



NON-INSURED HEALTH BENEFITS

First Nations and Inuit Health Branch

DRUG BENEFIT LIST

June 2019

The Non-Insured Health Benefits (NIHB) program provides supplementary health benefits, including prescription and non-prescription drugs, for registered First Nations and recognized Inuit throughout Canada.

Visit our Web site at: www.canada.gc.ca/nihb

**Department of Indigenous Services Canada
Non-Insured Health Benefits**

**INTRODUCTION
Drug Benefit List**

**Effective
June 2019**

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1. BACKGROUND ON NON-INSURED HEALTH BENEFITS (NIHB) PROGRAM

The Non-Insured Health Benefits (NIHB) Program of the Department of Indigenous Services Canada provides clients (registered First Nations and recognized Inuit) with coverage for a range of health benefits, including prescription drugs and over-the-counter medications, dental and vision care, medical supplies and equipment, mental health counselling, and transportation to access health services not available locally. These benefits complement provincial and territorial health care programs, such as physician and hospital care, as well as other First Nations and Inuit community-based programs and services. Benefits include drugs, medical transportation, dental care, medical supplies and equipment, crisis intervention counselling and vision care.

The authority for the NIHB Program is based on the 1979 Indian Health Policy which describes the responsibility for the health of First Nations as shared amongst various levels of government, the private sector and First Nations communities. As a result of this shared responsibility, when a benefit is covered under another plan, the federal government requires the coordination of benefits to ensure that the other plan meets its obligations.

2. PURPOSE OF THE NIHB DRUG BENEFIT LIST (DBL)

The Drug Benefit List (DBL) is a listing of the drugs provided as benefits by the NIHB Program. The DBL is updated regularly and published regularly. The listed drugs are those primarily used in a home or ambulatory setting. A prescription from a licensed practitioner is required for any listed drug to be processed as a benefit. Practitioners are health professionals authorized to prescribe drugs within the scope of practice in their province or territory. The DBL is a tool for prescribers and pharmacists that encourages the selection of optimal, cost-effective drug therapy.

3. DRUG REVIEW PROCESS

The review process for drug products that are considered for inclusion as a benefit under the NIHB Program varies depending on the type of drug submitted.

3.1 New Chemical Entities / New Combination Drug Products/ Existing Chemical Entities with New Indication

Submissions for new chemical entities, new combination drug products and existing chemical entities with new indications, must be sent to the Canadian Agency for Drugs and Technologies in Health (CADTH). Clinical and pharmacoeconomic reviews are coordinated by the Common Drug Review (CDR) Directorate, or by the pan-Canadian Oncology Drug Review (pCODR) for cancer therapies, and forwarded to their respective expert committees for recommendations on formulary listing. These recommendations are forwarded to participating drug plans, including the NIHB Program, for consideration. The NIHB Program and other drug plans make listing decisions based on these expert committee recommendations and other specific relevant factors, such as mandate, priorities and resources.

Please refer to CADTH for a list of requirements for manufacturers' submissions and a summary of procedures for the CDR or pCODR process. Inquiries should be directed to:

Canadian Agency for Drugs and Technologies in Health
865 Carling Avenue, Suite 600
Ottawa, Ontario K1S 5S8
Telephone: (613) 226-2553
Website: <http://www.cadth.ca>

Please ensure a copy of the complete submission is also sent to NIHB either electronically to NIHB.Drug.Submissions@hc-sc.gc.ca or on compact CD to the mailing address indicated in section 3.2.2.4. Paper (binder) versions of drug submissions are no longer accepted by the NIHB Program.

3.2 Line Extensions, Generics and All Other Submissions

Submissions for line extensions, generics and all other submissions are reviewed internally or by

the NIHB Drugs and Therapeutics Advisory Committee (DTAC). Generic drug products are considered for inclusion on the formulary based on provincial interchangeability lists and other relevant factors.

3.2.1 Drugs and Therapeutics Advisory Committee (DTAC)

The [DTAC](#) provides formulary listing recommendations for drug products to the NIHB Program. The NIHB Program makes listing decisions based on DTAC recommendations and other specific relevant factors, such as mandate, priorities and resources. The DTAC also contributes to the NIHB Drug Use Evaluation (DUE) Program which promotes safe, therapeutically effective and efficient use of drug therapy for First Nations and Inuit.

