reasons must be clearly explained, and other evidence of COPD severity provided (i.e. MRC Dyspnea Scale Score grade).
2. Inadequate control while being treated with a LABA/LAAC is defined as persistent symptoms for at least two months or experiencing 2 or more exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids or at least 1 exacerbation of COPD requiring hospitalization.
3. Patients should not be started on a LABA, LAAC and an inhaled corticosteroid (triple inhaled therapy) as initial therapy.



**FORMOTEROL, GLYCOPYRRONIUM BROMIDE AND BUDESONIDE (BREZTRI AEROSPHERE)  
5.8 mcg / 8.2 mcg / 182 mcg suspension for inhalation**

For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry, in patients who experience inadequate control while being treated with a long-acting beta-2 agonist/long-acting anticholinergic (LABA/LAAC).

Clinical Notes:

1. COPD is defined by spirometry as a post-bronchodilator FEV<sub>1</sub>/FVC ratio of less than 0.70. Spirometry reports from any point in time will be accepted. If spirometry cannot be obtained, reasons must be clearly explained, and other evidence of COPD severity provided (i.e. MRC Dyspnea Scale Score grade).
2. Inadequate control while being treated with a LABA/LAAC is defined as persistent symptoms for at least two months or experiencing two or more exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids or at least one exacerbation of COPD requiring hospitalization.
3. Patients should not be started on a LABA, LAAC and an inhaled corticosteroid (triple inhaled therapy) as initial therapy.

Claim Note:

- Approval period: Long term.

**FOSFOMYCIN (MONUROL and generic brand)  
3 g sachet**

For the treatment of uncomplicated urinary tract infections in adult female patients where:

- The infecting organism is resistant to other oral agents,  
OR
- Other less costly agents are not tolerated.

Clinical Note:

- Fosfomycin is not indicated in the treatment of pyelonephritis or perinephric abscess.

**FREMANEZUMAB (AJOVY)  
225 mg / 1.5 mL autoinjector and prefilled syringe**

For the prevention of migraine in patients with a confirmed diagnosis of episodic or chronic migraine who have experienced an inadequate response, intolerance, or contraindication to at least two classes of oral prophylactic migraine medications.

Renewal Criteria:

- A reduction of at least 50% in the average number of migraine days per month at the time of initial renewal compared with baseline.
- At subsequent renewals, the patient continues to maintain the reduction of at least 50% in average number of migraine days per month.

Clinical Notes:

1. The average number of headache and migraine days per month must be provided on initial and renewal requests.
2. According to the International Headache Society criteria, episodic or chronic migraine are defined as:
  - Episodic migraine: migraine headaches on at least 4 days per month and less than 15 headache days per month for more than 3 months.
  - Chronic migraine: headaches for at least 15 days per month for more than 3 months of which at least eight days per month are with migraine.

Claim Notes:

- Combined use with other calcitonin gene-related peptide (CGRP) antagonists will not be reimbursed
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**GALANTAMINE (generic brands)  
8 mg, 16 mg, and 24 mg extended release capsules**

For the treatment of patients with mild to moderate dementia who have had an intolerance to donepezil and who meet the following criteria:

- Mini-Mental State Exam (MMSE) score of 10 to 30
- Functional Assessment Staging Test (FAST) score of 4 to 5

Clinical Notes:

1. Requests must contain an updated MMSE and FAST score completed within 6 months of the request.
2. The nature of the intolerance must be described.

Claim Note:

- Approval period: 1 year.

**GALCANEZUMAB (EMGALITY)  
120 mg/mL autoinjector and prefilled syringe**

For the prevention of migraine in patients with a confirmed diagnosis of episodic or chronic migraine who have experienced an inadequate response, intolerance, or contraindication to at least two classes of oral prophylactic migraine medications.

Renewal Criteria:

- A reduction of at least 50% in the average number of migraine days per month at the time of initial renewal compared with baseline.
- At subsequent renewals, the patient continues to maintain the reduction of at least 50% in average number of migraine days per month.

Clinical Notes:

1. The average number of headache and migraine days per month must be provided on initial and renewal requests.
2. According to the International Headache Society criteria, episodic or chronic migraine are defined as:
  - Episodic migraine: migraine headaches on at least 4 days per month and less than 15 headache days per month for more than 3 months.
  - Chronic migraine: headaches for at least 15 days per month for more than 3 months of which at least eight days per month are with migraine.

Claim Notes:

- Combined use with other calcitonin gene-related peptide (CGRP) antagonists will not be reimbursed.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**GILTERITINIB (XOSPATA)  
40 mg tablet**

As monotherapy for the treatment of adult patients with relapsed or refractory FMS-like tyrosine kinase 3 (FLT3)-mutated acute myeloid leukemia who meet all of the following criteria:

- Confirmed positive for FLT3 mutation at the time of relapse or determination of refractory disease
- Presence of FLT3-ITD, FLT3-TKD/D835 or FLT3-TKD/I836 mutation

Renewal Criteria:

- Written confirmation that the patient is responding to treatment.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should continue as long as clinical benefit is observed or until unacceptable toxicity occurs.

Claim Notes:

- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**GLUCAGON (BAQSIMI)  
3 mg nasal powder**

For patients receiving insulin who are at high risk of hypoglycemia.

Claim Notes:

- A maximum of 2 doses will be reimbursed annually without special authorization for individuals who have had a claim for insulin in the previous 12 months.
- Special authorization requests for additional doses will be considered for up to one dose per month.

**GLYCEROL PHENYLBUTYRATE (RAVICTI)  
1.1 g/mL oral liquid**

For the treatment of patients with urea cycle disorders (UCDs).

Clinical Note:

- Diagnosis must be confirmed by blood, enzymatic, biochemical or genetic testing.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of UCs
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**GLECAPREVIR AND PIBRENTASVIR (MAVIRET)  
100 mg / 40 mg tablet**

For treatment-naïve or treatment-experienced adult patients with chronic hepatitis C virus (HCV) who meet the following criteria:

<b>Approval Period</b>	
<b>Genotypes 1, 2, 3, 4, 5 or 6</b> <ul style="list-style-type: none"><li>• Treatment-naïve</li></ul>	8 weeks
<b>Genotypes 1, 2, 4, 5 or 6</b> <ul style="list-style-type: none"><li>• Treatment-experienced with regimens containing peginterferon/ribavirin (PR) and/or sofosbuvir (SOF)</li></ul>	8 weeks (12 weeks with cirrhosis)
<b>Genotype 1</b> <ul style="list-style-type: none"><li>• NS5A inhibitor treatment-naïve and treatment-experienced with regimens containing:<ul style="list-style-type: none"><li>– Boceprevir/PR; or</li><li>– Simeprevir (SMV)/SOF; or</li><li>– SMV/PR; or</li><li>– Telaprevir/PR</li></ul></li></ul>	12 weeks
<b>Genotype 1</b> <ul style="list-style-type: none"><li>• NS3/4A inhibitor treatment-naïve and treatment-experienced with regimens containing:<ul style="list-style-type: none"><li>– Daclatasvir (DCV)/SOF; or</li><li>– DCV/PR; or</li><li>– Ledipasvir/SOF</li></ul></li></ul>	16 weeks
<b>Genotype 3</b> <ul style="list-style-type: none"><li>• Treatment-experienced with regimens containing PR and/or SOF</li></ul>	16 weeks

The following information is also required:

- Lab-confirmed hepatitis C genotype 1, 2, 3, 4, 5 or 6
- Quantitative HCV RNA value within the last 6 months
- Fibrosis stage

Clinical Note:

- Acceptable methods for the measurement of fibrosis score include Fibrotest, liver biopsy, transient elastography (FibroScan®), serum biomarker panels (such as AST-to-Platelet Ratio Index or Fibrosis-4 score) either alone or in combination.

Claim Notes:

- Must be prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating a patient with hepatitis C infection).
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**GOLIMUMAB (SIMPONI)  
50 mg / 0.5 mL and 100 mg/mL autoinjectors and prefilled syringes**

**Ankylosing Spondylitis**

- For the treatment of patients with moderate to severe ankylosing spondylitis (e.g., Bath AS Disease Activity Index (BASDAI) score greater than or equal to 4 on 10-point scale) who:
  - have axial symptoms and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each or in whom NSAIDs are contraindicated, or
  - have peripheral symptoms and who have failed to respond, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.
- Requests for renewal must include information demonstrating the beneficial effects of the treatment, specifically:

- a decrease of at least 2 points on the BASDAI scale, compared with the pre-treatment score, or
- patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as HAQ or “ability to return to work”).

Clinical Note:

- Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease do not require a trial of NSAIDs.

Claim Notes:

- Must be prescribed by a rheumatologist or internist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 50mg per month.
- Initial approval period: 4 months.
- Renewal approval period: 1 year.

**Psoriatic Arthritis**

- For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each.
- For the treatment of patients with predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each; and
  - methotrexate (oral or parenteral) at a dose of greater than or equal to 20mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 8 weeks; and
  - leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 50mg per month.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 50mg once a month.
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of continued response is required.

### **Ulcerative colitis**

- For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score greater than 4, and a rectal bleeding subscore greater than or equal to 2 and are:
  - refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks, and prednisone greater than or equal to 40mg daily for two weeks or IV equivalent for one week); or
  - corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year).
- Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease in the partial Mayo score greater than or equal to 2 from baseline, and
  - a decrease in the rectal bleeding subscore greater than or equal to 1.

### Clinical Notes:

1. Consideration will be given for patients who have not received a four week trial of aminosalicylates if disease is severe (partial Mayo score greater than 6).
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of the intolerance(s) must be clearly documented.

### Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 200mg at week 0, 100mg at week 2 then 100mg every four weeks thereafter.
- Initial approval period: 3 months.
- Renewal approval period: 1 year.

### **GRASS POLLEN ALLERGEN EXTRACT (ORALAIR) 100 IR and 300 IR sublingual tablets**

For the seasonal treatment of grass pollen allergic rhinitis in patients who have not adequately responded to, or tolerated, conventional pharmacotherapy.

### Clinical Notes:

- Treatment with grass pollen allergen extract must be initiated by physicians with adequate training and experience in the treatment of respiratory allergic diseases.
- Treatment should be initiated four months before the onset of pollen season and should only be continued until the end of the season
- Treatment should not be taken for more than three consecutive years

### **IBRUTINIB (IMBRUVICA) 140 mg capsule**

1. As monotherapy for adult patients with previously untreated chronic lymphocytic leukemia (CLL) / small lymphocytic lymphoma (SLL) for whom fludarabine-based treatment is inappropriate due to high-risk cytogenetic markers (i.e., del17p, TP53 mutation, or unmutated IGHV).
2. As monotherapy for the treatment of patients with CLL/SLL who have received at least one prior therapy.
3. As monotherapy for the treatment of patients with relapsed or refractory mantle cell lymphoma.

### Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

### Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

### Claim Notes:

- Requests will not be considered for patients who experience disease progression on a Bruton's tyrosine kinase (BTK) inhibitor or idelalisib.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

### **ICATIBANT (FIRAZYR) 30 mg / 3 mL prefilled syringe**

For the treatment of acute attacks of type I or type II hereditary angioedema (HAE) in adults with lab confirmed c1-esterase inhibitor deficiency if the following conditions are met:

- Non-laryngeal attacks of at least moderate severity,  
OR

- Acute laryngeal attacks.

Clinical Notes:

1. Using more than three doses in a 24 hour period is not recommended.
2. The safety of more than eight injections per month has not been investigated in clinical trials.

Claim Notes:

- Must be prescribed by, or in consultation with, physicians experienced in the treatment of HAE.
- Coverage is limited to a single dose per attack.
- The maximum quantity that may be dispensed at one time is two doses.

**IDELALISIB (ZYDELIG)  
100 mg and 150 mg film-coated tablets**

For the treatment of patients with relapsed chronic lymphocytic leukemia/small lymphocytic lymphoma, in combination with rituximab.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Note:

- Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will not be considered for patients who experience disease progression on a Bruton's tyrosine kinase (BTK) inhibitor, except as a bridge to transplant.
- Initial approval period: 6 months.
- Renewal approval period: 12 months.

**ICOSAPENT ETHYL (VASCEPA)  
1 g capsule**

To reduce the risk of cardiovascular events (cardiovascular death, non-fatal myocardial infarction, non-fatal stroke, coronary revascularization, or hospitalization for unstable angina) in statin treated patients with elevated triglycerides who meet all of the following criteria:

- 45 years of age and older
- Established cardiovascular disease
- Baseline fasting triglyceride between 1.7 mmol/L and 5.6 mmol/L measured within the three months prior to initiating treatment with Vascepa
- Baseline low-density lipoprotein cholesterol (LDL-C) between 1.0 mmol/L and 2.6 mmol/L
- Receiving a maximally tolerated statin dose for a minimum of 4 weeks, targeted to achieve an LDL-C lower than 2.0 mmol/L

Clinical Note:

- LDL-C and triglyceride levels must be provided.

Claim Notes:

- Approvals will be for a maximum of 4 g daily.
- Approval period: 1 year.

**IMIQUIMOD (ALDARA P and generic brand)  
5% cream**

1. For the treatment of external genital and external perianal/condyloma acuminata warts.

Claim Note:

- Approval Period: 16 weeks

2. For the treatment of actinic keratosis in patients who have failed treatment with 5-Fluorouracil (5-FU) and cryotherapy.

Claim Note:

- Approval Period: 16 weeks.

3. For the treatment of biopsy-confirmed primary superficial basal cell carcinoma:

- with a tumour diameter of ≤ 2 cm

AND

- located on the trunk, neck or extremities (excluding hands and feet)  
AND
- where surgery or irradiation therapy is not medically indicated
  - recurrent lesions in previously irradiated area  
OR
  - multiple lesions, too numerous to irradiate or remove surgically.

Clinical Note:

- Surgical management should be considered first-line for superficial basal cell carcinoma in most patients, especially for isolated lesions.

Claim Note:

- Approval Period: 6 weeks.

**INCOBOTULINUMTOXIN-A (XEOMIN)**

**50 LD<sub>50</sub> units per vial and 100 LD<sub>50</sub> units per vial**

- For the treatment of blepharospasm in patients 18 years of age and older.
- For the treatment of cervical dystonia (spasmodic torticollis) in patients 18 years of age or older.

**INDACATEROL, GLYCOPYRRONIUM BROMIDE, AND MOMETASONE (ENERZAI BREEZHALER)**

**160 mcg / 50 mcg / 150 mcg powder for inhalation**

For the treatment of asthma in patients who are inadequately controlled with a medium or high dose inhaled corticosteroid and a long-acting beta-2 agonist and have experienced one or more asthma exacerbations in the previous 12 months.

**INFLIXIMAB (AVSOLA, INFLECTRA, RENFLEXIS)**

**100 mg vial**

**Ankylosing Spondylitis**

- For the treatment of patients with moderate to severe ankylosing spondylitis (e.g., Bath AS Disease Activity Index (BASDAI) score greater than or equal to 4 on 10-point scale) who:
  - have axial symptoms and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each or in whom NSAIDs are contraindicated, or
  - have peripheral symptoms and who have failed to respond, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.
- Requests for renewal must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease of at least 2 points on the BASDAI scale, compared with the pre-treatment score, or
  - patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as HAQ or “ability to return to work”).

Clinical Note:

- Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease do not require a trial of NSAIDs.

Claim Notes:

- Must be prescribed by a rheumatologist or internist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of infliximab will be approved for the biosimilar versions only.
- Initial approval period: 6 months.
- Renewal approval period: Long term.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**Crohn’s Disease**

For the treatment of patients with moderately to severely active Crohn's disease who have contraindications, or are refractory, to therapy with corticosteroids and other immunosuppressants.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of infliximab will be approved for the biosimilar versions only.
- Initial approval period: 12 weeks.
- Renewal approval period: Long term. Confirmation of response is required.

- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

### **Plaque Psoriasis**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals, or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

#### Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented

#### Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of infliximab will be approved for the biosimilar versions only.
- Initial approval period: 16 weeks.
- Renewal approval period: Long term. Confirmation of response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

### **Psoriatic Arthritis**

- For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each.
- For the treatment of patients with predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each; and
  - methotrexate (oral or parenteral) at a dose of greater than or equal to 20mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 8 weeks; and
  - leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months.

#### Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

#### Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of infliximab will be approved for the biosimilar versions only.
- Initial approval period: 16 weeks.
- Renewal approval period: Long term. Confirmation of response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

### **Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

#### Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.



2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of infliximab will be approved for the biosimilar versions only.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**Ulcerative Colitis**

- For the treatment of patients with moderately to severely active ulcerative colitis who have a partial Mayo score greater than 4, and a rectal bleeding subscore greater than or equal to 2 and are:
  - refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks, and prednisone greater than or equal to 40mg daily for two weeks or IV equivalent for one week); or
  - corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year).
- Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease in the partial Mayo score greater than or equal to 2 from baseline, and
  - a decrease in the rectal bleeding subscore greater than or equal to 1.

Clinical Notes:

1. Consideration will be given for patients who have not received a four week trial of aminosalicylates if disease is severe (partial Mayo score greater than 6).
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- All new requests for coverage of infliximab will be approved for the biosimilar versions only.
- Initial approval period: 12 weeks.
- Renewal approval period: Long term.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**INOTERSEN (TEGSEDI)  
284 mg / 1.5 mL prefilled syringe**

For the treatment of polyneuropathy in adult patients with hereditary transthyretin-mediated amyloidosis (hATTR) who meet all of the following criteria:

- Confirmed genetic diagnosis of hATTR
- Symptomatic early-stage neuropathy
- Does not have New York Heart Association class III or IV heart failure
- Has not previously undergone a liver transplant

Discontinuation Criteria:

- The patient is permanently bedridden and dependent on assistance for basic activities of daily living, or
- The patient is receiving end-of-life care.

Clinical Note:

- Symptomatic early stage neuropathy is defined as Polyneuropathy disability stage I to IIIB or Familial amyloidotic polyneuropathy stage I or II.

Claim Notes:

- The patient must be under the care of a physician with experience in the diagnosis and management of hATTR.
- Combination therapy with other interfering ribonucleic acid drugs or transthyretin stabilizers used to treat hATTR will not be reimbursed.
- Initial approval period: 9 months.

- Renewal approval period: 12 months. Confirmation of continued response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

### **INSULIN DETEMIR (LEVEMIR)**

#### **100 U/mL penfill cartridge and FlexTouch prefilled pen**

1. For the treatment of patients with type 1 or type 2 diabetes who have taken other long acting insulin analogues (insulin glargine and insulin degludec), and have:
  - experienced unexplained nocturnal hypoglycemia at least once a month despite optimal management; or
  - documented severe or continuing systemic or local allergic reaction.
2. For the treatment of pediatric and adolescent patients with type 1 diabetes.
3. For the treatment of pregnant individuals with type 1 or type 2 diabetes requiring insulin.

### **INTERFERON BETA-1A (AVONEX PS)**

#### **30 mcg / 0.5 mL autoinjector and prefilled syringe**

1. For the treatment of adult patients who have experienced a clinically isolated syndrome.
2. For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet the following criteria:
  - Confirmed diagnosis based on McDonald criteria
  - Has experienced one or more disabling relapses or new MRI activity in the past two years
  - Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

#### Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

#### Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACDEFGHV.
- Prescriptions written by neurologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Combined use with other disease modifying therapies to treat MS will not be reimbursed.
- Approval Period: 2 years.

### **INTERFERON BETA-1A (REBIF)**

#### **22 mcg / 0.5 mL and 44 mcg / 0.5 mL prefilled syringes 66 mcg / 1.5 mL and 132 mcg / 1.5 mL prefilled cartridges**

1. For the treatment of adult patients who have experienced a clinically isolated syndrome.
2. For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet the following criteria:
  - Confirmed diagnosis based on McDonald criteria
  - Has experienced one or more disabling relapses or new MRI activity in the past two years
  - Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

#### Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

#### Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACDEFGHV.
- Prescriptions written by neurologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Combined use with other disease modifying therapies to treat MS will not be reimbursed.
- Approval Period: 2 years.

### **INTERFERON BETA-1B (BETASERON)**

#### **0.3 mg single-use vial**

1. For the treatment of adult patients who have experienced a clinically isolated syndrome.
2. For the treatment of adult patients with relapsing-remitting multiple sclerosis who meet the following criteria:
  - Confirmed diagnosis based on McDonald criteria
  - Has experienced one or more disabling relapses or new MRI activity in the past two years
  - Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)
3. For the treatment of adult patients with secondary progressive multiple sclerosis who meet the following criteria.
  - History of RRMS

- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Notes:

- Requests for Betaseron will be considered for individuals enrolled in Plans ACDEFGHV.
- Prescriptions written by neurologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Combined use with other disease modifying therapies to treat MS will not be reimbursed.
- Approval Period: 2 years.

**IPRATROPIUM BROMIDE (generic brands)  
125 mcg/mL and 250 mcg/mL solution for inhalation**

For patients who have tried using an inhaler with spacer device and

- Are unable to follow instructions, hold the spacer device or hold the device long enough to actuate it due to cognitive or physical limitations; or
- Have difficulty generating adequate inspiratory effort to achieve therapeutic benefit.

Claim Notes:

- Initial approval period: 1 year.
- Renewal approval period: Long term.

**FERRIC DERISOMALTOSE (MONOFERRIC)  
100 mg/mL single-use vial**

For the treatment of iron deficiency anemia in patients who

- are intolerant to oral iron replacement products, or
- have not responded to an adequate trial of oral iron.

**ISAVUCONAZOLE (CRESEMBA)  
100 mg capsule  
200 mg vial**

- For the treatment of adult patients with invasive aspergillosis who have a contraindication, intolerance or have failed to respond to oral voriconazole and caspofungin.
- For the treatment of adult patients with invasive mucormycosis.

Claim Notes:

- Must be prescribed by an infectious disease specialist or medical microbiologist.
- Initial requests will be approved for a maximum of 3 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**ITRACONAZOLE (SPORANOX and generic brands)  
10 mg/mL oral solution**

For the treatment of immunocompromised adult patients with oral and/or esophageal candidiasis.

Clinical Note:

- Itraconazole oral solution is not interchangeable with itraconazole capsules due to differences in bioavailability.

**IVABRADINE (LANCORA)  
5 mg and 7.5 mg film-coated tablets**

For the treatment of adult patients with New York Heart Association (NYHA) class II or III stable heart failure when administered in combination with standard care therapy to reduce the incidence of cardiovascular death and hospitalization who meet all of the following criteria:

- Left ventricular ejection fraction (LVEF) of less than or equal to 35%
- Sinus rhythm with a resting heart rate  $\geq 77$  beats per minute (bpm)
- NYHA class II to III symptoms despite at least four weeks of treatment with the following:
  - a stable dose of an angiotensin converting enzyme inhibitor (ACEI) or an angiotensin II receptor blocker (ARB)
  - a stable dose of a beta blocker
  - an aldosterone antagonist

Clinical Notes:

1. Resting heart rate must be documented as  $\geq 77$  bpm on average using either an ECG on at least three separate visits or by continuous monitoring.
2. For patients who have not received four weeks of therapy with an ACEI/ARB, beta blocker and aldosterone antagonist due to an intolerance or contraindication, details must be provided.
3. Initiation and up-titration should be under the supervision of a physician experienced in the treatment of heart failure.

**IVACAFTOR (KALYDECO)  
150mg tablet**

For the treatment of cystic fibrosis in patients who are:

- age 6 years and older and have one of the following cystic fibrosis transmembrane conductance regulator (CFTR) gene mutations: G551D, G1244E, G1349D, G178R, G551S, S1251N, S1255P, S549N or S549R; or
- age 18 years and older with an R117H mutation in the CFTR gene.

Renewal Criteria:

Renewal requests will be considered in patients with documented response to treatment as evidenced by the following:

In cases where the baseline sweat chloride levels were greater than 60 mmol/L:

- the patient's sweat chloride level fell below 60 mmol/L; or
- the patient's sweat chloride level falls by at least 30%

In cases where the baseline sweat chloride levels were below 60 mmol/L:

- the patient's sweat chloride level falls by at least 30%; or
- the patient demonstrates a sustained absolute improvement in FEV<sub>1</sub> of at least 5% when compared to the FEV<sub>1</sub> test conducted prior to starting therapy. FEV<sub>1</sub> will be compared with the baseline pre-treatment level one month and three months after starting treatment

Clinical Notes:

1. The patient's sweat chloride level and FEV<sub>1</sub> must be provided with each request.
2. A sweat chloride test must be performed within a few months of starting ivacaftor therapy to determine if sweat chloride levels are reducing.
  - If the expected reduction occurs, a sweat chloride test must be performed again 6 months after starting therapy to determine if the full reduction has been achieved. Thereafter, sweat chloride levels must be checked annually.
  - If the expected reduction does not occur, a sweat chloride test should be performed again one week later. If the criteria are not met, coverage will be discontinued.

Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- The patient must be under the care of a physician with experience in the diagnosis and management of CF.
- Combined use of more than one CFTR modulator will not be reimbursed.
- Approved dose: 150 mg every 12 hours.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**IVACAFTOR, TEZACAFTOR, ELEXACAFTOR (TRIKAFTA)  
75 mg / 50 mg / 100 mg and 150 mg tablet  
37.5 mg / 25 mg / 50 mg and 75 mg tablet**

For the treatment of patients 6 years of age and older who have at least one F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene.

Initial Renewal Criteria:

The patient must meet one of the following criteria:

- Increase in ppFEV<sub>1</sub> by at least 5% compared with baseline.
- Decrease in the total number of days for which the patient received treatment with oral and/or IV antibiotics for pulmonary exacerbations compared with the six month period prior to initiating treatment.
- Decrease in the total number of pulmonary exacerbations requiring oral and/or IV antibiotics compared with the six month period prior to initiating treatment.
- Decrease in the number of CF-related hospitalizations compared with the six month period prior to initiating treatment.
- No decrease in Body Mass Index (BMI) at six months compared with baseline.
- Increase of 4 points or more on the CF Questionnaire-Revised (CFQ-R) Respiratory Domain Scale compared with baseline.

Subsequent Renewal Criteria:

- Evidence of continued benefit must be provided (e.g., ppFEV<sub>1</sub>, CFQ-R, pulmonary exacerbations).

Clinical Notes:

1. The following baseline measurements must be provided prior to initiation of treatment
  - Spirometry of FEV<sub>1</sub> and ppFEV<sub>1</sub> measured within the 3 month period prior to initiation of treatment
  - Total number of days treated with oral and/or intravenous (IV) antibiotics for pulmonary exacerbations in the 6 months prior to initiation of treatment
  - Total number of pulmonary exacerbations requiring oral and/or IV antibiotics in the 6 months prior to initiation of treatment
  - Number of CF-related hospitalizations in the 6 months prior to initiation of treatment
  - BMI
  - CFQ-R Respiratory Domain score
2. Requests will not be considered for patients who have undergone lung transplantation.

Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- The patient must be under the care of a physician with experience in the diagnosis and management of CF.
- Combined use of more than one CFTR modulator will not be reimbursed.
- Initial approval period: 7 months.
- Renewal approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**IXEKIZUMAB (TALTZ)  
80 mg/mL autoinjector and prefilled syringe**

**Plaque Psoriasis**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals, or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.
4. Treatment should be discontinued if a response has not been demonstrated after 12 weeks.

Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for 160 mg at week 0, followed by 80 mg at weeks 2, 4, 6, 8, 10, and 12 then 80 mg every four weeks.
- Initial approval period: 12 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Psoriatic Arthritis**

- For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each.
- For the treatment of patients with predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each; and
  - methotrexate (oral or parenteral) at a dose of greater than or equal to 20mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 8 weeks; and
  - leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.

3. Intolerance is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for 160 mg at week 0, followed by 80 mg every four weeks.
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**LACTULOSE (various brands)  
667 mg/mL syrup**

For the treatment of hepatic encephalopathy in patients with liver disease.

Clinical Note:

- Please note requests for treatment of constipation will not be considered.

**LAMIVUDINE (generic brands)  
100 mg tablet**

For the treatment of Hepatitis B.

Claim Note:

- Must be prescribed by a hepatologist, gastroenterologist, infectious disease specialist or other physician with experience in the treatment of hepatitis B.

**LAMIVUDINE AND DOLUTEGRAVIR (DOVATO)  
50 mg / 300 mg tablets**

For the treatment of HIV-1 infection in patients 12 years of age or older and weighing at least 40kg, who meet the following criteria:

- HIV-1 treatment-naïve
- Viral load less than or equal to 500,000 copies/mL

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**LAMIVUDINE, TENOFOVIR DISOPROXIL AND DORAVIRINE (DELSTRIGO)  
300 mg / 300 mg / 100 mg tablet**

For the treatment of adult patients with HIV-1 infection with no known mutations associated with resistance to the individual components of Delstrigo.

Claim Notes:

- Prescriptions written for beneficiaries of Plans CU by infectious disease specialists and medical microbiologists who are licensed by the College of Physicians and Surgeons of New Brunswick, do not require special authorization.
- Approval period: Long term.

**LANADELUMAB (TAKHZYRO)  
300 mg vial and prefilled syringe**

For the prevention of attacks of type I or II hereditary angioedema (HAE) in patients 12 years of age and older who have experienced at least three HAE attacks within any four-week period and required the use of an acute injectable treatment.

Discontinuation Criteria:

- No reduction in the number of HAE attacks for which acute injectable treatment was received during the first three months of treatment with lanadelumab compared to the number of attacks observed before initiating treatment with lanadelumab; or
- Increase in the number of HAE attacks for which acute injectable treatment was received compared to the number of attacks before initiating treatment with lanadelumab.

Clinical Note:

- The pre-treatment attack rate must be provided for those patients who are already receiving long-term prophylactic treatment for HAE and intend to transition to lanadelumab.

Claim Notes:

- The patient must be under the care of a physician experienced in the diagnosis and treatment of HAE
- Not to be used in combination with other long-term prophylactic treatment of HAE (e.g., C1 esterase inhibitor).
- Approvals will be for a maximum of 300 mg every two weeks.
- Initial approval period: 3 months.
- Renewal approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**LANSOPRAZOLE (PREVACID and generic brands)  
15 mg and 30 mg delayed-release capsules**

- For patients who have had a therapeutic failure with all proton pump inhibitors listed as regular benefits (e.g. omeprazole, pantoprazole, rabeprazole).
- When compounded as an oral suspension for patients 18 years and younger, who require the use of a proton pump inhibitor and cannot use a tablet or capsule.

Claim Note:

- Approval period: Long term.

**LANSOPRAZOLE (PREVACID FASTAB)  
15 mg and 30 mg delayed-release tablets**

For patients who require drugs to be administered through a feeding tube or cannot use a tablet or capsule.

Claim Note:

- Approval period: Long term.

**LANTHANUM (FOSRENOL)  
250 mg, 500 mg, 750 mg and 1000 mg chewable tablets**

For the treatment of hyperphosphatemia (serum phosphate greater than 1.8 mmol/L) in patients with end-stage renal disease who are intolerant to, or have inadequate control of phosphate levels with, another phosphate binder.

Claim Notes:

- Approval period: Long term.

**LAPATINIB (TYKERB)  
250 mg tablet**

In combination with capecitabine for the treatment of patients with unresectable locally advanced or metastatic HER2-positive breast cancer when used as:

- first-line therapy following disease relapse during or within six months of completing adjuvant treatment with trastuzumab or trastuzumab emtansine; or
- second-line therapy following disease progression on trastuzumab, with or without pertuzumab, in the advanced setting.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and that there is no evidence of disease progression.

Clinical Note:

- Patients must have a good performance status.

Claim Note:

- Approval period: 6 months.

**LAROTRECTINIB (VITRAKVI)  
25 mg and 100 mg capsules  
20 mg / mL oral solution**

As monotherapy for the treatment of adult and pediatric patients with unresectable locally advanced or metastatic solid tumors who meet all of the following criteria:

- Tumors have a NTRK gene fusion without a known acquired resistance mutation
- No other satisfactory treatment options
- Not a candidate for surgery and/or radiation due to risk of substantial morbidity

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of radiographic disease progression.

**Clinical Notes:**

1. Patients must have a good performance status.
2. If brain metastases are present, patients must be asymptomatic.
3. Treatment should be discontinued upon radiographic disease progression or unacceptable toxicity.

**Claim Notes:**

- Requests will not be considered for patients who experience disease progression on a NTRK inhibitor.
- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**LENALIDOMIDE (REVLIMID and generic brands)  
2.5 mg, 5 mg, 10 mg, 15 mg, 20 mg and 25 mg capsules**

**Multiple Myeloma**

1. As first-line treatment for patients with newly diagnosed multiple myeloma who are not eligible for stem cell transplant when used:
  - in combination with dexamethasone, with or without bortezomib; or
  - in combination with daratumumab and dexamethasone.
2. For the treatment of patients with multiple myeloma when used in combination with bortezomib and dexamethasone as induction therapy prior to autologous stem cell transplant.
3. For the treatment of relapsed or refractory multiple myeloma when used:
  - in combination with dexamethasone for patients who have not progressed on lenalidomide; or
  - in combination with carfilzomib and dexamethasone for patients who have not progressed on bortezomib or lenalidomide; or
  - in combination with daratumumab and dexamethasone for patients who have not progressed on lenalidomide.
4. For the maintenance treatment of patients with newly diagnosed multiple myeloma who have stable or improved disease following stem cell transplant and no evidence of disease progression.

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Clinical Notes:**

1. Treatment should be discontinued upon disease progression or unacceptable toxicity.
2. Patients must have a good performance status.

**Claim Note:**

- Approval period: 1 year.

**Myelodysplastic Syndrome**

For the treatment of patients with anemia due to myelodysplastic syndrome who meet all of the following:

- Presence of deletion 5q cytogenetic abnormality
- International Prognostic Scoring System (IPSS) risk category low or intermediate-1
- Transfusion-dependent symptomatic anemia

**Renewal Criteria:**

- Patients who are transfusion-dependent must demonstrate at least fifty percent reduction in transfusion requirements.
- Renewal requests for patients who are not transfusion-dependent may be considered if the patient's serial CBC (pre- and post-lenalidomide) and any other objective evidence of response to therapy is included.

**Clinical Note:**

- Requests for patients who are not transfusion-dependent may be considered. Clinical evidence of symptomatic anemia affecting the patient's quality of life, rationale for why transfusions are not being used, and details pertaining to other therapies prescribed to manage anemia is required.



Claim Note:

- Approval period: 1 year.

**LENVATINIB (LENVIMA)**

**4 mg, 8 mg, and 12 mg per dose compliance packs**

**Advanced Hepatocellular Carcinoma**

For the treatment of unresectable hepatocellular carcinoma, as first-line or second-line therapy after progression on atezolizumab in combination with bevacizumab, for patients who meet all of the following criteria:

- Child-Pugh class status of A
- ECOG performance status of 0 or 1
- Less than 50% liver involvement and no invasion of the bile duct or main portal vein
- No prior liver transplant
- No brain metastases

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Note:

- Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests for lenvatinib will not be considered for patients who have progressed on sorafenib.
- Approval period: 6 months.

**LENVATINIB (LENVIMA)**

**10 mg, 14 mg, 20 mg and 24 mg per dose compliance packs**

**Differentiated Thyroid Cancer**

For the treatment of patients with locally recurrent or metastatic, progressive, differentiated thyroid cancer (DTC) who meet the following criteria:

- Pathologically confirmed papillary or follicular thyroid cancer, and
- Disease that is refractory or resistant to radioactive iodine therapy, and
- Radiological evidence of disease progression within the previous 13 months, and
- Previous treatment with no more than one tyrosine kinase inhibitor (TKI).

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**LETERMOVIR (PREVYMIS)**

**240 mg and 480 mg tablets**

**240 mg / 12 mL and 480 mg / 24 mL vials**

For the prevention of cytomegalovirus (CMV) infection in adult CMV-seropositive recipients [R+] of an allogeneic hematopoietic stem cell transplant (HSCT) who have undetectable CMV viremia at baseline and meet one of the following criteria:

- umbilical cord blood as a stem cell source
- recipient of a haploidentical transplant
- recipient of T-cell depleted transplant
- treated with antithymocyte globulin (ATG) for conditioning
- requiring high-dose steroids or other immunosuppression for acute graft versus host disease (GVHD)
- treated with ATG for steroid-refractory acute GVHD
- documented history of CMV disease prior to transplantation

Clinical Note:

- High-dose steroids is defined as the use of greater than or equal to 1 mg/kg/day of prednisone or equivalent dose of another corticosteroid.

Claim Notes:

- Must be prescribed by a medical oncologist, hematologist, or infectious disease specialist or other physician with experience in the management of HSCT.
- Approvals will be for a maximum dose of 480 mg per day.
- Approval period: 100 days per HSCT.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**LEVETIRACETAM (pdp-LEVETIRACETAM)  
100 mg/mL oral solution**

For use in patients who require administration through a feeding tube or in pediatric patients when oral tablets are not an option.

Claim Note:

- Approval period: 1 year.

**LEVOCARNITINE (CARNITOR and generic brand)  
100 mg/mL oral solution  
330 mg tablet**

1. For the treatment of patients with primary systemic carnitine deficiency.
2. For the treatment of patients with an inborn error of metabolism that results in secondary carnitine deficiency.

**LEVODOPA AND CARBIDOPA (DUODOPA)  
20 mg / 5 mg/mL intestinal gel**

For the treatment of adult patients with advanced levodopa-responsive Parkinson's disease who meet all the following criteria:

- Experiences severe, debilitating motor fluctuations and dyskinesia, with at least 25% of the waking day in the "off" state and/or ongoing levodopa-induced dyskinesias, despite having tried frequent dosing of levodopa (at least five doses per day)
- Received an adequate trial of maximally tolerated doses of levodopa, with demonstrated clinical response
- Failed an adequate trial of each of the following adjunctive medications, if not contraindicated and/or contrary to the clinical judgment of the prescriber: amantadine, a dopamine agonist, entacapone, and a monoamine oxidase (MAO-B) inhibitor

Renewal Criteria:

The patient has a significant reduction in time spent in the "off" state and/or in ongoing levodopa-induced dyskinesias along with improvement in the related disability.

Clinical Note:

- Time in the "off" state, frequency of motor fluctuations, and severity of associated disability should be assessed by a movement disorder subspecialist and be based on an adequate and reliable account (e.g. clinical interview of a patient or care partner, motor symptom diary).

Claim Notes:

- Must be prescribed by a movement disorder subspecialist who has appropriate training in the use of Duodopa and are practising in a movement disorder clinic that provides ongoing management and support for patients receiving treatment with Duodopa.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**LEVODOPA, CARBIDOPA AND ENTACAPONE (STALEVO)  
50 mg / 12.5 mg / 200 mg, 75 mg / 18.75 mg / 200 mg, 100 mg / 25 mg / 200 mg, 125 mg / 31.25 mg / 200 mg,  
and 150 mg / 37.5 mg / 200 mg tablets**

For the treatment of patients with Parkinson's disease

- who are currently receiving immediate-release levodopa/carbidopa and entacapone,  
OR
- who are not well controlled and are experiencing significant "wearing off" symptoms despite optimal therapy with levodopa/decarboxylase.

**LEVOFLOXACIN (generic brands)  
250 mg, 500 mg and 750 mg tablets**

1. For completion of treatment initiated in the hospital setting for patients with nosocomial pneumonia, community acquired pneumonia (CAP) or acute exacerbation of chronic obstructive pulmonary disease (AECOPD).

2. For the treatment of complicated AECOPD in patients who:
  - have failed treatment with at least one first-line therapy (doxycycline, beta-lactam, trimethoprim-sulfamethoxazole, or macrolide), or
  - are intolerant or have contraindication(s) to at least two first-line therapies.
3. For the treatment of CAP in patients with radiographic confirmation of pneumonia who:
  - have failed treatment with at least one first-line therapy (macrolide, doxycycline, beta-lactams), or
  - are intolerant or have contraindication(s) to at least two first-line therapies.
4. For the treatment of pulmonary infections in patients with cystic fibrosis.
5. For the treatment of severe pneumonia in nursing home patients.
6. For the treatment of patients with complicated osteomyelitis or joint infections.
7. For the treatment of patients with pyelonephritis.

Clinical Notes:

1. If the patient has been treated with an antibiotic within the past 3 months consider an antibiotic from a different class.
2. Complicated AECOPD is defined as patients with COPD (FEV<sub>1</sub>/FVC greater than 0.7) experiencing increased sputum purulence, and with increased dyspnea or sputum volume, and one of the following:
  - FEV<sub>1</sub> less than 50% predicted
  - At least 4 exacerbations per year
  - Ischemic heart disease
  - Home oxygen use
  - Chronic oral steroid use

Claim Notes:

- Prescriptions written by infectious disease specialists, internal medicine specialists, hematologists, medical microbiologists, oncologists, oncology clinical associates, general practitioners in oncology, respirologists or urologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Levofloxacin is a regular benefit for Plans BV.

**Tuberculosis**

For the treatment of tuberculosis in patients who have lab-verified drug resistance or a contraindication or intolerance to first-line drugs.

Claim Notes:

- Must be prescribed by, or in consultation with, an infectious disease specialist.
- Requests will only be considered under Plan P.

**LEVOFLOXACIN (QUINSAIR)  
240 mg / 2.4 mL solution for inhalation**

For the treatment of chronic pulmonary *Pseudomonas aeruginosa* infections, when used as a cyclic treatment, in adult patients with cystic fibrosis who have experienced treatment failure with inhaled tobramycin.

Clinical Note:

- Cyclic treatment measured in 28-day cycles is defined as 28 days of treatment, followed by 28 days without treatment.

Claim Notes:

- Combined use of inhaled levofloxacin, either concurrently or for antibiotic cycling during off-treatment periods, with other inhaled antibiotics (e.g. tobramycin, aztreonam) will not be reimbursed.
- Requests will be considered for individuals in Plans ACDEFGV.

**LINEZOLID (generic brands)  
600 mg tablet**

- For treatment of proven vancomycin-resistant *enterococci* (VRE) infections.
- For the treatment of proven methicillin-resistant *Staphylococcus aureus* (MRSA) / methicillin-resistant *Staphylococcus epidermidis* (MRSE) infections in patients who are unresponsive to, or intolerant of, intravenous vancomycin or in whom intravenous vancomycin is not appropriate.

Claim Note:

- The drug must be prescribed by, or in consultation with, an infectious disease specialist or medical microbiologist.

**LISDEXAMFETAMINE (VYVANSE)****10 mg, 20 mg, 30 mg, 40 mg, 50 mg and 60 mg capsules and chewable tablets**

For treatment of patients with Attention Deficit Hyperactivity Disorder who have tried extended release methylphenidate, dexamphetamine or mixed salts amphetamine with unsatisfactory results.

Claim Note:

- The maximum dose reimbursed is 60 mg daily.

**LIXISENATIDE (ADLYXINE)****0.05 mg/mL and 0.1 mg/mL prefilled pen**

For the treatment of type 2 diabetes mellitus when added to:

- basal insulin for patients who have inadequate glycemic control on basal insulin; or
- basal insulin and metformin for patients who have inadequate glycemic control on metformin and basal insulin.

**LONG-ACTING ANTICHOLINERGICS (LAAC)****Acclidinium bromide (Tudorza Genuair 400 mcg powder for inhalation)****Glycopyrronium bromide (Seebri Breezhaler 50 mcg powder for inhalation)****Tiotropium bromide (Spiriva 18 mcg powder for inhalation, Spiriva Respimat 2.5 mcg solution for inhalation)****Umeclidinium bromide (Incruse Ellipta 62.5 mcg powder for inhalation)**

- For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry, in patients who experience:
  - persistent symptoms, as defined by Medical Research Council (MRC) Dyspnea Scale of at least Grade 3 or a COPD Assessment test (CAT) score of at least 10, and have a post-bronchodilator FEV<sub>1</sub> less than 80% predicted; or
  - two or more moderate exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids; or
  - at least one acute severe exacerbation of COPD requiring hospitalization.
- For the treatment of COPD, as defined by spirometry, in combination with a long-acting beta-2 agonist/inhaled corticosteroid (LABA/ICS), for patients who have inadequate control while being treated with a LABA/ICS or a long-acting beta-2 agonist/long-acting anticholinergic (LABA/LAAC).