The [DTAC](#) is an advisory body of highly qualified health professionals who bring impartial and practical expert medical and pharmaceutical advice to the NIHB Program to promote improvement in the health outcomes of First Nations and Inuit clients through effective use of pharmaceuticals. The approach is evidence-based and the advice reflects medical and scientific knowledge, current utilization trends, current clinical practice, health care delivery and specific departmental client healthcare needs.

3.2.2 Submission Requirements

All submissions for drug products that are line extensions, generics and all other types of submissions must be submitted to the NIHB Program. Only drug products with a Health Canada Notice of Compliance (NOC) will be considered for provision as a benefit.

3.2.2.1 Letter of Authorization

The manufacturer will provide a letter authorizing the NIHB Program to gain access to all information with respect to the product in the possession of Health Canada or of the government of any provinces or territory in Canada, Patented Medicine Prices Review Board (PMPRB) or CADTH.

3.2.2.2 Justification for Consideration of Listing

The manufacturer will provide a statement indicating the rationale and evidence to justify the provision of the new product.

3.2.2.3 General Information

Additional information should include:

- Evidence of approval by Health Canada, such as a Notice of Compliance (NOC) and Drug Identification Number (DIN) and
- Two therapeutic Classifications:
 - American Hospital Formulary Service (AHFS) Pharmacologic Therapeutic Classification and;
 - The World Health Organization's Anatomical Therapeutic Chemical (ATC) Classification

3.2.2.4 Pricing and Marketing Information

The manufacturer must submit current price information for the drug product.

Manufacturers are required to notify the NIHB Program of any significant change to listed drug products. Significant changes include changes in DIN, product name, manufacturer or distributor, indication, product monograph, packaging, formulation, manufacturing specifications or discontinuation of a product. Notification of changes should be provided electronically to the NIHB Program.

All submissions for drug products, to be reviewed for inclusion on the NIHB DBL, must be sent to the NIHB Program electronically. Please send all drug submissions to the following email address: NIHB.Drug.Submissions@hc-sc.gc.ca. Submissions will also be accepted on compact CD when mailed to the

following address:

C/o Manager of Policy Development - Pharmacy
Non-Insured Health Benefits
First Nations and Inuit Health Branch, Department of Indigenous Services
Canada
200 Eglantine Driveway, 9th Floor
Postal Locator 1909D Tunney's Pasture
Ottawa, Ontario K1A 0K9

Only ONE copy of the submission is required. Receipt of submission will be acknowledged electronically with a confirmatory email message. Paper (binder) versions of drug submissions are no longer accepted by the NIHB Program.

4. BENEFIT CRITERIA

The following criteria are the framework for the NIHB Program DBL. The criteria provide the basis for decisions about drugs on the formulary relating to:

- A. Drug Benefit Listings
- B. Deletions
- C. Open Benefit
- D. Limited Use
- E. Exceptions
- F. Exclusions

All drugs that are to be either considered for listing or currently listed as Program benefits must, as a minimum:

1. be legally available for sale in Canada with an NOC;
2. sold in Canada (proof may include a copy of the completed notification form issued under the Food and Drug Regulations or listing on a provincial drug benefit formulary);
3. be administered in a home setting or in other ambulatory care settings;
4. not be provided in a provincially/territorially covered setting (hospital/institution) or provided through provincially/territorial covered programs or clinics according to provincial/territorial legislation; and
5. be in accordance with NIHB Program mandate and policies.

A. Drug Benefit Listings

The NIHB Program, with assistance from the CDR, pCPA, pCODR and the NIHB DTAC, balances a number of factors in making listing decisions about changes to the Drug Benefit List, such as:

- The needs of First Nations and Inuit clients;
- Accumulated scientific and clinical research on currently-listed drugs;
- Cost-benefit analysis;
- Availability of alternatives;
- Current health practices; and
- Policies and listings in provincial drug formularies.

New formulations and new strengths of listed products may be added or may replace previously approved products.

Generic products are added according to provincial/territorial interchangeability lists and other relevant factors.

Combination products are considered for listing if:

1. each component of the combination makes a contribution to the claimed effect;
2. a pharmacological or pharmaceutical rationale exists for the combination;
3. the dosage of each component (amount, frequency, duration) is safe and effective for a significant proportion of the patient population requiring such concurrent therapy as defined in the labeling of the drug; and
4. the cost is reduced, or scientific evidence indicates that the advantages outweigh any additional cost; or
5. an improvement in compliance, resulting in an increase in clinical effectiveness, is demonstrated.