Clinical Notes:

1. COPD is defined by spirometry as a post-bronchodilator FEV<sub>1</sub>/FVC ratio less than 0.70. Spirometry reports from any point in time will be accepted. If spirometry cannot be obtained, reasons must be clearly explained, and other evidence of COPD severity provided (i.e. MRC Dyspnea Scale grade).
2. Inadequate control while being treated with a LABA/LAAC or LABA/ICS is defined as persistent symptoms for at least two months or experiencing 2 or more exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids, or at least 1 exacerbation of COPD requiring hospitalization.

Claim Note:

- Requests for combination therapy of single agent long-acting bronchodilators, i.e. LABA and LAAC, will not be considered. Products which combine a LABA/LAAC in a single device are available as special authorization benefits with their own criteria.

**LONG-ACTING BETA-2 AGONISTS (LABA)****Formoterol (Oxeze Turbuhaler 6 mcg, 12 mcg powder for inhalation)****Formoterol (Foradil 12 mcg powder for inhalation)****Indacaterol (Onbrez Breezhaler 75 mcg powder for inhalation)****Salmeterol (Serevent Diskus 50 mcg powder for inhalation)****Asthma**

For the treatment of asthma in patients who are using optimal corticosteroid treatment but are still poorly controlled.

**Chronic Obstructive Pulmonary Disease**

For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry, in patients who experience:

- persistent symptoms, as defined by Medical Research Council (MRC) Dyspnea Scale of at least Grade 3 or a COPD Assessment test (CAT) score of at least 10, and have a post-bronchodilator FEV<sub>1</sub> less than 80% predicted; or
- two or more moderate exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids; or
- at least one acute severe exacerbation of COPD requiring hospitalization.

Clinical Note:

- COPD is defined by spirometry as a post-bronchodilator FEV<sub>1</sub>/FVC ratio less than 0.70. Spirometry reports from any point in time will be accepted. If spirometry cannot be obtained, reasons must be clearly explained, and other evidence of COPD severity provided (i.e. MRC Dyspnea Scale grade).

Claim Notes:

- Requests for combination therapy of single agent long-acting bronchodilators, i.e. long-acting beta-2 agonist (LABA) and long-acting anticholinergic (LAAC), will not be considered. Products which combine a LABA/LAAC in a single device are available as special authorization benefits with their own criteria.
- Oxeze Turbuhaler is not indicated for the treatment of COPD, therefore requests will only be considered for the treatment of asthma.
- Onbrez Breezhaler is not indicated for the treatment of asthma, therefore requests will only be considered for the treatment of COPD.

**LONG-ACTING BETA-2 AGONISTS/INHALED CORTICOSTEROID (LABA/ICS) COMBINATIONS**

**Formoterol and Budesonide (Symbicort Turbuhaler 6 mcg / 100 mcg, 6 mcg / 200 mcg powder for inhalation)**

**Formoterol and Mometasone (Zenhale 5 mcg / 100mcg, 5 mcg / 200 mcg suspension for inhalation)**

**Indacaterol and Mometasone (Atectura Breezhaler 150 mcg / 80 mcg, 150 mcg / 160 mcg, 150 mcg / 300 mcg powder for inhalation)**

**Salmeterol and Fluticasone (Advair 25 mcg / 125 mcg, 25 mcg / 250 mcg suspension for inhalation)**

**Salmeterol and Fluticasone (Advair Diskus and generic brands 50 mcg / 100 mcg, 50 mcg / 250 mcg, 50 mcg / 500 mcg powder for inhalation)**

**Vilanterol and Fluticasone (Breo Ellipta 25 mcg / 100 mcg, 25 mcg / 200 mcg powder for inhalation)**

**Asthma**

For the treatment of asthma in patients who are:

- Stabilized on an inhaled corticosteroid and a long-acting beta-2 agonist, or
- Using optimal doses of inhaled corticosteroids but are still poorly controlled.

**Chronic Obstructive Pulmonary Disease**

- For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry, in combination with a long-acting anticholinergic (LAAC), in patients who experience inadequate control while being treated with a long-acting beta-2 agonist/long-acting anticholinergic (LABA/LAAC).
- For the treatment of patients with asthma / chronic obstructive pulmonary disease (ACO) overlap, based on patient history and lung function studies indicating an ACO diagnosis.

Clinical Notes:

1. COPD is defined by spirometry as a post-bronchodilator FEV<sub>1</sub>/FVC ratio less than 0.70. Spirometry reports from any point in time will be accepted. If spirometry cannot be obtained, reasons must be clearly explained, and other evidence of COPD severity provided (i.e. MRC Dyspnea Scale grade).
2. Inadequate control while being treated with a LABA/LAAC is defined as persistent symptoms for at least two months or experiencing 2 or more exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids or at least 1 exacerbation of COPD requiring hospitalization.

Claim Note:

- Ateectura Breezhaler, Breo Ellipta 25mcg/200mcg and Zenhale are not indicated for the treatment of COPD, therefore requests for these products will only be considered for asthma.

**LONG-ACTING BETA-2 AGONIST/ LONG-ACTING ANTICHOLINERGIC (LABA/LAAC) COMBINATIONS**

**Formoterol and Aclidinium bromide (Duaklir Genuair 12 mcg / 400 mcg powder for inhalation)**

**Indacaterol and Glycopyrronium bromide (Ultibro Breezhaler 110 mcg / 50 mcg powder for inhalation)**

**Olodaterol and Tiotropium bromide (Inspiro Respimat 2.5 mcg / 2.5 mcg solution for inhalation)**

**Vilanterol and Umeclidinum bromide (Anoro Ellipta 25 mcg / 62.5 mcg powder for inhalation)**

For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry, in patients who experience inadequate control while being treated with either a long-acting beta-2 agonist (LABA) or long-acting anticholinergic (LAAC).

Clinical Notes:

1. COPD is defined by spirometry as a post-bronchodilator FEV<sub>1</sub>/FVC ratio less than 0.70. Spirometry reports from any point in time will be accepted. If spirometry cannot be obtained, reasons must be clearly explained, and other evidence of COPD severity provided (i.e. Medical Research Council (MRC) Dyspnea Scale grade).
2. Inadequate control is defined as persistent symptoms (e.g. MRC Dyspnea Scale of at least grade 3 or COPD Assessment test (CAT) score of at least 10) after at least one month of a LAAC or LABA.
3. LABA/LAAC combinations are not intended to be used with an inhaled corticosteroid (ICS) unless criteria for triple inhaled therapy (LABA/LAAC/ICS) is met.

**LUSPATERCEPT (REBLOZYL)  
25 mg and 75 mg vials**

**Beta-Thalassemia Anemia**

For the treatment of adult patients with red blood cell (RBC) transfusion-dependent anemia associated with beta-thalassemia who are receiving regular transfusions.

**Initial Renewal Criteria:**

- A reduction of 33% or greater in transfusion burden measured as the number of RBC units required in the initial 24 weeks of luspatercept treatment compared to the 24 weeks prior to luspatercept initiation.

**Subsequent Renewal Criteria:**

- Maintenance of a 33% or greater reduction in transfusion burden measured as the number of RBC units required in the past 24 weeks compared to the 24 weeks prior to luspatercept initiation.

**Clinical Notes:**

1. Regular transfusions are defined as receiving 6 to 20 RBC units and having no transfusion-free period greater than 35 days in the 24 weeks prior to initiating treatment.
2. History of transfusion burden must be provided with the initial and renewal requests.
3. Treatment should be discontinued if there is no response (as defined in renewal criteria) after 3 doses at the maximum dose.

**Claim Notes:**

- Must be prescribed by a hematologist.
- Approvals will be for a maximum of 1.25mg/kg (up to 120mg per dose) every three weeks.
- Approval period: 7 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**Myelodysplastic Syndromes (MDS) Associated Anemia**

For the treatment of adult patients with MDS-associated anemia who meet all of the following criteria:

- Diagnosed with very low- to intermediate-risk MDS with ringed sideroblasts in accordance with the Revised International Prognostic Scoring System (IPSS-R)
- Failed or are not suitable for erythropoietin stimulating agents (ESA)
- Red blood cell (RBC) transfusion-dependent anemia associated with MDS defined as having received at least 2 RBC units over 8 weeks
- Absence of deletion 5q cytogenetic abnormality
- Performance status of 0 to 2

**Initial Renewal Criteria:**

- Patient is RBC transfusion independent over a minimum of 16 consecutive weeks within the first 24 weeks of treatment with luspatercept.

**Subsequent Renewal Criteria:**

- Patient maintains transfusion independence with luspatercept treatment.

**Clinical Notes:**

1. History of transfusion burden must be provided with the initial and renewal requests.
2. Confirmation must be provided that the patient remains very low- to intermediate risk.
3. Details of ESA use (i.e. name of treatment, dose(s), duration of use, response) must be provided.

**Claim Notes:**

- Must be prescribed by a hematologist or oncologist.
- Approvals will be for a maximum of 1.75 mg/kg (up to 168 mg per dose) every three weeks.
- Approval period: 7 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**MACITENTAN (OPSUMIT)  
10 mg film-coated tablets**

For the treatment of patients with Group 1 pulmonary arterial hypertension (PAH) with World Health Organization (WHO) functional class II, III or IV.

**Clinical Note:**

- The diagnosis of PAH should be confirmed by right heart catheterization.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of PAH.
- Combined use of more than one endothelin receptor antagonists will not be reimbursed.
- The maximum dose of macitentan that will be reimbursed is 10 mg daily.
- Approval period: Long term.

**MARAVIROC (CELSENTRI)**  
**150 mg and 300 mg film-coated tablets**

For the treatment of HIV-1 infection in patients who have CCR5 tropic viruses and who have documented resistance to at least one agent from each of the three major classes of antiretrovirals (i.e. nucleoside/tide reverse transcriptase inhibitors, non-nucleoside reverse transcriptase inhibitors and protease inhibitors.)

Clinical Note:

- Requests for HIV-1 treatment-naïve patients will not be considered.

**MECASERMIN (INCRELEX)**  
**10 mg/mL multidose vial**

For the treatment of patients between 2 and 18 years of age with growth failure due to confirmed severe primary insulin-like growth factor-1 deficiency (SPIGFD) in whom epiphyseal closure has not yet occurred and meet the following criteria:

- Documented genetic mutation recognized as a cause of SPIGFD; or
- Clinical and biochemical features of SPIGFD.

Renewal Criteria:

- Height velocity is 1 cm or greater per 6 months or 2 cm or greater per year; and
- Bone age is 16 years or less in boys and 14 years or less in girls.

Clinical Notes:

1. Clinical and biochemical features of SPIGFD are defined as:
  - height standard deviation score less than or equal to  $-3.0$ ; and
  - basal insulin-like growth factor-1 (IGF-1) levels below the 2.5th percentile for age and gender; and
  - random or stimulated growth hormone (GH) level  $> 10$  ng/mL and failure to increase IGF-1 by 50 ug/L in response to exogenous GH during an IGF-1 generation test.
2. Exclusion of secondary forms of IGF-1 deficiency such as malnutrition, hypopituitarism, hypothyroidism, or chronic treatment with pharmacologic doses of anti-inflammatory steroids.

Claim Notes:

- Must be prescribed by a pediatric endocrinologist.
- Mecasermin will not be reimbursed in combination with recombinant growth hormone treatment.
- Approvals will be for a maximum of 0.12 mg/kg/dose twice daily.
- Approval period: 1 year
- Claims that exceed the maximum claim amount of \$9,999 must be divided and submitted as separate transactions as outlined [here](#).

**MEPOLIZUMAB (NUCALA)**  
**100 mg/mL single-use vial, autoinjector and prefilled syringe**

For the adjunctive treatment of severe eosinophilic asthma in adult patients who are inadequately controlled with high-dose inhaled corticosteroids (ICS) and one or more additional asthma controller(s) (e.g. a long-acting beta-agonist), and meets one of the following criteria:

- blood eosinophil count of  $\geq 0.3 \times 10^9/L$  and has experienced two or more clinically significant asthma exacerbations in the past 12 months, or
- blood eosinophil count of  $\geq 0.15 \times 10^9/L$  and is receiving treatment with daily oral corticosteroids (OCS).

Initial Discontinuation Criteria:

- Baseline asthma control questionnaire score has not improved at 12 months since the initiation of treatment, or
- No decrease in the daily maintenance OCS dose in the first 12 months of treatment, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months

Subsequent Discontinuation Criteria:

- Asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
- Reduction in the daily maintenance OCS dose achieved after the first 12 months of treatment is not maintained subsequently, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months

Clinical Notes:

1. A baseline and annual assessment of asthma symptom control using a validated asthma control questionnaire must be provided.
2. High-dose ICS is defined as greater than or equal to 500 mcg of fluticasone propionate or equivalent daily dose.
3. Significant clinical exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least 3 days or the patient visited an emergency department or was hospitalized.

Claim Notes:

- Must be prescribed by a respirologist, clinical immunologist, allergist or internist experienced in the treatment of severe eosinophilic asthma.
- Combined use of mepolizumab with other biologics used to treat asthma will not be reimbursed.
- Approvals will be for a maximum of 100mg every four weeks.
- Approval period: 1 year.

**METFORMIN AND LINAGLIPTIN (JENTADUETO)**  
**500 mg / 2.5 mg, 850 mg / 2.5 mg, and 1000 mg / 2.5 mg tablets**

For the treatment of type 2 diabetes mellitus in patients who are already stabilized on therapy with linagliptin and metformin, to replace the individual components of linagliptin and metformin.

**METFORMIN AND SAXAGLIPTIN (KOMBOGLYZE)**  
**500 mg / 2.5 mg, 850 mg / 2.5 mg, and 1000 mg / 2.5 mg tablets**

For the treatment of type 2 diabetes mellitus in patients who are already stabilized on therapy with saxagliptin and metformin, to replace the individual components of saxagliptin and metformin.

**METHADONE**  
**Compounded Oral Solution**

For the management of severe cancer-related or chronic non-malignant pain.

Claim Note:

- Claims submitted by pharmacies must be billed using PIN 00999801

**METHADONE (METADOL)**  
**1 mg, 5 mg, 10 mg and 25 mg tablets**  
**1 mg/mL oral solution and 10 mg/mL oral concentrate**

For the management of severe cancer-related or chronic non-malignant pain.

Claim Note:

- Requests will not be considered for the treatment of opioid use disorder.

**METHYLPHENIDATE (BIPHENTIN)**  
**10 mg, 15 mg, 20 mg, 30 mg, 40 mg, 50 mg, 60 mg and 80 mg controlled release capsules**

For the treatment of patients with Attention Deficit Hyperactivity Disorder who have tried extended-release methylphenidate with unsatisfactory results.

Claim Note:

- The maximum dose reimbursed is 80 mg daily.

**MIDOSTAURIN (RYDAPT)**  
**25 mg capsule**

For the treatment of adult patients with newly diagnosed FMS-like tyrosine kinase 3 (FLT3)-mutated acute myeloid leukemia (AML) when used in combination with standard cytarabine and daunorubicin (7+3) induction and cytarabine consolidation chemotherapy.

Claim Notes:

- Requests for midostaurin will not be considered when used as maintenance therapy, or as part of re-induction and/or re-consolidation.
- Requests for midostaurin in combination with idarubicin containing 7+3 induction and cytarabine consolidation chemotherapy will be considered.
- Approval period: Up to 6 cycles (maximum of 2 cycles of induction and 4 cycles of consolidation).
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).



**MIGALASTAT (GALAFOLD)**  
**123 mg capsule**

For the treatment of Fabry Disease in adults with a lab-confirmed alpha-galactosidase (alpha-Gal A) mutation, determined to be amenable by an in vitro assay.

Clinical Note:

- Eligibility for the treatment of Fabry Disease is determined by the Canadian Fabry Disease Initiative. Please contact the NB Drug Plans at 1-800-332-3691 for the request form.

Claim Notes:

- Combined use of more than one disease specific therapy (i.e. enzyme replacement therapy or chaperone therapy) will not be reimbursed.
- Approval period: 1 year. Confirmation of continued response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**MIRABEGRON (MYRBETRIQ)**  
**25 mg and 50 mg extended-release tablets**

For the treatment of overactive bladder (OAB) with symptoms of urgency, urgency incontinence, and urinary frequency in patients who have an intolerance or insufficient response to an adequate trial of a regular benefit OAB drug (e.g. immediate-release oxybutynin, solifenacin or tolterodine).

Clinical Note:

- Requests for the treatment of stress incontinence will not be considered.

**MIRTAZAPINE (REMERON RD and generic brand)**  
**15 mg, 30 mg, and 45 mg orally disintegrating tablets**

For use in patients when regular mirtazapine tablets are not an option.

**MODIFIED RAGWEED POLLEN TYROSINE ADSORBATE (POLLINEX-R)**  
**105 PNU / 0.5 ml, 250 PNU / 0.5 ml, 700 PNU / 0.5 ml, 2150 PNU / 0.5 ml prefilled syringes**

For the treatment of patients with severe, seasonal (lasting two or more years) IgE dependent allergic rhinoconjunctivitis when optimal therapy (i.e. intranasal corticosteroids and H<sub>1</sub> antihistamines) and allergen avoidance have not been sufficiently effective in controlling symptoms.

Clinical Notes:

1. Treatment with ragweed pollen allergen extract must be initiated by physicians with adequate training and experience in the treatment of respiratory allergic diseases.
2. Treatment should be initiated one month before the onset of ragweed season.
3. Optimal duration of therapy is unknown; therefore, if there is no improvement in symptoms after three years, treatment should be discontinued.

**MOXIFLOXACIN (generic brands)**  
**400 mg tablet**

1. For completion of treatment initiated in the hospital setting for patients with nosocomial pneumonia, community acquired pneumonia (CAP) or acute exacerbation of chronic obstructive pulmonary disease (AECOPD).
2. For the treatment of complicated AECOPD in patients who:
  - have failed treatment with at least one first-line therapy (doxycycline, beta-lactam, trimethoprim-sulfamethoxazole, or macrolide), or
  - are intolerant or have contraindication(s) to at least two first-line therapies.
3. For the treatment of CAP in patients with radiographic confirmation of pneumonia who:
  - have failed treatment with at least one first-line therapy (macrolide, doxycycline, beta-lactams), or
  - are intolerant or have contraindication(s) to at least two first-line therapies.
4. For the treatment of pulmonary infections in patients with cystic fibrosis.
5. For the treatment of severe pneumonia in nursing home patients.
6. For the treatment of patients with complicated osteomyelitis or joint infections.

Clinical Notes:

1. If the patient has been treated with an antibiotic within the past 3 months consider an antibiotic from a different class.
2. Complicated AECOPD is defined as patients with COPD (FEV<sub>1</sub>/FVC greater than 0.7) experiencing increased sputum purulence, and with increased dyspnea or sputum volume, and one of the following:
  - FEV<sub>1</sub> less than 50% predicted

- At least 4 exacerbations per year
- Ischemic heart disease
- Home oxygen use
- Chronic oral steroid use

Claim Notes:

- Prescriptions written by infectious disease specialists, internal medicine specialists, hematologists, medical microbiologists, oncologists, oncology clinical associates, general practitioners in oncology, or respirologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Moxifloxacin is a regular benefit for Plans BV.

**Tuberculosis**

For the treatment of tuberculosis in patients who have lab-verified drug resistance or a contraindication or intolerance to first-line drugs.

Claim Notes:

- Must be prescribed by, or in consultation with, an infectious disease specialist.
- Requests will only be considered under Plan P.

**NADROPARIN (FRAXIPARIN)**

**9,500 IU/mL prefilled syringe**

**NADROPARIN (FRAXIPARIN FORTE)**

**19,000 IU/mL prefilled syringe**

1. For the treatment of venous thromboembolism (VTE) and/or pulmonary embolism (PE) for a maximum of 30 days.
2. For the extended treatment of recurrent symptomatic venous thromboembolism (VTE) that has occurred while patients are on therapeutic doses of warfarin.
3. For the prophylaxis of venous thromboembolism (VTE) up to 35 days following elective hip replacement or hip fracture surgery.
4. For the prophylaxis of VTE up to 14 days following elective knee replacement surgery.
5. For the prophylaxis of venous thromboembolism (VTE) post abdominal or pelvic surgery for management of a malignant tumour for up to 28 days.
6. For the treatment and secondary prevention of symptomatic venous thromboembolism (VTE) or pulmonary embolism (PE) for a period of up to 6 months in patients with cancer for whom warfarin therapy is not an option.

Claim Note:

- An annual quantity of 35 days of therapy is available without special authorization.

**NARATRIPTAN (AMERGE and generic brands)**

**1 mg and 2.5 mg tablets**

For the treatment of patients with acute migraine attacks who have an intolerance or insufficient response to all triptans listed as regular benefits (e.g. almotriptan, eletriptan, rizatriptan, sumatriptan, zolmitriptan).

Claim Notes:

- Coverage limited to 6 doses per month.
- Requests for patients who have more than 3 migraines a month despite migraine prophylaxis therapy will be considered for a maximum of 12 doses per month.

**NATALIZUMAB (TYSABRI)**

**300 mg / 15 mL single-use vial**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet all the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Experienced one or more disabling relapses or new MRI activity in the past year
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)
- Refractory or intolerant to at least one disease modifying therapy (e.g., interferon, glatiramer, dimethyl fumarate, teriflunomide, ocrelizumab)

Renewal Criteria:

- Evidence of continued benefit must be provided (i.e. stability or reduction in the number of relapses in the past year or stability or improvement of EDSS score obtained within the previous 90 days).

Clinical Notes:

1. Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

2. A relapse is defined as the appearance of new or worsening neurological symptoms in the absence of fever or infection, lasting at least 24 hours yet preceded by stability for at least one month and accompanied by new objective neurological findings observed through evaluation by a neurologist.

Claim Notes:

- Must be prescribed by a neurologist.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Combined use with other disease modifying therapies to treat RRMS will not be reimbursed.
- Initial approval period: 1 year.
- Renewal approval period: 2 years.

**NETUPITANT AND PALONOSETRON (AKYNZEO)  
300 mg / 0.5 mg capsule**

In combination with dexamethasone for the prevention of acute and delayed nausea and vomiting in patients receiving:

- highly emetogenic chemotherapy, or
- moderately emetogenic chemotherapy who have had inadequate symptom control using a 5-HT3 antagonist and dexamethasone in a previous cycle.

Claim Note:

- Prescriptions written by hematologists, oncologists, oncology clinical associates, or general practitioners in oncology who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.

**NICOTINE (generic brands)  
2 mg gum  
7 mg, 14 mg and 21 mg patches  
1 mg, 2 mg and 4 mg lozenges**

For smoking cessation.

Clinical Notes:

1. The patient should be participating in a form of smoking cessation counselling.
2. For information on quitting smoking or to obtain the special authorization request form, visit our website [Smoking Cessation Therapies](#).

Claim Notes:

- A maximum of 24 weeks of standard therapy (168 patches and 960 pieces of nicotine gum or nicotine lozenges) will be reimbursed annually without special authorization.
- Patients being treated within a program or clinic that participates in the Ottawa Model may be approved for additional patches based on degree of dependence (e.g. number of cigarettes smoked prior to initiating cessation therapy).
- Requests for nicotine replacement therapy (patches/gum/lozenge) for use in combination with a non-nicotine prescription smoking cessation drug (varenicline or bupropion) will not be considered.
- Requests for special authorization should be submitted on the Smoking Cessation Therapy Special Authorization Request Form.

**NILOTINIB (TASIGNA)  
150 mg capsule**

For the first-line treatment of adult patients with Philadelphia chromosome positive chronic myeloid leukemia (Ph+ CML) in chronic phase.

**NILOTINIB (TASIGNA)  
200 mg capsule**

For the treatment of chronic phase (CP) and accelerated phase (AP) Philadelphia chromosome positive (Ph+) chronic myeloid leukemia (CML) in adult patients who:

- are resistant or intolerant to imatinib,  
OR
- intolerant to dasatinib

**NINTEDANIB (OFEV)**  
**100 mg and 150 mg capsules**

**Chronic Fibrosing Interstitial Lung Diseases**

For the treatment of adult patients with chronic fibrosing interstitial lung disease (ILD) with a progressive phenotype and a forced vital capacity (FVC) greater than or equal to 45% of predicted.

Renewal Criteria:

- Patients must not demonstrate progression of disease defined as an absolute decline in percent predicted FVC of greater than or equal to 10% over the preceding 12 months of treatment with nintedanib.

Claim Notes:

- Must be prescribed by, or in consultation with a physician experienced in the treatment of ILD.
- Combination therapy of pirfenidone with nintedanib will not be reimbursed.
- Approval period: 1 year.

**Idiopathic Pulmonary Fibrosis**

For the treatment of adult patients with mild to moderate idiopathic pulmonary fibrosis (IPF) confirmed by a respirologist and a high-resolution CT scan within the previous 24 months.

Initial Renewal Criteria:

- Patients must not demonstrate progression of disease defined as an absolute decline in percent predicted forced vital capacity (FVC) of greater than or equal to 10% from initiation of therapy until renewal (initial 6 month treatment period).

Subsequent Renewal Criteria:

- Patients must not demonstrate progression of disease defined as an absolute decline in percent predicted FVC of greater than or equal to 10% within any 12 month period.

Clinical Note:

- Mild to moderate IPF is defined as a FVC greater than or equal to 50% predicted.

Claim Notes:

- Must be prescribed by, or in consultation with, physicians experienced in the treatment of IPF.
- Combination therapy of pirfenidone with nintedanib will not be reimbursed.
- Initial approval period: 7 months.
- Initial renewal approval period: 6 months.
- Subsequent renewal approval period: 1 year.

**NIRAPARIB (ZEJULA)**  
**100 mg capsule**

1. As monotherapy maintenance treatment for adult patients with newly diagnosed epithelial ovarian, fallopian tube, or primary peritoneal cancer who meet all of the following criteria:
  - Complete or partial radiologic response after at least 4 cycles of first-line platinum-based chemotherapy
  - Last cycle of platinum-based chemotherapy was completed within the previous 12 weeks
  - High-grade serous or endometrioid tumors classified as Stage III or IV according to the International Federation of Gynecology and Obstetrics (FIGO) criteria

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.
- Requests for treatment beyond 3 years will not be considered.

Clinical Notes:

1. Patients must have a good performance status and no active or uncontrolled metastases to the central nervous system.
2. Treatment should continue until unacceptable toxicity, disease progression, or completion of 3 years of therapy, whichever occurs first.

Claim Notes:

- Requests for niraparib in combination with bevacizumab will not be considered.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

2. As monotherapy maintenance treatment for adult patients with recurrent, platinum-sensitive, epithelial ovarian, fallopian tube, or primary peritoneal cancer with high grade serous or endometrioid histology who meet all of the following criteria:
- Completed at least 2 prior lines of platinum-based chemotherapy
  - Received at least 4 cycles of the most recent platinum-based chemotherapy and in complete or partial radiologic response
  - Last cycle of platinum-based chemotherapy was completed within the previous 12 weeks

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Platinum-sensitive disease is defined as disease progression occurring at least 6 months after completion of platinum-based chemotherapy.
2. Patients should have good performance status and no active or uncontrolled metastases to the central nervous system.
3. Treatment should continue until unacceptable toxicity or disease progression.

Claim Notes:

- Requests for niraparib will not be considered for patients who experience disease progression on a PARP-inhibitor or who complete treatment with a PARP-inhibitor in a prior line of therapy.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**NITISINONE (ORFADIN and generic brand)  
2 mg, 5 mg, 10 mg and 20 mg capsules**

For the treatment of adult and pediatric patients with hereditary tyrosinemia type 1 (HT-1) in combination with dietary restriction of tyrosine and phenylalanine.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician with experience in the diagnosis and management of HT-1.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**NORETHINDRONE (NORLUTATE)  
5 mg tablet**

For the treatment of abnormal uterine bleeding in patients not able to be treated with other hormonal treatments.

**NUSINERSEN (SPINRAZA)  
2.4 mg/mL intrathecal injection**

For the treatment of 5q spinal muscular atrophy (SMA), if the following criteria are met:

- Genetic documentation of 5q SMA homozygous gene deletion, homozygous mutation, or compound heterozygous mutation; and
- Patient is not requiring permanent invasive ventilation; and
- Patient who:
  - is pre-symptomatic with genetic documentation of two or three copies of the survival motor neuron 2 (SMN2) gene, or
  - has had disease duration less than 6 months, two copies of the SMN2 gene, and symptom onset after the first week of birth and on or before 7 months of age, or
  - is under the age of 18 with symptom onset after 6 months of age.

Discontinuation Criteria:

Prior to the fifth dose or every subsequent dose:

- There is failure to demonstrate achievement or maintenance of motor milestone function as assessed using age-appropriate scales since treatment initiation in patients who were pre-symptomatic at the time of treatment initiation; or
- There is failure to demonstrate maintenance in motor milestone function as assessed using age-appropriate scales since treatment initiation in patients who were symptomatic at the time of treatment initiation; or
- Permanent invasive ventilation is required.

Clinical Notes:

1. An age-appropriate scale is defined as the Hammersmith Infant Neurological Examination (HINE) Section 2, Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND), or Hammersmith Functional Motor Scale-Expanded (HFMSE).
2. A baseline assessment using an age-appropriate scale must be completed prior to initiation of nusinersen treatment.
3. Permanent invasive ventilation is defined as the use of tracheostomy and a ventilator due to progression of SMA that is not due to an identifiable and reversible cause.

Claim Notes:

- The patient must be under the care of a specialist experienced in the treatment of SMA.
- Combination therapy with risdeplam will not be reimbursed.
- Requests for nusinersen will not be considered for patients who have received adeno-associated virus (AAV) vector-based gene therapy
- Patients currently receiving SMA drug therapy may be eligible to switch to an alternate SMA drug therapy; however, patients will not be permitted to switch back to a previously trialed SMA drug.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**OBETICHOLIC ACID (OCALIVA)  
5 mg and 10 mg tablets**

For the treatment of adult patients with primary biliary cholangitis (PBC) as either:

- combination therapy with ursodeoxycholic acid (UDCA) in patients who have experienced an inadequate response to a minimum of 12 months of UDCA treatment; or
- monotherapy in patients who have experienced unmanageable intolerance to UDCA.

Requirement for Initial Requests:

- Alkaline phosphatase (ALP) and bilirubin levels prior to initiation of treatment with obeticholic acid must be provided.

Renewal Criteria:

- Requests for renewal will be considered if the patient achieved:
  - a reduction in the ALP to less than 1.67 times the upper limit of normal (ULN); or
  - at least a 15% reduction in the ALP level from baseline (i.e. prior to initiation of treatment with obeticholic acid).

Clinical Notes:

1. Diagnosis confirmed by positive antimitochondrial antibodies or liver biopsy results consistent with PBC.
2. An inadequate response is defined as:
  - ALP  $\geq$  1.67 times ULN, or
  - bilirubin  $>$  ULN and  $<$  2 times the ULN, or
  - evidence of compensated cirrhosis.
3. For patients who experience unmanageable intolerance to UDCA, details must be provided.

Claim Notes:

- Must be prescribed by, or in consultation with, a gastroenterologist, hepatologist or other physician experienced in the treatment of PBC.
- Approval period: 12 months.

**OCRELIZUMAB (OCREVUS)  
30 mg/mL single-use vial**

**Primary Progressive Multiple Sclerosis**

For the treatment of adult patients with early primary progressive multiple sclerosis (PPMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Recent Expanded Disability Status Scale (EDSS) score between 3.0 and 6.5
- Recent Functional Systems Scale (FSS) score of at least 2 for the pyramidal functions component due to lower extremity findings
- Disease duration of 10 years for those with an EDSS of less than or equal to 5 or disease duration less than 15 years for those with an EDSS greater than 5
- Diagnostic imaging features characteristic of inflammatory activity

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Note:

- Must be prescribed by a neurologist.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Approval Period: 2 years.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**Relapsing Remitting Multiple Sclerosis**

For the treatment of adult patients with relapsing remitting multiple sclerosis (RRMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Experienced one or more disabling relapses or new MRI activity in the last two years
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Notes:

- Must be prescribed by a neurologist.
- Combined use with other disease modifying therapies to treat RRMS will not be reimbursed.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Approval Period: 2 years.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**OFATUMUMAB (KESIMPTA)  
20 mg / 0.4 mL autoinjector**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Experienced one or more disabling relapses or new MRI activity in the past two years
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Notes:

- Must be prescribed by a neurologist.
- Combined use with other disease modifying therapies to treat RRMS will not be reimbursed.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Approval Period: 2 years.

**OFLOXACIN (OCUFLOX)  
0.3% ophthalmic solution**

- For the treatment of ophthalmic infections caused by susceptible bacteria.
- For the prevention of ophthalmic infections associated with non-elective eye surgery.

Claim Note:

- Prescriptions written by ophthalmologists and prescribing optometrists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.

**OLAPARIB (LYNPARZA)  
100 mg and 150 mg tablet**

**Metastatic Castration-Resistant Prostate Cancer**

For the treatment of patients with metastatic castration-resistant prostate cancer (mCRPC) who meet all of the following criteria:

- deleterious or suspected deleterious germline and/or somatic mutations in the homologous recombination repair (HRR) genes BRCA1, BRCA2 or ATM; and
- Disease progression on prior treatment with androgen-receptor-axis-targeted (ARAT) therapy.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**Ovarian Cancer**

1. As monotherapy maintenance treatment for adult patients with newly diagnosed BRCA-mutated (germline or somatic) epithelial ovarian, fallopian tube, or primary peritoneal cancer who meet all of the following criteria:
  - Complete or partial radiologic response after at least 4 cycles of first-line platinum-based chemotherapy
  - Last cycle of platinum-based chemotherapy was completed within the previous 12 weeks
  - High-grade serous or endometrioid tumors classified as Stage III or IV according to the International Federation of Gynecology and Obstetrics (FIGO) criteria

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.
- Requests for treatment beyond 2 years will not be considered if there is no evidence of disease.

Clinical Notes:

1. Patients must have a good performance status and no active or uncontrolled metastases to the central nervous system.
2. Treatment should continue until unacceptable toxicity, disease progression, or completion of 2 years of therapy, whichever occurs first.

Claim Notes:

- Requests for olaparib in combination with bevacizumab will not be considered.
- Approval period: 1 year.

2. As monotherapy maintenance treatment for patients with recurrent, platinum-sensitive, BRCA-mutated (germline or somatic) epithelial ovarian, fallopian tube, or primary peritoneal cancer with high grade serous or endometrioid histology who meet all of the following criteria:
  - Completed at least 2 previous lines of platinum-based chemotherapy
  - Received at least 4 cycles of the most recent platinum-based chemotherapy and in complete or partial radiologic response
  - Last cycle of platinum-based chemotherapy was completed within the previous 12 weeks

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Platinum-sensitive disease is defined as disease progression occurring at least 6 months after completion of platinum-based chemotherapy.
2. Patients must have a good performance status and no active or uncontrolled metastases to the central nervous system.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests for olaparib will not be considered for patients who experience disease progression on a PARP-inhibitor or who complete treatment with a PARP-inhibitor in a prior line of therapy.
- Approval period: 1 year.

**OMALIZUMAB (XOLAIR)**

**150 mg single-use vial**

**150 mg/mL prefilled syringe**

For the treatment of patients 12 years of age and older with moderate to severe chronic idiopathic urticaria (CIU) who remain symptomatic (presence of hives and/or associated itching) despite optimum management with H<sub>1</sub> antihistamines.



Requirement for Initial Requests:

- Documentation of the most recent Urticaria Activity Score over 7 days (UAS7) must be provided on the submitted request.

Renewal Criteria:

- Requests for renewal will be considered if the patient has achieved:
  - complete symptom control for less than 12 consecutive weeks; or
  - partial response to treatment, defined as at least a  $\geq 9.5$  point reduction in baseline UAS7.

Clinical Notes:

1. Moderate to severe CIU is defined as a UAS7  $\geq 16$ .
2. Treatment cessation could be considered for patients who experience complete symptom control for at least 12 consecutive weeks at the end of a 24 week treatment period.
3. In patients who discontinue treatment due to temporary symptom control, re-initiation can be considered if CIU symptoms reappear.

Claim Notes:

- Approvals will be for a maximum dose of 300mg every four weeks.
- Initial approval period: 24 weeks.

**ONABOTULINUMTOXINA (BOTOX)  
50 and 100 Allergan units per vial**

1. For the treatment of equinus foot deformity in cerebral palsy in patients 2 years of age and older.
2. To reduce the subjective symptoms and objective signs of cervical dystonia (spasmodic torticollis) in adults.
3. For the treatment of blepharospasm, hemifacial spasm (VII nerve disorder) and strabismus in patients 12 years of age and older.
4. For the treatment of upper and lower limb (at or below the knee) focal spasticity following stroke in adults. Initial approval period for focal spasticity following stroke will be 6 months.

Renewal Criteria:

- Continued approval will require documented benefit of improved passive and/or active range of motion, muscle tone, or improved gait (in the case of lower limb spasticity).
5. For the treatment of overactive bladder (OAB) with symptoms of urgency, urgency incontinence, and urinary frequency, in adult patients who have an intolerance or insufficient response to an adequate trial of at least two other pharmacologic treatments (e.g. anticholinergics, mirabegron).

Renewal Criteria:

- Requests for renewal should provide objective evidence of a treatment response, defined as a reduction of at least 50% in the frequency of urinary incontinence episodes.

Claim Notes:

- Must be prescribed and administered by a urologist.
- Initial approval period: 12 weeks (one dose).
- Renewal approval period: Maximum of 3 doses per year in responders, at a frequency of no more than once every twelve weeks.

Exclusion Criteria:

The following conditions are excluded from coverage:

- Chronic migraine
- Chronic pain
- Hyperhidrosis
- Muscle contracture for support of perineal care

**ONABOTULINUMTOXINA (BOTOX)  
200 Allergan units per vial (PIN 00999505)**

For the treatment of urinary incontinence due to neurogenic detrusor overactivity resulting from neurogenic bladder associated with multiple sclerosis (MS) or subcervical spinal cord injury (SCI) if the following conditions are met:

- patient failed to respond to behavioural modification and anticholinergics and/or is intolerant to anticholinergics
- subsequent treatments are provided at intervals no less than every 36 weeks.

Clinical Note:

- Patients who fail to respond to initial treatment with onabotulinumtoxinA should not be retreated.

**ONASEMNOGENE ABEPARVOVEC (ZOLGENSMA)**  
**2 x 10<sup>13</sup> vector genomes/mL solution for infusion**

For the treatment of spinal muscular atrophy (SMA) in individuals who meet all of the following criteria:

- Genetic documentation of 5q SMA with biallelic mutations in the survival motor neuron 1 (SMN1) gene; and
- Patient is 180 days of age or younger at the time onasemnogene abeparvec is administered; and
- Patient is pre-symptomatic or symptomatic with one to three copies of the survival motor neuron 2 (SMN2) gene; and
- Patient does not require permanent ventilatory support (invasive or non-invasive) or a permanent feeding tube.

Clinical Note:

- Permanent ventilatory support is defined as the need for a tracheostomy or requirement of 16 hours or more of respiratory assistance per day (via non-invasive ventilatory support) for 14 or more consecutive days in the absence of an acute reversible illness excluding perioperative ventilation.

Claim Notes:

- The patient must be under the care of a specialist experienced in the diagnosis and treatment of SMA.
- No treatment with nusinersen, risdiplam or other medications indicated for the treatment of SMA will be considered after the patient has received a dose of onasemnogene abeparvec.
- Approvals will be limited to one lifetime administration of 1.1 x 10<sup>14</sup> vector genomes/kg.
- Patients who have received a prior dose of onasemnogene abeparvec accessed by any mechanism (e.g. private insurance plan, clinical trial, compassionate access) will not be funded.
- Patients with 4 or more copies of the SMN2 gene will not be funded.

**ONDANSETRON (ZOFTRAN, ZOFTRAN ODT and generic brands)**  
**2 mg/mL injection**  
**4 mg / 5 mL oral solution**  
**4 mg and 8 mg tablets and orally disintegrating tablets**

1. For the prevention of nausea and vomiting in patients receiving:
  - highly or moderately emetogenic chemotherapy / radiation therapy, or
  - chemotherapy / radiation therapy who have had inadequate symptom control with other available antiemetics.

Claim Note:

- Prescription written for tablets and orally disintegrating tablets by oncologists, oncology clinical associates, or a general practitioner in oncology who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
2. For the treatment of nausea and vomiting in pediatric patients (under 18 years of age) receiving chemotherapy (e.g., methotrexate) for chronic non-oncology conditions who have experienced an episode of nausea and vomiting.
  3. For the management of nausea and vomiting in patients receiving palliative care.

**OSELTAMIVIR (TAMIFLU and generic brands)**  
**30 mg, 45 mg and 75 mg capsules**

For residents of long-term care facilities during an influenza outbreak when one of the following criteria is met:

- For treatment of long-term care residents with clinically suspected or lab confirmed influenza A or B. A clinically suspected case is one in which the patient meets the criteria of influenza-like illness and there is confirmation of influenza A or B circulating within the facility or surrounding community.
- For prophylaxis of long-term care residents where the facility has an influenza A or B outbreak. Prophylaxis should be continued until the outbreak is over. An outbreak is declared over 7 days after the onset of the last case in the facility.

Clinical Note:

- Long-term care facilities are licensed nursing homes and do not include special care homes.

Claim Notes:

- Coverage is limited to individuals enrolled in Plan V, when recommended by a Medical Officer of Health as outlined in the [policy](#).
- Oseltamivir is a regular benefit for individuals who meet eligibility criteria of the Avian Flu Drug Therapy program (Plan I), as outlined [here](#).

**OSELTAMIVIR (TAMIFLU and generic brand)  
6 mg/mL powder for suspension**

1. For residents of long-term care facilities during an influenza outbreak when oral capsules are not an option and who otherwise meet special authorization criteria for oseltamivir capsules.
2. For the prevention and treatment of avian influenza when oral capsules are not an option, for individuals who meet eligibility criteria of the Avian Flu Drug Therapy program.

Clinical Note:

- Long-term care facilities are licensed nursing homes and do not include special care homes.

Claim Notes:

- Requests will be considered for individuals enrolled in Plan V, when recommended by a Medical Officer of Health as outlined in the [policy](#).
- Requests will be considered for individuals who meet eligibility criteria of the Avian Flu Drug Therapy program (Plan I) as outlined [here](#).

**OSIMERTINIB (TAGRISSO)  
40 mg and 80 mg tablets**

**Adjuvant Non-Small Cell Lung Cancer**

For the adjuvant treatment of patients with completely resected stage IB to IIIA (AJCC 7<sup>th</sup> edition or equivalent) non-small cell lung cancer (NSCLC) whose tumours have EGFR exon 19 deletions or exon 21 (L858R) substitution mutations.

Renewal Criteria:

- Written confirmation that the patient has not experienced disease recurrence.

Clinical Notes:

- Patients must have a good performance status.
- Patients should initiate treatment within 26 weeks of complete surgical resection if treated with adjuvant chemotherapy, or within 10 weeks if chemotherapy was not given.
- Treatment should continue until disease recurrence, unacceptable toxicity, or until a maximum treatment duration of 3 years, regardless of dose reduction and dose interruption.