Long Acting (Sustained-Extended Release) Products may be listed when:

1. clinical studies have demonstrated the safety and efficacy of the active ingredient when administered in the long acting form; and
2. a therapeutic advantage is demonstrated in the treatment of the disease entity for which the product is indicated (therapeutic advantage is defined as: improved efficacy relative to the conventional dosage with no increase in toxicity; or less toxicity with improved or similar efficacy); or
3. there is demonstrated improvement in compliance resulting in an increase in clinical effectiveness; or
4. there is evidence that the long acting product is at least as cost-effective as the best price alternative in the conventional form that is currently covered; or
5. there is no suitable conventional dosage form(s) of the drug listed that is readily available.

Injectable Drug Products will be considered if they are:

1. self-administered in a home or other ambulatory setting;
2. not part of a physician's standard office supply;
3. not provided in a provincially/territorially covered hospital or institution; or
4. not provided through provincially/territorial covered programs or clinics according to provincial/territorial legislation.

B. Deletion Criteria

The following deletion criteria guide the removal or delisting of a drug product from the NIHB drug benefit list. Drugs are deleted:

1. when a product is discontinued from the Canadian market;
2. when new products possessing clearly demonstrated therapeutic and safety advantages or improvements have been listed;
3. when new toxicity data shift the risk/benefit ratio to make the continued listing of the product inappropriate;
4. when new information demonstrates that the product does not have the anticipated therapeutic benefit;
5. when the purchase cost is disproportionate to the benefits provided; or
6. when the drug has a high potential for misuse or abuse.

NOTE: Drugs may also be removed at the discretion of the Director General, NIHB Program when there are undesirable financial, supply or administrative implications to the continued listing of a product.

C. Open Benefits

Open benefits are the drugs listed in the NIHB DBL which do not have established criteria or prior approval requirements.

D. Limited Use Benefits

Limited use drugs are drug products listed on the NIHB DBL that may be inappropriate for

general listing, but have value in specific circumstances. These products will have specific criteria for provision as a benefit under the NIHB Program. A product will be designated for limited use when:

1. it has the potential for widespread use outside the indications for which benefit has been demonstrated;
2. it has proven effectiveness, but is associated with predictable severe adverse effects;
3. it is usually a second or third line choice for treatment and is required because of allergies, intolerance, treatment failure or noncompliance with a first line alternative; or
4. it is very costly and a therapeutically effective alternative is available as a benefit.

There are three types of limited use benefits:

1. Limited use benefits which do not require prior approval. These include but are not limited to:
 - Multivitamins (which are benefits for children up to 19 years of age); and
 - Prenatal and postnatal vitamins (which are benefits for women of childbearing age (12 to 50 years)).

2. Benefits which have a quantity and/or frequency limit. A maximum quantity of drug is allowed within a specified period of time. No prior approval is required for the recipient to obtain the allowable quantity of drug within the specified period. An example of a category of drugs with a quantity and frequency limit is smoking cessation products. Recipients are eligible to receive up to three treatment courses of nicotine replacement therapy (NRT) within a 12-month period with quantity limits, which include two courses of NRT patches and one course of NRT products used PRN (i.e. gums, lozenges, inhalers).

3. Limited use benefits which require prior approval (using the "Limited Use Drugs Request Form"). Limited use benefits and the criteria for their coverage are identified in the Drug Benefit List and also in Appendix A. The criteria are also listed on the forms faxed to prescribers for completion.

E. Exceptions

Exception drugs are drug products which are not listed in the DBL. These drug products may be approved in special circumstances upon receipt of a completed "Exception Drugs Request Form" from the attending licensed practitioner.

- when the prescription is for a recognized clinical indication and dose which is supported by published evidence or authoritative opinion; and
- when there is significant evidence that the requested drug is superior to drugs already listed as program benefits; or
- when a patient has experienced an adverse reaction with a best-price alternative drug, and a higher cost alternative is requested by the prescriber; or
- when there is supporting evidence that available alternatives are ineffective, toxic, or contraindicated (personal preference alone does not justify an exception).

F. Exclusions

Exclusions are items not listed as benefits on the DBL and are not available through the exception or appeal processes. These include certain drug therapies for particular conditions which fall outside of the NIHB mandate and are not provided as benefits under the NIHB Program.

Examples of categories of drugs or drug products* that are not considered for coverage under the NIHB Program under any circumstances are listed in Appendix G

- Anti-obesity drugs;
- Household products (e.g. regular soaps and shampoos);
- Cosmetics;

- Alternative therapies, including glucosamine and evening primrose oil;
- Megavitamins;
- Drugs with investigational/experimental status;
- Vaccines
- Medications for travel
- Hair growth stimulants;
- Fertility agents and impotence drugs;
- Selected over-the-counter products;
- Opioid containing cough preparations.

*Note: List of excluded drugs or drug products is not exhaustive and may be modified as necessary

5. POLICIES

A. Best Price Alternative and Interchangeability

The NIHB Program will reimburse only the best price (lowest cost) alternative product in a group of interchangeable drug products. Pharmacists must follow their provincial/territorial pharmacy legislation/policies to identify interchangeable products and to select the lowest-priced brand. (NIHB may not necessarily reimburse at the cost listed in the provincial drug plan formulary).

B. “No Substitution” Claims

NIHB will consider reimbursement for a higher-cost interchangeable product when a patient has experienced an adverse reaction with a lower-cost alternative. In such circumstances, the prescriber must provide the NIHB Program with:

1. a completed and signed Canada Vigilance Adverse Reaction Reporting Form: ‘Report of suspected adverse reactions to health products in Canada’ and,
2. the prescription with “No Substitution” or “No Sub” written by hand or typed on the prescription.

Upon receipt, the pharmacist will forward a copy of the prescription to NIHB for review. The prescriber is responsible for sending a copy of the form to the Canada Vigilance Program. Forms can be obtained by calling the Canada Vigilance Program at 1-866-234-2345 or by downloading a copy from Health Canada website at: http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/ar-ei_form-eng.php

NOTE: The Canada Vigilance Adverse Reaction Reporting Form will not need to be resubmitted for renewals or new prescriptions of the same drug for the patient, although “No Sub” will still have to be written or typed on the prescription.

C. Prescription Quantities

The normal quantity dispensed shall be the entire quantity of the drug prescribed. A maximum 100-day supply should be considered for those circumstances where the patient has been stabilized on a medication and the prescriber feels that further adjustment during the prescribed period is unlikely. Prescriptions for opioids and benzodiazepines have a maximum 30-day supply. The physician may continue to prescribe a smaller quantity with repeats at certain intervals when it is in the patient’s best interest.

D. Short Term Dispensing Policy

It is the Program’s expectation that certain medications required for long-term maintenance therapy should be prescribed and dispensed in up to 100 days supplies. For refills for medications requiring short-term dispensing for a shorter time than 28 days due to compliance concerns, the Program will only reimburse a total of one dispensing fee per 28 days up to the regional maximum of the Program, These medications include (but are not limited to) the following:

Antihistamines	Anticoagulants	Immunosuppressants
Antiemetics for cancer chemotherapy (excluding nabilone)		Prokinetic agents
Synthetic antidiuretic hormone	Respiratory smooth muscle relaxants	
Alpha-adrenoreceptor Antagonists	Anti-dementia Drugs	Anti-gout Drugs
Anti-Parkinsonian Drugs	Anti-platelet aggregation Drugs	BPH Drugs
Cardiovascular Drugs	Enzyme Preparations	Drugs for Diabetes
Drugs for Treatment of Bone Diseases	GI Anti-inflammatory Drugs	Thyroid Therapy
Proton Pump Inhibitors	Urinary Anti-Spasmotics	NSAIDs
H2-Receptor Antagonists	OTCs (including vitamins)	
Other Drugs for Peptic Ulcer and Gastro-esophageal Reflux Disease (GERD)		

Note: This list may be amended as required and changes will be communicated through the quarterly on-line updates to the DBL. Medications on the Short term Dispensing list are identified in the DBL using the symbol ST beside the medication strength and dosage form.