Claim Notes:

- Requests for treatment beyond 3 years will not be considered.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**Advanced Non-Small Cell Lung Cancer**

1. For the first-line treatment of patients with locally advanced (not amenable to curative intent therapy) or metastatic NSCLC whose tumours have EGFR exon 19 deletions or exon 21 (L858R) substitution mutations.
2. For the treatment of patients with locally advanced (not amenable to curative therapy) or metastatic EGFR T790M mutation-positive NSCLC who have progressed on EGFR tyrosine kinase inhibitor therapy.

Renewal Criteria:

- Written confirmation that the patient is responding to treatment.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

Claim Notes:

- Requests for first line therapy will be considered for patients with de novo EGFR T790M mutation-positive NSCLC.
- Requests will not be considered for patients who progress on, or within 6 months of, treatment with adjuvant EGFR targeted therapy.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**OXCARBAZEPINE (TRILEPTAL and generic brand)  
150 mg, 300 mg and 600 mg tablets  
60 mg/mL oral suspension**

For the treatment of epilepsy in patients who have had an inadequate response or are intolerant to at least 3 other

antiepileptics including carbamazepine.

**OXYBUTYNIN (DITROPAN XL)  
5 mg and 10 mg extended-release tablets**

For the treatment of overactive bladder (OAB) with symptoms of urgency, urgency incontinence, and urinary frequency in patients who have an insufficient response to an adequate trial of immediate-release oxybutynin.

Clinical Notes:

1. Requests for the treatment of stress incontinence will not be considered.
2. Not to be used in combination with other pharmacological treatments of OAB.

**OXYCODONE (OXY IR and generic brand and SUPEUDOL)  
5 mg, 10 mg and 20 mg immediate release tablets**

For the treatment of moderate to severe cancer-related or chronic non-malignant pain.

**PALBOCICLIB (IBRANCE)  
75 mg, 100 mg, and 125 mg capsules and tablets**

1. In combination with an aromatase inhibitor for the treatment of patients with hormone receptor positive, HER2 negative advanced or metastatic breast cancer who:
  - have not received prior endocrine therapy for advanced or metastatic disease, and
  - are not resistant to prior (neo)adjuvant non-steroidal aromatase inhibitor (NSAI) therapy, and
  - do not have active or uncontrolled metastases to the central nervous system.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. For patients who received (neo)adjuvant NSAI therapy, a minimum disease-free interval of twelve months after stopping therapy is required.
2. Pre- and peri-menopausal patients must be treated with a luteinizing hormone-releasing hormone agonist.
3. Patients must have a good performance status.
4. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will be considered for patients who have received up to one prior chemotherapy for advanced or metastatic disease.
  - Approval period: 1 year.
2. In combination with fulvestrant for the treatment of patients with hormone receptor positive, HER2 negative advanced or metastatic breast cancer who:
    - have not received prior endocrine therapy or have experienced disease progression on endocrine therapy, and
    - have received up to one prior chemotherapy for advanced or metastatic disease, and
    - do not have active or uncontrolled metastases to the central nervous system.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Pre- and peri-menopausal patients must be treated with a luteinizing hormone-releasing hormone agonist.
2. Patients must have a good performance status.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will not be considered for patients who experience disease progression on a CDK4/6 inhibitor, fulvestrant or everolimus.
- Approval period: 1 year.

**PALIPERIDONE (INVEGA SUSTENNA)  
50 mg / 0.5 mL, 75 mg / 0.75 mL, 100 mg/mL and 150 mg / 1.5 mL prefilled syringes**

For the maintenance treatment of schizophrenia and related psychotic disorders (not dementia related) in patients who:

- are not adherent to an oral antipsychotic, or

- are currently receiving a long-acting injectable antipsychotic and require an alternative long-acting injectable antipsychotic.

Claim Note:

- Approval period: Long term.

**PALIPERIDONE PALMITATE (INVEGA TRINZA)**

**175 mg / 0.875 mL, 263 mg / 1.315 mL, 350 mg / 1.75 mL, 525 mg / 2.625 mL prefilled syringes**

For the maintenance treatment of schizophrenia and related psychotic disorders (not dementia related) in patients who have been stabilized on therapy with injectable paliperidone for at least four months.

Claim Note:

- Approval period: Long term.

**PATISIRAN (ONPATTRO)**

**2 mg/mL vial**

For the treatment of polyneuropathy in adult patients with hereditary transthyretin-mediated amyloidosis (hATTR) who meet all of the following criteria:

- Confirmed genetic diagnosis of hATTR
- Symptomatic early-stage neuropathy
- Does not have New York Heart Association class III or IV heart failure
- Has not previously undergone a liver transplant

Discontinuation Criteria:

- The patient is permanently bedridden and dependent on assistance for basic activities of daily living, or
- The patient is receiving end-of-life care.

Clinical Note:

- Symptomatic early stage neuropathy is defined as Polyneuropathy disability stage I to IIIB or Familial amyloidotic polyneuropathy stage I or II.

Claim Notes:

- The patient must be under the care of a physician with experience in the diagnosis and management of hATTR.
- Combination therapy with other interfering ribonucleic acid drugs or transthyretin stabilizers used to treat hATTR will not be reimbursed.
- Initial approval period: 9 months.
- Renewal approval period: 12 months. Confirmation of continued response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**PAZOPANIB (VOTRIENT and generic brand)**

**200 mg tablet**

For the treatment of patients with advanced or metastatic renal cell carcinoma when used as:

- first-line therapy, or
- second-line therapy following disease progression on nivolumab and ipilimumab combination therapy.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**PEGFILGRASTIM (FULPHILA, LAPELGA, NYVEPRIA, ZIEXTENZO)**

**6 mg / 0.6 mL prefilled syringe**

For the prevention of febrile neutropenia in patients receiving myelosuppressive chemotherapy with curative intent who:

- are at high risk of febrile neutropenia due to chemotherapy regimen, co-morbidities or pre-existing severe neutropenia; or

- have had an episode of febrile neutropenia, neutropenic sepsis or profound neutropenia in a previous cycle of chemotherapy; or
- have had a dose reduction, or treatment delay greater than one week due to neutropenia.

Clinical Note:

- Patients with non-curative cancer receiving chemotherapy with palliative intent are not eligible for coverage of pegfilgrastim for prevention of febrile neutropenia.

**PEGINTERFERON-BETA 1A (PLEGRIDY)  
63 mcg / 0.5 mL, 94 mcg / 0.5 mL, and 125 mcg / 0.5 mL prefilled syringes and pen**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Has experienced one or more disabling relapses of MS in the past two years
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Notes:

- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Prescriptions written by neurologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Combined use with other disease modifying therapies to treat RRMS will not be reimbursed.
- Approval Period: 2 years

**PERAMPANEL (FYCOMPA)  
2 mg, 4 mg, 6 mg, 8 mg, 10 mg, 12 mg tablets**

For the adjunctive treatment of refractory partial-onset seizures or primary generalized tonic-clonic seizures in patients who are currently receiving two or more antiepileptic drugs, and who have had an inadequate response to at least three other antiepileptic drugs.

Claim Notes:

- The patient must be under the care of a physician experienced in the treatment of epilepsy.

**PILOCARPINE (SALAGEN)  
5 mg tablet**

- For the treatment of the symptoms of xerostomia (dry mouth) due to salivary gland hypofunction caused by radiotherapy for cancer of the head and neck.
- For the treatment of the symptoms of xerostomia (dry mouth) and xerophthalmia (dry eyes) in patients with Sjögren's syndrome.

**PIRFENIDONE (ESBRIET and generic brands)  
267 mg capsule  
267 mg and 801 mg tablets**

For the treatment of adult patients with mild to moderate idiopathic pulmonary fibrosis (IPF) confirmed by a respirologist and a high-resolution CT scan within the previous 24 months.

Initial Renewal Criteria:

Patients must not demonstrate progression of disease defined as an absolute decline in percent predicted forced vital capacity (FVC) of  $\geq 10\%$  from initiation of therapy until renewal (initial 6 month treatment period). If a patient has experienced progression as defined above, then the results should be validated with a confirmatory pulmonary function test conducted 4 weeks later.

Subsequent Renewal Criteria:

Patients must not demonstrate progression of disease defined as an absolute decline in percent predicted FVC of  $\geq 10\%$  within any 12 month period. If a patient has experienced progression as defined above, then the results should be validated with a confirmatory pulmonary function test conducted 4 weeks later.

Clinical Notes:

1. Mild to moderate IPF is defined as a FVC  $\geq 50\%$  predicted.
2. All other causes of restrictive lung disease (e.g. collagen vascular disorder or hypersensitivity pneumonitis) should be excluded before initiating treatment.

Claim Notes:

- Must be prescribed by, or in consultation with, physicians experienced in the treatment of IPF.
- Combination therapy of pirfenidone with nintedanib will not be reimbursed.
- Initial approval period: 7 months (allow 4 weeks for repeat pulmonary function tests).
- Initial renewal approval period: 6 months.
- Subsequent renewal approval period: 12 months.

**PLERIXAFOR (MOZOBIL)**  
**24 mg / 1.2 mL solution for injection**

For use in combination with filgrastim to mobilize hematopoietic stem cells for subsequent autologous transplantation in patients who meet one of the following criteria:

- PBCD34+ count of less than 10 cells/ $\mu$ L after 4 days of filgrastim, or
- Less than 50% of the target CD34+ yield is achieved on the first day of apheresis (after being mobilized with filgrastim alone or following chemotherapy), or
- Failed a previous attempt for stem cell mobilization with filgrastim alone or following chemotherapy.

Claim Notes:

- Reimbursement is limited to a maximum of 4 doses (0.24mg/kg given daily) for a single mobilization attempt and to prescriptions written by an oncologist or hematologist.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**POMALIDOMIDE (POMALYST)**  
**1 mg, 2 mg, 3 mg and 4 mg capsules**

For the treatment of patients with relapsed and/or refractory multiple myeloma who:

- Have previously failed at least two treatments including both bortezomib and lenalidomide, and
- Demonstrated disease progression on the last treatment.

Clinical Note:

- Requests for pomalidomide will be considered in rare instances where bortezomib is contraindicated or when patients are intolerant to it; however, in all cases patients should have failed lenalidomide which they may have received in the maintenance setting.

**PONATINIB (ICLUSIG)**  
**15 mg and 45 mg film-coated tablets**

For the treatment of patients with chronic, accelerated or blast phase chronic myelogenous leukemia (CML) or Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ ALL) who have:

- resistance or intolerance to two or more tyrosine kinase inhibitors (TKIs), or
- confirmed T315i mutation positive disease.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have an ECOG performance status of 0-2.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**PRASUGREL (generic brand)**  
**10 mg tablet**

In combination with ASA for patients with:

- unstable angina (UA) or non-ST-segment elevation myocardial infarction (NSTEMI) managed with percutaneous coronary intervention (PCI); or
- ST-segment elevation myocardial infarction (STEMI) managed with primary or delayed PCI; or
- failure on clopidogrel and ASA therapy as defined by definite stent thrombosis, or recurrent STEMI, NSTEMI or UA after revascularization with PCI.

Clinical Note:

- Definite stent thrombosis, according to the Academic Research Consortium, is a total occlusion originating in or within 5 mm of the stent or is a visible thrombus within the stent or is within 5 mm of the stent in the presence of an acute ischemic clinical syndrome within 48 hours.

Claim Notes:

- Prescriptions written by cardiologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Approval period: 1 year.

**PROPIVERINE (MICTORYL PEDIATRIC)  
5 mg tablet**

For the treatment of overactive bladder with symptoms of urgency incontinence and/or urinary frequency and urgency in pediatric patients under 18 years of age.

**PROPRANOLOL (HEMANGIOL)  
3.75 mg/mL oral solution**

For the treatment of patients with proliferating infantile hemangioma that is:

- Life- or function-threatening, or
- Ulcerated with pain or not responding to simple wound care measures, or
- At risk of permanent scarring or disfigurement

**RANIBIZUMAB (BYOOVIZ)  
10 mg/mL solution for intravitreal injection**

1. For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD).
2. For the treatment of patients with choroidal neovascularization secondary to pathologic myopia (PM).
3. For the treatment of patients with choroidal neovascularization secondary to ocular conditions other than AMD and PM.
3. For the treatment of patients with diabetic macular edema (DME).
4. For the treatment of macular edema secondary to central retinal vein occlusion (CRVO) or branch retinal vein occlusion (BRVO).

Claim Notes:

- An initial claim of up to two vials (1 vial per eye treated) will be automatically reimbursed when prescribed by a New Brunswick ophthalmologist. If continued treatment is required, a request must be made through special authorization.
- Approvals will be for a maximum of 1 vial per eye every 30 days.
- Approval Period: 1 year.

**REGORAFENIB (STIVARGA)  
40 mg film-coated tablet**

**Advanced Hepatocellular Carcinoma**

For the second-line treatment of patients with unresectable hepatocellular carcinoma who meet all of the following criteria:

- Disease progression on sorafenib or lenvatinib
- Child-Pugh class status of A
- ECOG performance status of 0 or 1

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Note:

- Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Patients with disease progression on sorafenib must have tolerated a minimum dose of 400 mg per day for at least 20 of the last 28 days of treatment.
- Requests for regorafenib will not be considered for patients who experience disease progression on cabozantinib or atezolizumab in combination with bevacizumab.
- Initial approval period: 4 months.
- Renewal approval period: 6 months.



### **Gastrointestinal Stromal Tumour**

For the treatment of patients with metastatic and/or unresectable gastrointestinal stromal tumours (GIST) who have had disease progression on, or intolerance to, imatinib and sunitinib, and who have an ECOG performance status of 0 or 1.

#### **Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

#### **Clinical Note:**

- Treatment should be discontinued upon disease progression or unacceptable toxicity.

#### **Claim Note:**

- Approval period: 6 months.

### **RIBAVIRIN (IBAVYR) 200mg tablets**

For use in combination with other drugs for the treatment of chronic hepatitis C. The applicable criteria for the combination regimen must be met.

#### **Claim note:**

- Must be prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating a patient with hepatitis C infection).
- Requests will be considered for individuals enrolled in Plans ACDEFGV.

### **RIBOCICLIB (KISQALI) 200 mg tablet**

1. In combination with an aromatase inhibitor for the treatment of patients with hormone receptor positive, HER2 negative advanced or metastatic breast cancer who:
  - have not received prior endocrine therapy for advanced or metastatic disease, and
  - are not resistant to prior (neo)adjuvant non-steroidal aromatase inhibitor (NSAI) therapy, and
  - do not have active or uncontrolled metastases to the central nervous system.

#### **Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

#### **Clinical Notes:**

1. For patients who received (neo)adjuvant NSAI therapy, a minimum disease-free interval of twelve months after stopping therapy is required.
2. Pre- and peri-menopausal patients must be treated with a luteinizing hormone-releasing hormone agonist.
3. Patients must have a good performance status.
4. Treatment should be discontinued upon disease progression or unacceptable toxicity.

#### **Claim Notes:**

- Requests will be considered for patients who have received up to one prior chemotherapy for advanced or metastatic disease.
  - Approval period: 1 year.
2. In combination with fulvestrant for the treatment of patients with hormone receptor positive, HER2 negative advanced or metastatic breast cancer who:
    - have not received prior endocrine therapy or have experienced disease progression on endocrine therapy, and
    - have received up to one prior chemotherapy for advanced or metastatic disease, and
    - do not have active or uncontrolled metastases to the central nervous system.

#### **Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

#### **Clinical Notes:**

1. Pre- and peri-menopausal patients must be treated with a luteinizing hormone-releasing hormone agonist.
2. Patients must have a good performance status.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will not be considered for patients who experience disease progression on a CDK4/6 inhibitor, fulvestrant or everolimus.
- Approval period: 1 year.

**RIFABUTIN (MYCOBUTIN)**  
**150 mg capsule**

For the prevention of disseminated Mycobacterium avium complex (MAC) disease in patients with advanced HIV infection.

Claim Notes:

- Must be prescribed by, or in consultation with, an infectious disease specialist.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.

**RIFAXIMIN (ZAXINE)**  
**550 mg tablet**

For reducing the risk of overt hepatic encephalopathy (HE) recurrence in patients who have had two or more episodes and are unable to achieve adequate control of HE with maximum tolerated doses of lactulose alone.

Clinical Note:

- Must be used in combination with lactulose unless lactulose is not tolerated.

**RIOCIGUAT (ADEMPAS)**  
**0.5 mg, 1 mg, 1.5 mg, 2 mg, and 2.5 mg film-coated tablets**

For the treatment of inoperable chronic thromboembolic pulmonary hypertension (CTEPH) World Health Organization [WHO] Group 4) or persistent or recurrent CTEPH after surgical treatment in adult patients (18 years of age or older) with WHO Functional Class II or III pulmonary hypertension.

Clinical Note:

- Requests will be considered from physicians with experience in the diagnosis and treatment of CTEPH.

Claim Note:

- Approval period: 1 year.

**RISANKIZUMAB (SKYRIZI)**  
**75 mg / 0.83 mL prefilled syringe**  
**150 mg/mL autoinjector and prefilled syringe**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals, or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 150 mg at weeks 0 and 4, then every 12 weeks thereafter.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**RISDIPLAM (EVRYSDI)**  
**60 mg powder for oral solution**

For the treatment of 5q spinal muscular atrophy (SMA), if the following criteria are met:

- Genetic documentation of 5q SMA homozygous gene deletion, or compound heterozygous mutation; and

- Patient is not requiring permanent invasive ventilation; and
- Patient who is symptomatic with two or three copies of the SMN2 gene and is:
  - 2 months to 7 months of age, or
  - 8 months to 25 years of age and non-ambulatory.

**Discontinuation Criteria:**

- There is failure to demonstrate maintenance in motor milestone function as assessed using age-appropriate scales since treatment initiation; or
- permanent invasive ventilation is required.

**Clinical Notes:**

1. An age-appropriate scale is defined as the Hammersmith Infant Neurological Examination (HINE) Section 2, Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND), or Hammersmith Functional Motor Scale-Expanded (HFMSE).
2. A baseline assessment using an age-appropriate scale must be completed prior to initiation of treatment.
3. Yearly assessments must be completed using an age-appropriate scale no more than 12 weeks prior to the renewal date.
4. Permanent invasive ventilation is defined as the use of tracheostomy and a ventilator due to progression of SMA that is not due to an identifiable and reversible cause.

**Claim Notes:**

- The patient must be under the care of a specialist experienced in the treatment of SMA.
- Combination therapy with nusinersen will not be reimbursed.
- Requests for risdiplam will not be considered for patients who have received adeno-associated virus (AAV) vector-based gene therapy.
- Patients currently receiving SMA drug therapy may be eligible to switch to an alternate SMA drug therapy; however, patients will not be permitted to switch back to a previously trialed SMA drug.
- Approvals will be for a maximum of 0.2 mg/kg/day for patients 2 months to less than 2 years of age, 0.25 mg/kg/day for patients greater than or equal to 2 years of age weighing less than 20 kg, or 5 mg/day for patients greater than or equal to 2 years of age and weighing greater than or equal to 20 kg.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**RISEDRONATE (generic brand)  
30 mg film-coated tablet**

For the treatment of Paget's disease.

**Claim Notes:**

- A maximum of 60 tablets will be reimbursed annually without special authorization.
- Requests for re-treatment may be considered through special authorization following a two month post-treatment observation period.

**RISPERIDONE (RISPERDAL CONSTA)  
12.5 mg, 25 mg, 37.5 mg and 50 mg vials**

For the treatment of patients who are:

- not adherent to an oral antipsychotic, or
- currently receiving a long-acting injectable antipsychotic and require an alternative long-acting injectable antipsychotic.

**Claim Notes:**

- Requests will not be considered for the treatment of psychotic symptoms related to dementia.
- Approval period: Long term.

**RITUXIMAB (RIXIMYO, RUXIENCE, TRUXIMA)  
10 mg/mL vial**

For the treatment of patients with rheumatoid arthritis, vasculitis, or other autoimmune disease.

**Claim Notes:**

- Must be prescribed by a specialist.
- Initial approval period: 6 months.
- Renewal approval period: Long term. Confirmation of response is required.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**RIVAROXABAN (XARELTO)  
2.5 mg tablet**

For use in combination with acetylsalicylic acid (75 mg to 100 mg) for the prevention of atherothrombotic events in patients with concomitant coronary artery disease (CAD) and peripheral artery disease (PAD) who meet the following criteria:

- CAD defined as having one or more of the following:
  - Myocardial infarction within the last 20 years
  - Multi-vessel CAD with symptoms or history of angina
  - Multi-vessel percutaneous coronary intervention
  - Multi-vessel coronary artery bypass graft surgery
- PAD defined as having one or more of the following:
  - Previous aorto-femoral bypass surgery, limb bypass surgery, or percutaneous transluminal angioplasty revascularization of the iliac or infrainguinal arteries
  - Previous limb or foot amputation for arterial vascular disease
  - History of intermittent claudication and one or more of the following: an ankle-brachial index of less than 0.90 or peripheral artery stenosis greater than 50% as documented by angiography or duplex ultrasound
  - Previous carotid revascularization or asymptomatic carotid artery stenosis greater than or equal to 50% diagnosed by angiography or duplex ultrasound

Clinical Notes:

1. Atherothrombotic events include stroke, myocardial infarction, cardiovascular death, acute limb ischemia and mortality
2. Multivessel CAD is defined as stenosis of more than 50% in two or more coronary arteries, or in one coronary artery territory if at least one other territory has been revascularized

Claim Note:

- The maximum dose of rivaroxaban that will be reimbursed is 2.5 mg twice daily.

**RIVAROXABAN (XARELTO)  
10 mg tablet**

**Venous thromboembolic event prophylaxis following total knee or total hip replacement surgery**

- For the prevention of venous thromboembolic events in patients who have undergone elective total knee replacement (TKR) surgery or total hip replacement (THR) surgery.

Clinical Note:

- The total duration of therapy includes the period during which doses are administered post-operatively in an acute care (hospital) setting, and the approval period is for the balance of the total duration after discharge.

Claim Notes:

- Maximum reimbursement without special authorization will be limited to 14 days of therapy (14 tablets) for TKR or 35 days of therapy (35 tablets) for THR, within a 6 month period.
- Subsequent reimbursement for prophylaxis within a 6 month period (i.e. second joint replacement procedure within the 6 month period) will require special authorization.