The following are exceptions to the STD policy:

- Refills for intermittent treatment of a chronic disorder or refills of a medication which is prescribed to be taken on an “as needed” (PRN) basis. Note: Medications prescribed to be taken on an “as needed” (PRN) basis and dispensed chronically may be subject to audit and recovery.
- Prescriptions for dose changes.
- The following dosage forms: injectable and suppository.
- Refills or new prescriptions when prescribed/dispensed in accordance with a court order.
- Others as identified by the NIHB Program

Compensation

The compensation will be the lesser of the usual and customary fee up to the maximum negotiated NIHB regional dispensing fee for each 28 days supplied. NIHB will continue to audit and recover in instances where quantity reduction occurs.

Less than 28 Day Supply

For the medications listed below in which short-term dispensing is deemed medically necessary, the Program will compensate up to one full dispensing fee every seven days, up to the regional maximum of the Program. If these medications are dispensed daily, the Program will compensate 1/7th of this fee:

Anticonvulsants	Hormonal Contraceptives
Antidepressants	Needles & Syringes
Antipsychotics	Drug used in nicotine dependence
Benzodiazepines	Antimanic agents
Stimulants	Estrogens
Nicotine Replacement Therapy	Progestins

Implementation

When filling a new prescription for a chronic use drug, the Program will pay a full dispensing fee regardless of the days supply. A new prescription may include a dosage change or an intermittent treatment, based on an assessment by a prescriber.

When refilling a prescription for a chronic use drug that is for less than a 28 day supply or when a need for compliance packaging is identified by the prescriber, the Program will pay no more than one full dispensing fee per 28 day period. For the medications listed above the Program will pay no more than full dispensing fee per 7 day period.

A refill is defined as the second and all subsequent fills for a given strength and dosage of a drug.

6. FORMULARY FOR CHRONIC RENAL FAILURE PATIENTS

Clients with chronic renal failure are eligible to receive a list of supplemental benefits that are not included in the NIHB DBL but which are required on a long-term basis. Some supplemental benefits include: darbepoetin alfa products (except in provinces where NIHB clients are eligible to receive darbepoetin alfa through the provincial programs), calcium products, multivitamins formulated for renal patients and select nutritional supplements to support management of chronic renal failure.

New clients requiring drugs on the special formulary will be identified for coverage through the usual prior approval process. Once the client is confirmed as eligible, coverage will automatically be extended to all drugs in the special formulary for as long as needed.

7. PALLIATIVE CARE FORMULARY

Clients diagnosed with a terminal illness and are near the end of life will be eligible to receive a list of supplemental benefits that are not included in the NIHB Drug Benefit List. The Palliative Care Formulary includes medications and nutritional supplements used to provide comfort to those near the end of life.

Requests for any of the DINs on the Palliative Care Formulary will generate a Palliative Care Application Form, faxed to the prescriber. Once completed and submitted, the recipient will be eligible for all medications on the Palliative Care Formulary for six months if the following criteria are met:

The client:

1. is not receiving care in a provincially covered hospital or provincially covered long-term care facility; and
2. has been diagnosed with a terminal illness or disease which is expected to be the primary cause of death within six months or less

If coverage is required beyond the initial six months, an additional six months will be granted upon receipt of another completed Palliative Care Application Form.

8. ADJUNCT CANCER FORMULARY

The NIHB Program has established a new formulary to streamline access to adjunctive (non-chemotherapy) medications frequently used by clients undergoing active cancer treatment.

Clients who are approved for oral chemotherapy drugs are given access to all of the medications and nutritional supplements in the formulary. Additionally, clients who request, and are approved, for one of the medications on the formulary for a cancer-related indication are also granted access.

Clients are automatically enrolled for a period of six months. If cancer treatment is of a longer duration, access to the formulary will be granted to align with the treatment duration. In the event that treatment duration is not known and the treatment plan extends beyond six months, access to this formulary may be extended upon request.

9. DRUG UTILIZATION EVALUATION

A drug utilization evaluation, which is part of the point-of-service or on-line adjudication system, provides an analysis of both previous claims data and current claims data to identify potential drug-related problems. Messages are returned to pharmacists to alert them of the potential problems. These messages are intended to enhance pharmacy practice with additional information. Currently, the system monitors for:

- potential drug/drug interactions
- duplicate drugs
- duplicate therapy

As part of the NIHB Drug Use Evaluation (DUE) Program, DTAC reviews utilization patterns of medications billed to the NIHB program and provides advice to promote effective, efficient and optimal drug therapy to First Nations and Inuit recipients.