**RIVASTIGMINE (EXELON and generic brands)  
1.5 mg, 3 mg, 4.5 mg and 6 mg capsules**

For the treatment of patients with mild to moderate dementia who have had an intolerance to donepezil and who meet the following criteria:

- Mini-Mental State Exam (MMSE) score of 10 to 30
- Functional Assessment Staging Test (FAST) score of 4 to 5

Clinical Notes:

1. Requests must contain an updated MMSE and FAST score completed within 6 months of the request.
2. The nature of the intolerance must be described.

Claim Note:

- Approval period: 1 year.

**RIVASTIGMINE (EXELON)  
2 mg/mL oral solution**

For the treatment of patients with mild to moderate dementia for whom oral tablets or capsules are not an option and who meet the following criteria:

- Mini-Mental State Exam (MMSE) score of 10 to 30
- Functional Assessment Staging Test (FAST) score of 4 to 5

Clinical Note:

- Requests must contain an updated MMSE and FAST score completed within 6 months of the request.

Claim Note:

- Approval period: 1 year.

**ROTIGOTINE (NEUPRO)**  
**2 mg, 4 mg, 6 mg, 8 mg transdermal patch**

For adjunctive treatment of patients with advanced stage Parkinson's disease who are currently receiving a levodopa-decarboxylase inhibitor combination.

**RUFINAMIDE (BANZEL)**  
**100 mg, 200 mg and 400 mg film-coated tablets**

For the adjunctive treatment of seizures associated with Lennox-Gastaut syndrome for patients who meet all of the following criteria:

- are under the care of a physician experienced in treating Lennox-Gastaut syndrome-associated seizures,  
AND
- are currently receiving two or more antiepileptic drugs,  
AND
- in whom less costly antiepileptic drugs are ineffective or not appropriate.

**RUXOLITINIB (JAKAVI)**  
**5 mg, 10 mg, 15 mg, 20 mg tablets**

**Myelofibrosis**

For the treatment of splenomegaly and/or disease-related symptoms in adult patients with primary myelofibrosis, post-polycythemia vera myelofibrosis, or post-essential thrombocythemia myelofibrosis who meet all of the following criteria:

- Intermediate to high risk disease, or low risk disease with symptomatic splenomegaly, as assessed using DIPSS Plus
- Previously untreated or refractory to other treatment

Renewal Criteria:

- Confirmation that the patient has responded to treatment as evidenced by a reduction in spleen size or symptom improvement.

Clinical Notes:

1. Patients must have an ECOG performance status of less than or equal to 3.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will not be considered for patients who experience disease progression following treatment with fedratinib.
- Approval period: 6 months

**Polycythemia Vera**

For the treatment of patients with polycythemia vera who have demonstrated resistance or intolerance to hydroxyurea (HU).

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.
3. Resistance is considered if, after at least 3 months of HU therapy at the maximum tolerated dose, patients experience at least one of the following:
  - Need for phlebotomy to maintain hematocrit (HCT) < 45%
  - Uncontrolled myeloproliferation (i.e. platelet count > 400 x 10<sup>9</sup>/L and white blood cell count > 10 x 10<sup>9</sup>/L)
  - Failure to reduce massive splenomegaly by greater than 50%, as measured by palpation
4. Intolerance to HU is considered if patients experience at least one of the following:
  - Absolute neutrophil count < 1.0 x 10<sup>9</sup>/L, platelet count < 100 x 10<sup>9</sup>/L or hemoglobin < 100 g/L at the lowest dose of HU required to achieve a response. A response to HU is defined as HCT < 45% without phlebotomy, and/or all of the following: platelet count ≤ 400 x 10<sup>9</sup>/L, white blood cell count ≤ 10 x 10<sup>9</sup>/L, and non-palpable spleen.

- Presence of leg ulcers or other unacceptable HU-related non-hematological toxicities (defined as grade 3 or 4 or, more than one week of grade 2) such as mucocutaneous manifestations, gastrointestinal symptoms, pneumonitis, or fever
- Toxicity requiring permanent discontinuation of HU, interruption of HU until toxicity resolved, or hospitalization due to HU toxicity

Claim Notes:

- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**SACUBITRIL AND VALSARTAN (ENTRESTO)**

**24 mg / 26 mg, 49 mg / 51 mg and 97 mg / 103 mg film-coated tablets**

For the treatment of patients with New York Heart Association (NYHA) class II or III heart failure to reduce the incidence of cardiovascular death and heart failure hospitalization who meet all of the following criteria:

- Left ventricular ejection fraction (LVEF) of less than or equal to 40%.
- NYHA class II to III symptoms despite at least four weeks of treatment of the following:
  - a stable dose of an angiotensin-converting enzyme inhibitor (ACEI) or an angiotensin II receptor antagonist (ARB); and
  - a stable dose of a beta-blocker and other recommended therapies, including an aldosterone antagonist.
- Plasma B-type natriuretic peptide (BNP)  $\geq$  150 pg/mL or N-terminal prohormone B-type natriuretic peptide (NT-proBNP)  $\geq$  600 pg/mL.

Clinical Notes:

1. A plasma BNP  $\geq$  100 pg/mL or NT-proBNP  $\geq$  400 pg/mL will be considered if the patient has been hospitalized for heart failure within the past 12 months.
2. For patients who have not received four weeks of therapy with a beta blocker or aldosterone antagonist due to an intolerance or contraindication, details must be provided.

**SALBUTAMOL (VENTOLIN and generic brands)**

**0.5 mg/mL, 1 mg/mL, 2 mg/mL and 5 mg/mL solution for inhalation**

For patients who have tried using an inhaler with spacer device and

- Are unable to follow instructions, hold the spacer device or hold the device long enough to actuate it due to cognitive or physical limitations; or
- Have difficulty generating adequate inspiratory effort to achieve therapeutic benefit.

Claim Notes:

- Initial approval period: 1 year.
- Renewal approval period: Long term.

**SALBUTAMOL AND IPRATROPIUM BROMIDE (generic brands)**

**2.5 mg / 0.5 mg / 2.5 mL solution for inhalation**

For patients who have tried using an inhaler with spacer device and

- are unable to follow instructions, hold the spacer device or hold the device long enough to actuate it due to cognitive or physical limitations; or
- have difficulty generating adequate inspiratory effort to achieve therapeutic benefit.

Claim Notes:

- Initial approval period: 1 year.
- Renewal approval period: Long term.

**SAPROPTERIN (KUVAN)**

**100 mg tablet**

**100 mg and 500 mg sachet**

For the ongoing treatment of hyperphenylalaninemia due to tetrahydrobiopterin (BH4)-responsive phenylketonuria (PKU) in conjunction with a phenylalanine (Phe)-restricted diet in patients who meet all of the following criteria:

- Confirmed diagnosis based on genetic testing.
- Response to Kuvan as demonstrated by a Kuvan responsiveness test.
- Baseline blood Phe levels greater than 360  $\mu$ mol/L despite compliance with a low protein diet and formulas (non-pregnant patients require at least 2 baseline levels and pregnant patients require at least 1 baseline level during a 3 to 6 month time frame).
- Achievement of the following during a 6-month trial of treatment:
  - For pregnant or non-pregnant patients, normal sustained blood Phe levels of 120  $\mu$ mol/L to 360  $\mu$ mol/L; or
  - For non-pregnant patients, sustained blood Phe reduction of at least 30% compared to baseline if the baseline blood Phe level is less than 1200  $\mu$ mol/L; or

- For non-pregnant patients, sustained blood Phe reduction of at least 50% compared to baseline if the baseline blood Phe level is greater than 1200 umol/L.
- For non-pregnant patients, documented increase in dietary protein tolerance based on targets set between the clinician and patient.

**Renewal Criteria:**

- Confirmation of continued response to Kuvan based on Phe levels achieved during the 6-month trial. Two Phe levels taken at least 1 month apart must be provided.

**Clinical Notes:**

1. Patients must be initiated on treatment and followed in a specialized clinic with expertise in the diagnosis and management of PKU.
2. Phe blood levels and Phe tolerance levels must be provided.
3. Pregnant patients who have maintained a decrease in Phe levels below 360 umol/L during the 6-month trial period will be eligible for coverage of Kuvan for the duration of the pregnancy.

**Claim Notes:**

- Approvals will be for a maximum of 20mg/kg per day.
- Renewals for Kuvan in pregnant patients will not be considered.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SARILUMAB (KEVZARA)  
150 mg / 1.14 mL and 200 mg / 1.14 mL prefilled pen**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

**Clinical Notes:**

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

**Claim Notes:**

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 200 mg every other week.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**SATRALIZUMAB (ENSPRYNG)  
120 mg/mL prefilled syringe**

For the treatment of patients 12 years of age and older with neuromyelitis optica spectrum disorder (NMOSD) who meet all of the following criteria:

- Aquaporin-4 antibody positive
- Expanded Disability Status Scale (EDSS) score of 6.5 points or less
- Experienced at least one relapse in the previous 12 months
- Relapse occurred despite an adequate trial of rituximab, or there has been an intolerance to rituximab

**Renewal Criteria:**

- Requests for renewal will be considered for patients who maintain an EDSS score of less than 8 points.

**Clinical Note:**

- Satralizumab should not be initiated during a NMOSD relapse.

Claim Notes:

- Must be prescribed by a neurologist with experience in the treatment of NMOSD.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 120 mg at week 0, 2 and 4, then 120 mg every four weeks thereafter.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SAXAGLIPTIN (ONGLYZA and generic brands)  
2.5 mg and 5 mg tablets**

For the treatment of type 2 diabetes mellitus when added to metformin and a sulfonylurea for patients with inadequate glycemic control on metformin and a sulfonylurea and in whom insulin is not an option.

Clinical Note:

- For patients who cannot take metformin and/or a sulfonylurea due to contraindications or intolerances, details must be provided.

**SEBELIPASE ALFA (KANUMA)  
20 mg vial**

For the treatment of patients with lysosomal acid lipase (LAL) deficiency. For the complete criteria, please contact the NB Drug Plans at 1-800-332-3691.

Claim Note:

- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SECUKINUMAB (COSENTYX)  
150 mg/mL autoinjector and prefilled syringe**

**Ankylosing Spondylitis**

- For the treatment of patients with moderate to severe ankylosing spondylitis (e.g., Bath AS Disease Activity Index (BASDAI) score greater than or equal to 4 on 10-point scale) who:
  - have axial symptoms and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each or in whom NSAIDs are contraindicated, or
  - have peripheral symptoms and who have failed to respond, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 2 weeks each and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.
- Requests for renewal must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease of at least 2 points on the BASDAI scale, compared with the pre-treatment score, or
  - patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as HAQ or “ability to return to work”).

Clinical Note:

- Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease do not require a trial of NSAIDs.

Claim Notes:

- Must be prescribed by a rheumatologist or internist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 150 mg given at weeks 0, 1, 2, 3, and 4, then monthly.
- Requests for 300 mg monthly will be considered for patients who continue to have active disease while on the recommended monthly maintenance dose of 150 mg.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**Plaque Psoriasis**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals, or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.



2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for 300 mg given at weeks 0, 1, 2, 3, and 4, then monthly.
- Initial approval period: 12 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Psoriatic Arthritis**

- For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each.
- For the treatment of patients with predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each; and
  - methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 8 weeks; and
  - leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 150 mg given at weeks 0, 1, 2, 3, and 4, then monthly.
- Requests for 300 mg given at weeks 0, 1, 2, 3, and 4 then monthly will be considered for patients who have previously had an inadequate response to TNF-inhibitors.
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**SELEXIPAG (UPTRAVI)**

**200 mcg, 400 mcg, 600 mcg, 800 mcg, 1000 mcg, 1200 mcg, 1400 mcg, and 1600 mcg tablets**

For the treatment of pulmonary arterial hypertension (PAH) in patients with World Health Organization functional class II to IV, if the following clinical criteria are met:

- Inadequate control with a first-line (i.e. phosphodiesterase-5 inhibitor) and second-line (i.e. endothelin receptor antagonist) PAH therapy.
- Diagnosis of PAH should be confirmed by right heart catheterization.

Claim Notes:

- Combination therapy with prostacyclin or prostacyclin analogs will not be reimbursed.
- Must be prescribed by a clinician with experience in the diagnosis and treatment of PAH.

**SEMAGLUTIDE (OZEMPIC)**

**2 mg / 1.5 mL and 4 mg / 3 mL prefilled pens**

For the treatment of type 2 diabetes mellitus when added to:

- metformin for patients who have inadequate glycemic control on metformin; or
- metformin and a sulfonylurea for patients who have inadequate glycemic control on metformin and a sulfonylurea.

Clinical Note:

- For patients who cannot take metformin due to contraindications or intolerances, details must be provided.

Claim Note:

- Approvals will be for a maximum of 1 prefilled pen every 4 weeks.

**SEVELAMER (RENVELA)  
0.8 g and 2.4 g sachets**

For use in patients who have difficulty swallowing tablets.

Claim Note:

- Approval Period: 1 year

**SILDENAFIL (REVATIO and generic brands)  
20 mg film-coated tablet**

For the treatment of patients with Group 1 pulmonary arterial hypertension (PAH) with World Health Organization (WHO) functional class II, III or IV.

Clinical Note:

- The diagnosis of PAH should be confirmed by right heart catheterization.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of PAH.
- The maximum dose of sildenafil that will be reimbursed is 20 mg three times daily.
- Approval period: Long term.

**SIPONIMOD (MAYZENT)  
0.25 mg and 2 mg tablets**

For the treatment of patients with active secondary progressive multiple sclerosis (SPMS) who meet all of the following criteria:

- History of relapsing-remitting multiple sclerosis and current active SPMS
- Recent Expanded Disability Status Scale (EDSS) score of 3.0 to 6.5

Clinical Notes:

1. Active SPMS is defined as having had relapses in the past 2 years and/or having at least one T1 gadolinium-enhancing lesion prior to treatment initiation with siponimod.
2. Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

Claim Notes:

- Must be prescribed by a neurologist.
- Approvals will be for a maximum of 2 mg daily.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Approval Period: 2 years.

**SODIUM BICARBONATE (generic brands)  
500 mg tablets**

For the treatment of metabolic acidosis in patients with chronic kidney disease who have a serum bicarbonate (CO<sub>2</sub>) < 22mmol/L.

**SODIUM FERRIC GLUCONATE COMPLEX (FERRLECIT)  
12.5 mg/mL ampoule and vial**

For the treatment of iron deficiency anemia in patients who

- are intolerant to oral iron replacement products, or
- have not responded to an adequate trial of oral iron.

**SODIUM PHENYLBUTYRATE (PHEBURANE)  
483 mg/g coated granules**

For the treatment of patients with urea cycle disorders (UCDs).

Clinical Note:

- Diagnosis must be confirmed by blood, enzymatic, biochemical or genetic testing.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of UCDs.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SODIUM PHENYLBUTYRATE AND URSODOXICOLTAURINE (ALBRIOZA)  
3 g / 1 g powder for suspension**

For the treatment of patients with definite amyotrophic lateral sclerosis (ALS) who meet all the following criteria:

- Forced vital capacity (FVC) greater than or equal to 60% of predicted
- ALS symptoms for 18 months or less
- Permanent non-invasive or invasive ventilation is not required

Discontinuation Criteria:

- The patient requires permanent non-invasive or invasive ventilation; or
- The patient becomes non-ambulatory and is unable to cut food and feed themselves without assistance, irrespective of whether a gastrostomy is in place.

Clinical Note:

- FVC must be provided with initial request.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician with experience in the diagnosis and management of ALS.
- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SOFOSBUVIR (SOVALDI)  
400 mg tablet**

For treatment-naïve or treatment-experienced adult patients with chronic hepatitis C virus (HCV) who meet the following criteria:

<b>Approval Period and Regimen</b>	
<b>Genotype 2</b> <ul style="list-style-type: none"> <li>• Without cirrhosis</li> <li>• With compensated cirrhosis</li> </ul>	12 weeks in combination with ribavirin (RBV)
<b>Genotype 3</b> <ul style="list-style-type: none"> <li>• Without cirrhosis</li> <li>• With compensated cirrhosis</li> </ul>	24 weeks in combination with RBV

The following information is also required:

1. Lab-confirmed hepatitis C genotype 2 and 3
2. Quantitative HCV RNA value within the last 6 months
3. Fibrosis stage

Clinical Notes:

1. Treatment-experienced is defined as a patient who has been previously treated with a peginterferon/ribavirin regimen and has not experienced an adequate response.
2. Acceptable methods for the measurement of fibrosis score include Fibrotest, liver biopsy, transient elastography (FibroScan®), serum biomarker panels (such as AST-to-Platelet Ratio Index or Fibrosis-4 score) either alone or in combination.
3. Compensated cirrhosis is defined as a CTP score of 5 to 6 (Class A) and decompensated cirrhosis as a CTP score of 7 or above (Class B or C).
4. Re-treatment for direct-acting antiviral failures will be considered on a case-by-case basis under the formulary exception process.

Claim Notes:

- Must be prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating a patient with hepatitis C infection).
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SOFOSBUVIR AND LEDIPASVIR (HARVONI)  
400 mg / 90 mg tablet**

For treatment-naïve or treatment-experienced adult patients with chronic hepatitis C virus (HCV) who meet the following criteria:

<b>Approval Period and Regimen</b>	
<b>Genotype 1</b> <ul style="list-style-type: none"> <li>Treatment-naïve without cirrhosis, who have pre-treatment HCV RNA level &lt; 6 million IU/mL and mono-HCV infected only</li> </ul>	8 weeks
<b>Genotype 1</b> <ul style="list-style-type: none"> <li>Treatment-naïve without cirrhosis, who have pre-treatment HCV RNA level ≥ 6 million IU/mL</li> <li>Treatment-naïve with compensated cirrhosis</li> <li>Treatment-naïve with advanced liver fibrosis (Fibrosis stage F3-F4)</li> <li>Treatment-experienced without cirrhosis</li> <li>HCV/HIV co-infected without cirrhosis or with compensated cirrhosis</li> </ul>	12 weeks
<b>Genotype 1</b> <ul style="list-style-type: none"> <li>Treatment-experienced with compensated cirrhosis</li> </ul>	24 weeks
<b>Genotype 1</b> <ul style="list-style-type: none"> <li>Decompensated cirrhosis</li> <li>Liver transplant recipients without cirrhosis or with compensated cirrhosis</li> </ul>	12 weeks in combination with ribavirin

The following information is also required:

1. Lab-confirmed hepatitis C genotype 1
2. Quantitative HCV RNA value within the last 6 months
3. Fibrosis stage

**Clinical Notes:**

1. Treatment-experienced is defined as a patient who has been previously treated with a peginterferon/ribavirin regimen, including regimens containing HCV protease inhibitors and who has not experienced an adequate response.
2. Acceptable methods for the measurement of fibrosis score include Fibrotest, liver biopsy, transient elastography (FibroScan®), serum biomarker panels (such as AST-to-Platelet Ratio Index or Fibrosis-4 score) either alone or in combination.
3. Compensated cirrhosis is defined as a CTP score of 5 to 6 (Class A) and decompensated cirrhosis as a CTP score of 7 or above (Class B or C).

**Claim Notes:**

- Must be prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating a patient with hepatitis C infection).
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SOFOSBUVIR AND VELPATASVIR (EPCLUSA)  
400 mg / 100 mg tablet**

For treatment-naïve or treatment-experienced adult patients with chronic hepatitis C virus (HCV) who meet the following criteria:

<b>Approval Period and Regimen</b>	
<b>Genotypes 1, 2, 3, 4, 5, 6 or mixed genotypes</b> <ul style="list-style-type: none"> <li>Patients with compensated cirrhosis</li> <li>Patients without cirrhosis</li> </ul>	12 weeks
<b>Genotypes 1, 2, 3, 4, 5, 6 or mixed genotypes</b> <ul style="list-style-type: none"> <li>Patients with decompensated cirrhosis</li> </ul>	12 weeks in combination with ribavirin

The following information is also required:

1. Lab-confirmed hepatitis C genotype 1, 2, 3, 4, 5, 6 or mixed genotypes
2. Quantitative HCV RNA value within the last 6 months
3. Fibrosis stage

Clinical Notes:

1. Treatment-experienced is defined as a patient who has been previously treated with a peginterferon/ribavirin regimen, including regimens containing HCV protease inhibitors and who has not experienced an adequate response.
2. Acceptable methods for the measurement of fibrosis score include Fibrotest, liver biopsy, transient elastography (FibroScan®), serum biomarker panels (such as AST-to-Platelet Ratio Index or Fibrosis-4 score) either alone or in combination.
3. Compensated cirrhosis is defined as a CTP score of 5 to 6 (Class A) and decompensated cirrhosis as a CTP score of 7 or above (Class B or C).

Claim Notes:

- Must be prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating a patient with hepatitis C infection).
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SOFOBUVIR, VELPATASVIR AND VOXILAPREVIR (VOSEVI)  
400 mg / 100 mg / 100 mg tablet**

For treatment-experienced adult patients with chronic hepatitis C virus (HCV) who meet the following criteria:

	<b>Approval Period</b>
<b>Genotypes 1, 2, 3, 4, 5, 6 or mixed genotypes</b> <ul style="list-style-type: none"><li>• Patients with compensated cirrhosis</li><li>• Patients without cirrhosis</li></ul>	12 weeks

The following information is also required:

- Lab-confirmed hepatitis C genotype 1, 2, 3, 4, 5, 6 or mixed genotypes.
- Quantitative HCV RNA value within the last 6 months.

Clinical Notes:

1. Treatment experienced is defined as a patient who has been previously treated with an NS5A inhibitor for genotype 1, 2, 3, 4, 5 or 6 or sofosbuvir without an NS5A inhibitor for genotype 1, 2, 3 or 4 and who has not experienced an adequate response.
2. Compensated cirrhosis is defined as a CTP score of 5 to 6 (Class A).

Claim Notes:

- Must be prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating a patient with hepatitis C infection).
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**SOMATROPIN (GENOTROPIN)  
0.6 mg, 0.8 mg, 1 mg, 1.2 mg, 1.4 mg, 1.6 mg, 1.8 mg, and 2 mg MiniQuick® prefilled syringes  
5.3 mg, and 12 mg GoQuick® prefilled pens**

**1. Growth Hormone Deficiency in Children**

For the treatment of growth hormone deficiency in children under the age of 19.

Claim Notes:

- Must be prescribed by, or in consultation with, an endocrinologist.
- Somatropin is a regular benefit for Plan T

**2. Turner Syndrome**

For the treatment of short stature associated with Turner Syndrome in patients whose epiphyses are not closed.

Claim Note:

- Must be prescribed by, or in consultation with, an endocrinologist.

**SOMATROPIN (HUMATROPE)**  
**6 mg, 12 mg and 24 mg cartridges**

**1. Growth Hormone Deficiency in Children**

For the treatment of growth hormone deficiency in children under the age of 19.

Claim Notes:

- Must be prescribed by, or in consultation with, an endocrinologist.
- Somatropin is a regular benefit for Plan T.

**2. Turner Syndrome**

For the treatment of short stature associated with Turner Syndrome in patients whose epiphyses are not closed.

Claim Note:

- Must be prescribed by, or in consultation with, an endocrinologist.

**SOMATROPIN (NORDITROPIN NORDIFLEX)**  
**5 mg / 1.5 mL, 10 mg / 1.5 mL and 15 mg / 1.5 mL prefilled pens**

**Growth Hormone Deficiency in Children**

For the treatment of growth hormone deficiency in children under the age of 19.

Claim Notes:

- Must be prescribed by, or in consultation with, an endocrinologist.
- Somatropin is a regular benefit for Plan T

**SOMATROPIN (NUTROPIN AQ NuSpin)**  
**5 mg / 2 mL, 10 mg / 2 mL, and 20 mg / 2 mL prefilled cartridges**  
**SOMATROPIN (SAIZEN)**  
**5 mg vials**  
**6 mg, 12 mg and 20 mg cartridges**

**1. Growth Hormone Deficiency in Children**

For the treatment of growth hormone deficiency in children under the age of 19.

Claim Notes:

- Must be prescribed by, or in consultation with, an endocrinologist.
- Somatropin is a regular benefit for Plan T.

**2. Turner Syndrome**

For the treatment of short stature associated with Turner Syndrome in patients whose epiphyses are not closed.

Claim Note:

- Must be prescribed by, or in consultation with, an endocrinologist.

**3. Chronic Renal Insufficiency**

For the treatment of children with growth failure associated with chronic renal insufficiency, up to the time of renal transplantation, who meet the following criteria:

- A glomerular filtration rate less than or equal to 1.25 mL/s/1.73m<sup>2</sup> (75 mL/min/1.73m<sup>2</sup>)
- Evidence of growth impairment:
  - Z score (HSDS) less than -1.88 (HSDS = height standard deviation score, a statistical comparison to the average of normal values for age and sex) or height-for-age at the 3rd percentile
  - OR
  - Height velocity-for-age SDS less than -1.88 or height velocity-for-age less than 3<sup>rd</sup> percentile, persisting for greater than 3 months despite treatment of nutritional deficiencies and metabolic abnormalities.

Claim Note:

- Somatropin must be prescribed by, or in consultation with, a specialist in pediatric nephrology.

**SOMATROPIN (OMNITROPE)**  
**5 mg / 1.5 mL, 10 mg / 1.5 mL and 15 mg / 1.5 mL cartridges**

For the treatment of growth hormone deficiency in children under the age of 19.

Claim Notes:

- Must be prescribed by, or in consultation with, an endocrinologist.
- Somatropin is a regular benefit for Plan T.

**SORAFENIB (NEXAVAR)  
200 mg film-coated tablet**

**Advanced Hepatocellular Carcinoma**

For the treatment of unresectable hepatocellular carcinoma, as first-line or second-line therapy after progression on atezolizumab in combination with bevacizumab, for patients who meet all of the following criteria:

- Child-Pugh class status of A
- ECOG performance status of 0-2
- Progressed on trans-arterial chemoembolization (TACE) or not suitable for the TACE procedure

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Claim Notes:**

- Requests for sorafenib will not be considered for patients who have progressed on lenvatinib.
- Approval period: 6 months.

**Metastatic Renal Cell Carcinoma (MRCC)**

For the treatment of patients with advanced or metastatic renal cell carcinoma when used as a second-line therapy following disease progression on cytokine therapy.

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Clinical Notes:**

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Claim Note:**

- Approval period: 1 year.

**STIRIPENTOL (DIACOMIT)  
250 mg and 500 mg capsules  
250 mg and 500 mg powder for suspension**

For use in combination with clobazam and valproate as adjunctive therapy of refractory generalized tonic-clonic seizures in patients with severe myoclonic epilepsy in infancy (Dravet syndrome), whose seizures are not adequately controlled with clobazam and valproate alone.

**Clinical Note:**

- The patient must be under the care of a neurologist or a pediatrician.

**SUCROFERRIC OXYHYDROXIDE (VELPHORO)  
500 mg iron chewable tablet**

For the treatment of hyperphosphatemia (serum phosphate greater than 1.8 mmol/L) in patients with end-stage renal disease who are on dialysis.

**Claim Note:**

- Approval period: Long term.

**SUMATRIPTAN (IMITREX NASAL SPRAY)  
5 mg and 20 mg nasal sprays**

For the treatment of patients with acute migraine attacks who have an intolerance or insufficient response to oral triptans listed as regular benefits.

**Claim Notes:**

- Coverage limited to 6 doses per month.
- Requests for patients who have more than 3 migraines a month despite migraine prophylaxis therapy will be considered for a maximum of 12 doses per month.

**SUMATRIPTAN (IMITREX INJECTION and generic brand)  
6 mg / 0.5 mL prefilled syringe**

For the treatment of patients with acute migraine attacks who have had an insufficient response to oral and nasal triptans, or nausea and/or vomiting precludes their use.

Claim Notes:

- Coverage limited to 6 doses per month.
- Requests for patients who have more than 3 migraines a month despite migraine prophylaxis therapy will be considered for a maximum of 12 doses per month.

**SUNITINIB (SUTENT and generic brand)  
12.5 mg, 25 mg and 50 mg capsules**

**Gastrointestinal Stromal Tumour**

For the treatment of patients with unresectable or metastatic gastrointestinal stromal tumour who experience disease progression on, or intolerance to, imatinib.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Note:

- Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 6 months.

**Metastatic Renal Cell Carcinoma**

For the treatment of patients with advanced or metastatic renal cell carcinoma when used as:

- first-line therapy, or
- second-line therapy following disease progression on nivolumab and ipilimumab combination therapy.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**Pancreatic Neuroendocrine Tumours**

For the treatment of patients with progressive, unresectable, locally advanced or metastatic, well or moderately differentiated pancreatic neuroendocrine tumours.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**TACROLIMUS (PROTOPIC)  
0.03% ointment**

For children over 2 years of age with refractory atopic dermatitis.

Claim Note:

- Approvals will be given for up to twelve months at a time.



**TACROLIMUS (PROTOPIC)****0.1% ointment**

For the treatment of adults with moderate to severe atopic dermatitis who have failed or are intolerant to a site appropriate strength of corticosteroid therapy (i.e. low potency for the face versus intermediate to high potency for the trunk and extremities).

**TAFAMIDIS (VYNDAMAX)****61 mg capsule****TAFAMIDIS MEGLUMINE (VYNDAQEL)****20 mg capsule**

For the treatment of cardiomyopathy in adult patients with documented hereditary or wild-type transthyretin-mediated amyloidosis (ATTR) who meet all of the following criteria:

- New York Heart Association (NYHA) class I to III heart failure
- At least one prior hospitalization for heart failure or clinical evidence of heart failure that required treatment with a diuretic
- Has not previously undergone a heart or liver transplant
- Does not have an implanted cardiac mechanical assist device (CMAD)

**Discontinuation Criteria:**

The patient has:

- NYHA class IV heart failure, or
- received an implanted CMAD, or
- received a heart or liver transplant.

**Clinical Notes:**

1. Wild-type ATTR-cardiomyopathy (CM) consists of all of the following:
  - absence of a variant transthyretin (TTR) genotype
  - TTR precursor protein identification by immunohistochemistry, scintigraphy, or mass spectrometer
  - evidence of cardiac involvement by echocardiography with end-diastolic interventricular septal wall thickness greater than 12 mm
  - positive findings on technetium-99m pyrophosphate (Tc-99m-PYP) scintigraphy with single-photon emission computerized tomography (SPECT) scanning or presence of amyloid deposits in biopsy tissue (fat aspirate, salivary gland, median nerve connection tissue sheath, or cardiac tissue)
2. Hereditary ATTR-CM consists of all of the following:
  - presence of a variant TTR genotype associated with CM and presenting with a CM phenotype
  - evidence of cardiac involvement by echocardiography with end-diastolic interventricular septal wall thickness greater than 12 mm
  - positive findings on technetium-99m pyrophosphate (Tc-99m-PYP) scintigraphy with single-photon emission computerized tomography (SPECT) scanning or presence of amyloid deposits in biopsy tissue (fat aspirate, salivary gland, median nerve connective tissue sheath, or cardiac tissue)

**Claim Notes:**

- The patient must be under the care of a physician with experience in the diagnosis and treatment of ATTR-CM.
- Combination therapy with other interfering ribonucleic acid drugs or transthyretin stabilizers used to treat ATTR-CM will not be reimbursed.
- Initial approval period: 9 months.
- Renewal approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**TALIGLUCERASE ALFA (ELELYSO)****200 units per vial**

For the treatment of patients with symptomatic Gaucher disease type 1 (GD1) for whom treatment with velaglucerase alfa is not tolerated or contraindicated.

**Clinical Notes:**

1. Velaglucerase alfa is the preferred reimbursed enzyme replacement therapy for GD1. Requests for patients currently using taliglucerase alfa who do not have a contraindication or intolerance to velaglucerase alfa will be considered for coverage of velaglucerase alfa only.
2. Requests for coverage must meet the criteria for diagnosis of GD1, indication for therapy and expected response to enzyme replacement therapy. These criteria are consistent with the Ontario Guidelines for the Treatment of Gaucher Disease. Please contact the NB Drug Plans at 1-800-332-3691 for the criteria.

**Claim Notes:**

- Approvals will be for a maximum of 60 units/kg every 2 weeks.
- Initial approval period: 6 months.

- Renewal approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

### **TAZAROTENE (TAZORAC CREAM)**

**0.05% cream**

### **TAZAROTENE (TAZORAC GEL)**

**0.1% gel**

For the treatment of patients with plaque psoriasis in whom conventional therapies have been ineffective or are inappropriate.

### **TEDUGLUTIDE (REVESTIVE)**

**5 mg vial**

For the ongoing treatment of patients with Short Bowel Syndrome (SBS) as a result of major intestinal resection (e.g. volvulus, vascular disease, cancer, Crohn's disease, injury, congenital disease) who meet the following criteria:

- For pediatric patients:
  - Cumulative lifetime duration of parenteral support (PS) must be at least 12 months
  - PS must provide more than 30% of caloric and/or fluid and electrolyte needs
  - Prior to initiating teduglutide, PS frequency and volume must be stable for at least three months or there must be no improvement in enteral feeding for at least three months
- For adult patients:
  - Dependency on parenteral support (PS) for a least 12 months
  - Prior to initiating teduglutide, PS required at least three times weekly to meet caloric, fluid and electrolyte needs and stable PS frequency and volume for at least one month

A request for coverage for continued treatment will be considered if the patient has achieved at least a 20% reduction in PS volume compared to baseline, while on teduglutide therapy.

Renewal Criteria:

- Has maintained at least a 20% reduction in PS volume from baseline at 12 months.

#### Clinical Note:

- PS is defined as parenteral nutrition which encompasses parenteral delivery of lipids, protein and/or carbohydrates to address caloric needs, and/or intravenous fluids which addresses fluid and electrolyte needs of patients

#### Claim Notes:

- Must be prescribed by a gastroenterologist or an internal medicine specialist with a specialty in gastroenterology.
- Approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

### **TERIFLUNOMIDE (AUBAGIO and generic brands)**

**14 mg film-coated tablet**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Experienced one or more disabling relapses or new MRI activity in the past two years
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)

#### Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

#### Claim Notes:

- Prescriptions written by neurologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Combined use with other disease modifying therapies to treat RRMS will not be reimbursed.
- Requests will be considered for individuals enrolled in Plans ACDEFGV.
- Approval Period: 2 years

**TESTOSTERONE (ANDROGEL, TESTIM and generic brand)  
1% gel (2.5 g and 5 g packets)  
TESTOSTERONE UNDECANOATE (generic brands)  
40 mg capsule**

For the treatment of congenital and acquired primary or secondary hypogonadism in males with a specific diagnosis of:

- Primary: cryptorchidism, Klinefelter's, orchiectomy, and other established causes
- Secondary: Pituitary-hypothalamic injury due to tumours, trauma, radiation

Testosterone deficiency should be clearly demonstrated by clinical features and confirmed by two separate free testosterone measurements before initiating any replacement therapy

Clinical Note:

- Older males with non-specific symptoms of fatigue, malaise, or depression who have low testosterone levels do not satisfy these criteria.

**THYROTROPIN (THYROGEN)  
0.9 mg/mL vial**

1. For on-going evaluation in patients who have documented evidence of thyroid cancer, have undergone appropriate surgical and/or medical management, and require monitoring for recurrence and metastatic disease. This includes:
  - The patient has failed to respond to, or relapsed during:
    - Primary use in patients with inability to raise an endogenous TSH level ( $\geq 25$  mu/L) with thyroid hormone withdrawal.
    - Primary use in patients with one of the following documented comorbidities in whom severe hypothyroidism could be life threatening:
      - unstable angina
      - recent myocardial infarction
      - class III-IV congestive heart failure
      - uncontrolled psychiatric illness
      - other medical condition in which the clinical course could lead to a potential life threatening situation
    - Secondary use in patients with previous thyroid hormone withdrawal resulting in a documented life threatening event.
2. As an adjunctive treatment as pre-therapeutic stimulation for radioiodine ablation of thyroid tissue remnants in patients maintained on thyroid hormone suppression therapy who have undergone near-total or total thyroidectomy for well-differentiated thyroid cancer without evidence of distant metastatic thyroid cancer.

**TICAGRELOR (generic brands)  
60 mg tablet**

In combination with ASA for patients with a history of ST elevation myocardial infarction (STEMI) or non-ST elevation acute coronary syndrome (NSTEMI) in the previous 3 years who are at high risk for subsequent cardiovascular events.

Clinical Note:

- High risk for subsequent cardiovascular events is defined as age 65 years or older, diabetes, second prior spontaneous myocardial infarction, multivessel coronary artery disease, or chronic renal dysfunction (creatinine clearance  $< 60$  mL/min).

Claim Notes:

- Prescriptions written by cardiologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Approval period: 3 years.

**TICAGRELOR (BRILINTA and generic brands)  
90 mg tablet**

1. In combination with ASA for patients with ST elevation myocardial infarction (STEMI) or non-ST elevation acute coronary syndrome (NSTEMI) who receive percutaneous coronary intervention (PCI).

Claim Notes:

- Prescriptions written by cardiologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Approval period: 1 year.

2. For the treatment of patients who have recurrent cardiovascular events (STEMI or NSTEMI), or definite stent thrombosis, while on clopidogrel and ASA therapy.

Clinical Note:

- Definite stent thrombosis, according to the Academic Research Consortium, is a total occlusion originating in or within 5 mm of the stent or is a visible thrombus within the stent or is within 5 mm of the stent in the presence of an acute ischemic clinical syndrome within 48 hours.

Claim Notes:

- Prescriptions written by cardiologists who are licensed by the College of Physicians and Surgeons of New Brunswick do not require special authorization.
- Approval period: Long term.

**TIGECYCLINE (TYGACIL)**  
**50 mg vial**

For the treatment of patients with multi-drug resistant infections when alternative agents are not an option.

Claim Note:

- Must be prescribed by, or in consultation with, an infectious disease specialist or medical microbiologist.

**TINZAPARIN (INNOHEP)**  
**10,000 IU/mL multidose vials and prefilled syringes**  
**20,000 IU/mL multidose vials and prefilled syringes**

1. For the treatment of venous thromboembolism (VTE) and/or pulmonary embolism (PE) for a maximum of 30 days.
2. For the extended treatment of recurrent symptomatic venous thromboembolism (VTE) that has occurred while patients are on therapeutic doses of warfarin.
3. For the prophylaxis of venous thromboembolism (VTE) up to 35 days following elective hip replacement or hip fracture surgery.
4. For the prophylaxis of VTE up to 14 days following elective knee replacement surgery.
5. For the prophylaxis of venous thromboembolism (VTE) post abdominal or pelvic surgery for management of a malignant tumour for up to 28 days.
6. For the treatment and secondary prevention of symptomatic venous thromboembolism (VTE) or pulmonary embolism (PE) for a period of up to 6 months in patients with cancer for whom warfarin therapy is not an option.

Claim Note:

- An annual quantity of 35 days of therapy is available without special authorization.

**TILDRAKIZUMAB (ILUMYA)**  
**100 mg/mL prefilled syringe**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended dose and for duration of treatment specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum of 100 mg at weeks 0 and 4, then every 12 weeks thereafter.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of response is required.

**TIPRANA VIR (APTIVUS)**  
**250 mg capsule**

For the treatment of adult patients with HIV-1 infection who are treatment experienced, have demonstrated failure to

multiple protease inhibitors and in whom no other protease inhibitor is a treatment option.

**TOBRAMYCIN (TOBI PODHALER)  
28 mg powder for inhalation**

For the treatment of chronic pulmonary *Pseudomonas aeruginosa* infections, when used as a cyclic treatment, in patients with cystic fibrosis.

Clinical Note:

- Cyclic treatment measured in 28-day cycles is defined as 28 days of treatment, followed by 28 days without treatment.

Claim Note:

- Combined use of tobramycin either concurrently or for antibiotic cycling during off-treatment periods, with other inhaled antibiotics (e.g. aztreonam, levofloxacin) will not be reimbursed.
- Requests will be considered for individuals enrolled in Plans ABCDEFGV.

**TOCILIZUMAB (ACTEMRA)  
80 mg / 4 mL, 200 mg / 10 mL, and 400 mg / 20 mL single-use vial  
162 mg / 0.9 mL autoinjector and prefilled syringe**

**Giant Cell Arteritis**

- For the treatment of adult patients with new onset or relapsed giant cell arteritis (GCA) in combination with oral glucocorticoids.
- Requests for renewal must include:
  - confirmation of response to treatment (e.g. absence of flares, normalization of C-reactive protein), and
  - description of attempts to taper or discontinue glucocorticoids, and
  - rationale for the need for ongoing treatment.

Clinical Note:

- A flare is defined as the recurrence of signs or symptoms and/or erythrocyte sedimentation rate greater than or equal to 30 mm/hour.

Claim Notes:

- Must be prescribed by, or in consultation with, a rheumatologist or other physician experienced in the treatment of GCA.
- Combined use of more than one biologic drug will not be reimbursed.
- Subcutaneous injection: Approvals will be for up to 162 mg every week.
- Approval period: 1 year

**Polyarticular Juvenile Idiopathic Arthritis**

For the treatment of children (age 2-17) with moderately to severely active polyarticular juvenile idiopathic arthritis (pJIA) who have had inadequate response to one or more disease-modifying antirheumatic drugs (DMARDs).

Claim Notes:

- Must be prescribed by, or in consultation with, a rheumatologist who is familiar with the use of biologic DMARDs in children.
- Combined use of more than one biologic drug will not be reimbursed.
- Intravenous infusion: Approvals will be for 10mg/kg for patients less than 30kg or 8mg/kg for patients greater than or equal to 30kg, to a maximum of 800mg, administered every four weeks.
- Subcutaneous injection: Approvals will be for a maximum of 162 mg once every three weeks for patients weighing less than 30 kg or 162 mg once every two weeks for patients weighing more than 30 kg.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.

2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Intravenous infusion: Initial approvals will be for 4mg/kg/dose every four weeks, with a maximum maintenance dose escalation up to 8mg/kg, to a maximum of 800mg per infusion for patients greater than 100kg.
- Subcutaneous injection: Initial approvals will be for 162mg every other week for patients less than 100kg, with a maximum maintenance dose escalation to weekly dosing permitted. Patients greater than or equal to 100kg will be approved for 162mg every week, with no dose escalation permitted.
- Initial approval period: 16 weeks
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Systemic Juvenile Idiopathic Arthritis**

For the treatment of active systemic juvenile idiopathic arthritis (sJIA), in patients 2 years of age or older, who have responded inadequately to non-steroidal anti-inflammatory drugs (NSAIDs) and systemic corticosteroids (with or without methotrexate) due to intolerance or lack of efficacy.

Claim Notes:

- Must be prescribed by, or in consultation with, a rheumatologist, who is familiar with the use of biologic DMARDs in children.
- Combined use of more than one biologic drug will not be reimbursed.
- Intravenous infusion: Approvals will be for 12 mg/kg for patients less than 30kg or 8 mg/kg for patients greater than or equal to 30kg, to a maximum of 800mg, administered every two weeks.
- Subcutaneous injection: Approvals will be for a maximum of 162 mg once every three weeks for patients weighing less than 30 kg or 162 mg once every two weeks for patients weighing more than 30 kg.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**TOFACITINIB (XELJANZ and generic brands and XELJANZ XR)  
5 mg and 10 mg film-coated tablet  
11 mg extended-release tablet**

**Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, alone or in combination with methotrexate, in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another disease-modifying antirheumatic drug (DMARD), at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and
- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum dose of 5 mg twice daily (Xeljanz) or 11 mg once daily (Xeljanz XR).
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of continued response is required.

### **Ulcerative Colitis**

- For the treatment of patients with moderately to severely active ulcerative colitis who have a partial Mayo score greater than 4, and a rectal bleeding subscore greater than or equal to 2 and are:
  - refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks, and prednisone greater than or equal to 40mg daily for two weeks or IV equivalent for one week); or
  - corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year).
- Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease in the partial Mayo score greater than or equal to 2 from baseline, and
  - a decrease in the rectal bleeding subscore greater than or equal to 1.

#### Clinical Notes:

1. Consideration will be given for patients who have not received a four week trial of aminosalicylates if disease is severe (partial Mayo score greater than 6).
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of the intolerance(s) must be clearly documented.

#### Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for a maximum dose of 10 mg twice daily (Xeljanz).
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year.

### **TOPIRAMATE (TOPAMAX) 15 mg and 25 mg sprinkle capsules**

For patients who cannot take the tablet form of topiramate and require sprinkle capsules for proper administration.

### **TRAMETINIB (MEKINIST) 0.5 mg and 2 mg tablets**

#### **Adjuvant Melanoma**

In combination with dabrafenib for the adjuvant treatment of patients with cutaneous melanoma who meet all of the following criteria:

- Stage IIIA (limited to lymph node metastases of greater than 1 mm) to stage IIID disease (AJCC 8<sup>th</sup> edition)
- BRAF V600-mutation positive
- Completely resected disease including in-transit metastases

#### Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should continue until disease recurrence, unacceptable toxicity, or up to a maximum of 12 months.

#### Claim Notes:

- Requests will be considered for patients with regional lymph nodes with micrometastases after sentinel lymph node biopsy.
- Requests will not be considered for patients who received adjuvant immunotherapy for greater than three months. Patients may switch to BRAF targeted therapy within the first three months of initiating immunotherapy to complete a total of 12 months of adjuvant treatment.
- Approval period: Up to 12 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

#### **Metastatic Melanoma**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used alone or in combination with dabrafenib.

#### Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

#### Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Trametinib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy.
- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**TREPROSTINIL (REMODULIN)  
1 mg/mL, 2.5 mg/mL, 5 mg/mL and 10 mg/mL multi-use vials**

For the treatment of patients with Group 1 pulmonary arterial hypertension (PAH) with World Health Organization (WHO) functional class III or IV who have failed to respond to non-prostanoid therapies.

Clinical Note:

- The diagnosis of PAH should be confirmed by right heart catheterization.

Claim Notes:

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of PAH.
- Approval period: Long term.

**TRIAMCINOLONE HEXACETONIDE (TRISPAN)  
20 mg/mL suspension for injection**

For the treatment of juvenile idiopathic arthritis.

**TRIENTINE (generic brands)  
250 mg capsule**

For the treatment of patients with Wilson's disease (WD) who are intolerant, or have contraindications, to d-penicillamine.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment. Supporting documentation must be provided.

Clinical Note:

- Details of d-penicillamine intolerances and/or contraindications must be provided.

Claim Notes:

- In adult patients, trientine therapy must be initiated by a clinician experienced in the management of WD.
- In pediatric patients, initiation and renewal of trientine therapy must be overseen by a clinician experienced in the management of WD.
- Approvals will be for a maximum of 2000 mg per day.
- Approval period: 1 year.

**TRIFLURIDINE / TIPIRACIL (LONSURF)  
15 mg / 6.14 mg and 20 mg / 8.19 mg tablets**

For the treatment of adult patients with metastatic gastric cancer or adenocarcinoma of the gastroesophageal junction who meet the following criteria:

- Previously treated with at least two prior lines of chemotherapy including a fluoropyrimidine, a platinum, and either a taxane or irinotecan and if appropriate, with HER2-targeted therapy
- ECOG performance status of 0 or 1

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Trifluridine / tipiracil should be used in combination with best supportive care.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will be considered for patients who have an intolerance or contraindication to platinum-based therapy.
- Approval period: 6 months.



**TROSPIUM (TROSEC and generic brand)  
20 mg tablet**

For the treatment of overactive bladder (OAB) with symptoms of urgency, urgency incontinence, and urinary frequency in patients who have an intolerance or insufficient response to an adequate trial of a regular benefit OAB drug (e.g. immediate-release oxybutynin, solifenacin or tolterodine).

Clinical Notes:

1. Requests for the treatment of stress incontinence will not be considered.
2. Not to be used in combination with other pharmacological treatments of OAB.

**TUCATINIB (TUKYSA)  
50 mg and 150 mg film-coated tablets**

In combination with trastuzumab and capecitabine for the treatment of patients with locally advanced unresectable or metastatic HER2-positive breast cancer who have received prior treatment with trastuzumab, pertuzumab and a HER2-targeted antibody-drug conjugate (e.g., Kadcyla, Enhertu), where at least one was given in the advanced or metastatic setting.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression, unacceptable toxicity, or if both trastuzumab and capecitabine are discontinued.

Claim Note:

- Approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**UPADACITINIB (RINVOQ)  
15 mg extended-release tablets**

**Psoriatic Arthritis**

- For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each.
- For the treatment of patients with predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - the sequential use of at least two NSAIDs at maximal tolerated dose for a minimum of two weeks each; and
  - methotrexate (oral or parenteral) at a dose of greater than or equal to 20mg weekly (greater than or equal to 15mg if patient is greater than or equal to 65 years of age) for a minimum of 8 weeks; and
  - leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use with other biologic drugs will not be reimbursed.
- Approvals will be for a maximum of 15 mg daily.
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, alone or in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory, intolerant or have contraindications to:

- methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks; and

- methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Optimal treatment response to DMARDs may take up to 24 weeks, however coverage can be considered if no improvement is seen after 12 weeks of triple DMARD use.
3. For patients who have intolerances preventing the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
4. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
5. Intolerant is defined as demonstrating serious adverse effects. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use with other biologic drugs or janus kinase inhibitors will not be reimbursed.
- Approvals will be for a maximum of 15 mg daily.
- Initial approval period: 6 months.
- Renewal approval period: 1 year. Confirmation of response is required.

**USTEKINUMAB (STELARA)  
45 mg / 0.5 mL and 90 mg/mL prefilled syringes**

For the treatment of patients with chronic moderate to severe plaque psoriasis who meet all of the following criteria:

- Psoriasis Area Severity Index (PASI) greater than 10 and Dermatology Life Quality Index (DLQI) greater than 10, or major involvement of visible areas, scalp, genitals, or nails
- Refractory, intolerant or unable to access phototherapy
- Refractory, intolerant or have contraindications to methotrexate (oral or parenteral) at a dose of greater than or equal to 20 mg weekly (greater than or equal to 15 mg if patient is greater than or equal to 65 years of age) for a minimum of 12 weeks

Clinical Notes:

1. For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a dermatologist.
- Combined use of more than one biologic drug will not be reimbursed.
- Approvals will be for 90 mg given at weeks 0, 4 and 16, then every 12 weeks thereafter
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**VALGANCICLOVIR (VALCYTE)  
50 mg/mL oral suspension**

For the prevention and treatment of cytomegalovirus (CMV) in patients for whom oral tablets are not an option.

**VANDETANIB (CAPRELSA)  
100 mg and 300 mg tablets**

For the treatment of symptomatic and/or progressive medullary thyroid cancer (MTC) in patients with unresectable locally advanced or metastatic disease.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**VARENICLINE (CHAMPIX and generic brands)  
0.5 mg and 1 mg tablets**

For smoking cessation in adults 18 years of age and older.

Clinical Notes:

1. The patient should be participating in a form of smoking cessation counselling.
2. For information on quitting smoking visit our website [Smoking Cessation Therapies](#).

Claim Notes:

- A maximum of 24 weeks of standard therapy (336 tablets) will be reimbursed annually without special authorization. Special authorization requests for additional tablets will not be considered.
- Requests for nicotine replacement therapy (patches/gum/lozenge) for use in combination with a non-nicotine prescription smoking cessation drug (varenicline or bupropion) will not be considered.

**VEDOLIZUMAB (ENTYVIO)  
108 mg / 0.68 mL prefilled syringe and prefilled pen  
300 mg vial**

**Crohn's Disease**

For the treatment of adult patients with moderately to severely active Crohn's disease who have contraindications, or are refractory, to therapy with corticosteroids and other immunosuppressants.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- Intravenous infusion: Approvals will be for maximum of 300 mg at week 0, 2, and 6, then 300 mg every eight weeks.
- Subcutaneous injection: Approvals will be for a maximum of 108 mg every two weeks following at least two intravenous infusions of vedolizumab.
- Initial approval period: 14 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required.

**Ulcerative Colitis**

- For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score greater than 4, and a rectal bleeding subscore greater than or equal to 2 and are:
  - refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks, and prednisone greater than or equal to 40mg daily for two weeks or IV equivalent for one week); or
  - corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year).
- Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease in the partial Mayo score greater than or equal to 2 from baseline, and
  - a decrease in the rectal bleeding subscore greater than or equal to 1.

Clinical Notes:

1. Consideration will be given for patients who have not received a four week trial of aminosalicylates if disease is severe (partial Mayo score greater than 6).
2. Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
3. Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic drug will not be reimbursed.
- Intravenous infusion: Approvals will be for maximum of 300 mg at week 0, 2, and 6, then 300 mg every eight weeks.
- Subcutaneous injection: Approvals will be for a maximum of 108 mg every two weeks following at least two intravenous infusions of vedolizumab.
- Initial approval period: 14 weeks.
- Renewal approval period: 1 year.

**VELAGLUCERASE ALFA (VPRIV)  
400 units per vial**

For the treatment of patients with symptomatic Gaucher disease type 1 (GD1).

Clinical Note:

- Requests for coverage must meet the criteria for diagnosis of GD1, indication for therapy and expected response to enzyme replacement therapy. These criteria are consistent with the Ontario Guidelines for the Treatment of Gaucher Disease. Please contact the NB Drug Plans at 1-800-332-3691 for the criteria.

Claim Notes:

- Approvals will be for a maximum of 60 units/kg every 2 weeks.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions as outlined [here](#).

**VEMURAFENIB (ZELBORAF)  
240 mg film-coated tablet**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used alone or in combination with cobimetinib.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Vemurafenib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy.
- Approval period: 6 months.

**VENETOCLAX (VENCLEXTA)  
10 mg, 50 mg, 100 mg film-coated tablets**

**Acute Myeloid Leukemia**

In combination with azacitidine for the treatment of patients with newly diagnosed acute myeloid leukemia who are 75 years of age or older, or who have comorbidities that preclude use of intensive induction chemotherapy.

Renewal Criteria:

- Written confirmation that the patient is responding to treatment and there is no evidence of disease progression.

Clinical Note:

- Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests for patients previously treated with a hypomethylating agent or chemotherapy for myelodysplastic syndrome will not be considered.
- Requests for patients with high-risk myelodysplastic syndrome will not be considered.
- Approval period: 1 year.

**Chronic Lymphocytic Leukemia / Small Cell Lymphoma**

1. In combination with obinutuzumab for adult patients with previously untreated chronic lymphocytic leukemia (CLL) / small lymphocytic lymphoma (SLL) whom fludarabine-based treatment is inappropriate.

Clinical Notes:

1. Patient must have a good performance status.
2. Treatment should be given for a total of 12 months (six 28-day cycles in combination with obinutuzumab, followed by six months of monotherapy), or until disease progression or unacceptable toxicity, whichever occurs first.

Claim Notes:

- Requests for re-treatment with venetoclax in combination with obinutuzumab will not be considered.
  - Approval period: 1 year.
2. In combination with rituximab for the treatment of patients with CLL / SLL who have received at least one prior therapy.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patient must have a good performance status.
2. Treatment should be continued until disease progression or unacceptable toxicity, up to a maximum of 2 years.

Claim Notes:

- Requests will not be considered for patients previously treated with anti-CD20 therapy if relapse occurs less than 6 months following completion of therapy. However, for patients previously treated with venetoclax, the relapse-free interval must be 12 months or greater.
  - Approval period: 1 year.
3. As monotherapy for the treatment of patients with CLL / SLL who have received at least one prior therapy which must include disease progression on or intolerance to a B-cell receptor inhibitor.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Requests will not be considered for patients previously treated with venetoclax-based therapy if relapse occurs less than 12 months following completion of therapy.
- Approval period: 1 year.

**VIGABATRIN (SABRIL)**

**500 mg tablet**

**500 mg sachet**

1. For the treatment of epilepsy in those patients who respond inadequately to alternative treatment combinations or in whom other drug combinations have not been tolerated.
2. For the treatment of infantile spasms.

Clinical Note:

- Potential benefits conferred by the use of vigabatrin should outweigh the risk of ophthalmologic abnormalities.

**VISMODEGIB (ERIVEDGE)**

**150 mg capsule**

**Initial Requests:**

- For patients with metastatic basal cell carcinoma (BCC) or with locally advanced BCC (including patients with basal cell nevus syndrome, i.e. Gorlin syndrome) who have measurable metastatic disease or locally advanced disease, which is considered inoperable or inappropriate for surgery<sup>1</sup> AND inappropriate for radiotherapy<sup>2</sup>  
AND
- Patient 18 years of age or older;  
AND
- Patient has ECOG  $\leq$  2
- Patient preference for oral therapy will not be considered

Information Required

- Physicians must provide rationale for why surgery<sup>1</sup> AND radiation<sup>2</sup> cannot be considered
- The request must include a surgical consultation report that provides a preoperative/surgical evaluation why surgery is not appropriate for the patient;  
AND
- A consultation report as to why radiation therapy is not appropriate for the patient
- Both of the above evaluations must come from a physician who is not the requesting physician
- Confirmation that the patient has been discussed at a multi-disciplinary cancer conference or equivalent (e.g. Regional Tumour Board).

Renewal Criteria:

- The physician has confirmed that the patient has not experienced disease progression while on Erivedge therapy.

Clinical Notes:

- <sup>1</sup>Considered inoperable or inappropriate for surgery for one of the following reasons:
  - Technically not possible to perform surgery due to size/location/invasiveness of BCC (either lesion too large or can be several small lesions making surgery not feasible)
  - Recurrence of BCC after two or more surgical procedures and curative resection unlikely
  - Substantial deformity and/or morbidity anticipated from surgery
- <sup>2</sup>Considered inappropriate for radiation for one of the following reasons:
  - Contraindication to radiation (e.g. Gorlin syndrome)
  - Prior radiation to lesion
  - Suboptimal outcomes expected due to size/location/invasiveness of BCC
- Dose: 150mg orally once daily taken until disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**VITAMINS B AND C (REPLAVITE)**

**Tablet**

For the replacement of water-soluble vitamins in patients with end-stage renal disease who are on dialysis.

Claim Note:

- Approval Period: Long term.

**VORICONAZOLE (VFEND and generic brands)**

**50 mg and 200 mg tablets**

- For the management of invasive aspergillosis.
- For culture proven invasive candidiasis with documented resistance to fluconazole.

Claim Notes:

- Must be prescribed by a hematologist, infectious disease specialist or medical microbiologist.
- Initial requests will be approved for a maximum of 3 months.

**ZANAMIVIR (RELENZA)**

**5 mg powder for inhalation**

For beneficiaries residing in long-term care facilities meeting the same criteria as for oseltamivir and for whom there is suspected or confirmed oseltamivir resistance, or for whom oseltamivir is contraindicated.

**ZANUBRUTINIB (BRUKINSA)**

**80 mg capsule**

For the treatment of adult patients with relapsed or refractory Waldenström macroglobulinemia who have received at least one prior therapy and have not experienced disease progression on a Bruton's tyrosine kinase inhibitor.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must meet at least one criterion for treatment as per IWWM consensus panel.
2. Patients must have a good performance status and no evidence of disease transformation.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Note:

- Approval period: 1 year.

**ZOLMITRIPTAN (ZOMIG NASAL SPRAY)**

**2.5 mg and 5 mg nasal sprays**

For the treatment of patients with acute migraine attacks who have an intolerance or insufficient response to oral triptans listed as regular benefits.

Claim Notes:

- Coverage limited to 6 doses per month.
- Requests for patients who have more than 3 migraines a month despite migraine prophylaxis therapy will be considered for a maximum of 12 doses per month.

## APPENDIX IV

### Provisional Benefits

The following drugs are provisional benefits according to the criteria specified below.

#### **ACETYLSALICYLIC ACID, CAFFEINE AND BUTALBITAL (FIORINAL and generics)**

**330 mg / 40 mg / 50 mg capsule and tablet**

#### **ACETYLSALICYLIC ACID, CAFFEINE, CODEINE AND BUTALBITAL (FIORINAL C1/4 and generic)**

**330 mg / 40 mg / 15 mg / 50 mg capsule**

#### **ACETYLSALICYLIC ACID, CAFFEINE, CODEINE AND BUTALBITAL (FIORINAL C1/2 and generic)**

**330 mg / 40 mg / 30 mg / 50 mg capsule**

Effective March 20, 2018, requests for coverage of butalbital-containing products which include Fiorinal, Fiorinal C<sup>1</sup>/<sub>4</sub>, Fiorinal C<sup>1</sup>/<sub>2</sub> and generic brands are no longer considered. Patients who had coverage prior to this date will continue to remain eligible for coverage if a special authorization request, documenting the rationale for continued use, is submitted on an annual basis.

#### **ADEFOVIR (HEPSERA and generic brand)**

**10 mg tablet**

Effective January 22, 2018 requests for coverage of adefovir (Hepsera) are no longer considered. Patients who had coverage of Hepsera prior to this date will continue to have coverage.

#### **ATORVASTATIN / AMLODIPINE (CADUET and generic brands)**

**10 mg / 5 mg, 10 mg / 10 mg, 20 mg / 5 mg, 10 mg / 10 mg, 40 mg / 5 mg, 40 mg / 10 mg, 80 mg / 5 mg, 80 mg / 10 mg tablet**

Effective May 13, 2021, atorvastatin/amlodipine tablets are no longer listed as a regular benefit. Patients who have had a claim paid for atorvastatin/amlodipine between November 13, 2020 and May 13, 2021 will continue to have coverage. Requests for special authorization will not be considered.

#### **BETAHISTINE (generic brands)**

**8 mg tablets**

Effective March 18, 2021, requests for coverage of betahistine 8 mg tablets are no longer considered. Patients who had a claim paid for betahistine 8 mg between September 18, 2020 and March 18, 2021 will continue to have coverage.

#### **CHLORAL HYDRATE (Chloral hydrate syrup Odan)**

**100 mg/mL syrup**

Effective June 26, 2023, chloral hydrate syrup is no longer listed as a regular benefit. For patients who had a claim paid for chloral hydrate between December 26, 2022 and June 26, 2023, chloral hydrate syrup will continue to be a benefit until January 26, 2024. After January 26, 2024, a special authorization request, documenting the rationale for continued use, will be required on an annual basis for coverage to be considered. New requests for special authorization will not be considered.

#### **CLIDINIUM / CHLORDIAZEPOXIDE (LIBRAX)**

**5 mg / 2.5 mg capsules**

Effective April 22, 2021, chlordiazepoxide/clidinium is no longer listed as a regular benefit. For patients who had a claim paid for chlordiazepoxide/clidinium between October 22, 2020 and April 22, 2021, chlordiazepoxide/clidinium will continue to be a benefit until October 22, 2021. After October 22, 2021, a special authorization request, documenting the rationale for continued use, will be required on an annual basis for coverage to be considered. New requests for special authorization will not be considered.

#### **GLIMEPIRIDE (Sandoz Glimepiride)**

**1 mg, 2 mg, 4 mg tablets**

Effective June 17, 2021, Sandoz glimepiride 1 mg, 2 mg and 4 mg tablets are no longer listed as a regular benefit. Patients who had a claim paid between June 17, 2020 and June 17, 2021, will continue to have coverage. Requests for special authorization will not be considered.

**HYDROCORTISONE / PRAMOXINE / ZINC (PROCTODAN-HC)****0.5% / 1% / 0.5% ointment****10 mg / 20 mg / 10 mg suppositories**

Effective June 29, 2023, Proctodan-HC ointment and suppositories are no longer listed as a regular benefit. Patients who had a claim paid between December 29, 2022 and June 28, 2023 will continue to have coverage. Requests for special authorization will not be considered.

**OXYBUTYNIN (generic brand)****2.5 mg tablet**

Effective April 22, 2021, pms-Oxybutynin 2.5 mg tablets are no longer listed as a regular benefit. Patients who had a claim paid between October 22, 2020 and April 22, 2021 will continue to have coverage. Requests for special authorization will not be considered.

**PLACEBO****100 mg capsule**

Effective June 29, 2023, placebo 100 mg capsules are no longer listed as a regular benefit. Patients who had a claim paid between December 29, 2022 and June 28, 2023 will continue to have coverage. Requests for special authorization will not be considered.

**QUININE SULFATE (generic brands)****200 mg and 300 mg capsules****300 mg tablet**

Effective September 1, 2017, quinine is no longer listed as a regular benefit. For patients who have had a claim paid for quinine between September 1, 2016 and August 31, 2017, quinine will continue to be a benefit until March 1, 2018. After March 1, 2018, a special authorization request, documenting the rationale for continued use, will be required for coverage to be considered. Requests for special authorization will not be considered for new patients or patients who have not had a claim paid for quinine between September 1, 2016 and August 31, 2017.

**RANIBIZUMAB (LUCENTIS)****10 mg/mL solution for intravitreal injection**

Effective August 28, 2023, ranibizumab (Lucentis) is no longer listed as a special authorization (SA) benefit. Patients who had coverage of Lucentis prior to this date will continue to have coverage until their current SA approval expires, or February 28, 2024, whichever occurs first. Patients must switch to the biosimilar brand of ranibizumab to maintain their coverage under the New Brunswick Drug Plans.

For patients who are unable to switch for medical reasons, an SA request for exceptional coverage of the originator biologic may be submitted. Exceptional requests are reviewed on a case-by-case basis.

The biosimilar brand of ranibizumab is listed as a SA benefit.

**ROSIGLITAZONE (AVANDIA and generic brand)****2 mg, 4 mg, 8 mg tablets**

Effective April 2, 2012, requests for coverage of rosiglitazone (Avandia) are no longer considered. Patients who had coverage of Avandia prior to this date will continue to have coverage.