10. GENERAL INFORMATION

Sources of information about the NIHB Program include:

- The NIHB section of the Government of Canada website which provides background information on the Program and a copy of the DBL. This can be found at: <http://www.canada.ca/nihb>

Information about the NIHB Program can also be obtained by contacting:

Non-Insured Health Benefits
First Nations and Inuit Health Branch
200 Eglantine Driveway, 9th Floor
Postal Locator 1909D
Tunney's Pasture
Ottawa, Ontario K1A 0K9

11. NIHB PRIVACY CODE

The NIHB Program is committed to protecting an individual's privacy and safeguarding the personal information in its possession. When a benefit request is received, the NIHB Program collects, uses, discloses and retains an individual's personal information according to the applicable federal privacy legislation. The information collected is limited to only that information required for the NIHB Program to administer and verify benefits.

As a program of the federal government, the NIHB Program must comply with the Privacy Act, the Canadian Charter of Rights and Freedoms, the Access to Information Act, the Treasury Board of Canada Privacy and Data Protection Policies, the Government Security Policy, and Health Canada's Security Policy.

12. PHARMACOLOGIC-THERAPEUTIC CLASSIFICATION OF DRUGS

The drugs in the NIHB DBL are classified according to the AHFS Pharmacologic-Therapeutic classification developed by the American Society of Health-System Pharmacists for the purposes of the AHFS Drug Information.

Permission to use this system has been granted by the American Society of Health-System Pharmacists. The Society is not responsible for the accuracy of transpositions from the original context.

Drugs are listed alphabetically within each therapeutic classification according to their chemical names. Under each drug, acceptable products are listed.

LEGEND

1. Pharmacologic-Therapeutic classification
2. Pharmacologic-Therapeutic sub-classification
3. Nonproprietary or generic name of the drug
4. Drug strength and dosage form. ST indicates the drug is identified as a chronic medication under the Short-Term Dispensing Policy.
5. Drug Identification Number (DIN), assigned by the Therapeutic Products Directorate of Health Canada, to uniquely identify the drug product as to its manufacturer, name and strength of active ingredients, route of administration and pharmaceutical dosage form
6. Brand name of the drug
7. List of all active ingredients in a combination product
8. Strengths of active ingredients in a combination product, listed in the same order as the ingredients
9. List of available brands of drugs. Provincial or territorial drug plan formularies should be consulted to determine interchangeable products and to identify best price (lowest cost) alternatives
10. Three letter identification code assigned to manufacturer

1	→	04:00 ANTIHISTAMINE DRUGS		
2	→	04.00.00 ANTIHISTAMINE DRUGS		
3	→	CETIRIZINE HCL		
4	→	ST 10mg Tablet		
5	→	02231603	APO-CETIRIZINE	APX
6	→	↑		
7	→	28:08.08 ACETAMINOPHEN, CAFFEINE, CODEINE PHOSPHATE		
8	→	300mg & 15mg & 15mg Tablet		
9	→	00706515 00653241 02163934	PMS-ACET 2 RATIO-LENOLTEC NO.2 TYLENOL WITH CODEINE NO.2	PMS RPH JNO
		300mg & 15mg & 30mg Tablet		
		00653276 02163926	RATIO-LENOLTEC NO.3 TYLENOL WITH CODEINE NO.3	RPH JNO
10	→	↑		

DRUG BENEFIT LIST

04:00 ANTIHISTAMINE DRUGS

04:04.04 ANTIHISTAMINE DRUGS

DIPHENHYDRAMINE HYDROCHLORIDE

ST 25MG CAPSULE			
00757683	PDP-DIPHENHYDRAMINE		PMS
ST 50MG CAPSULE			
00757691	PDP-DIPHENHYDRAMINE		PMS
ST 2.5MG/ML ELIXIR			
00833266	ALLERGY ELIXIR		TAN
00804193	ALLERNIX ELIXIR		TEV
02019736	BENADRYL		MCL
00792705	PMS-DIPHENHYDRAMINE		PMS
ST 12.5MG/5ML ELIXIR			
02298503	DIPHENHYDRAMINE		JMP
ST 1.25MG/ML LIQUID			
02019698	BENADRYL CHILDRENS		MCL
50MG/ML LIQUID			
00596612	DIPHENHYDRAMINE		SDZ
02219336	DIPHENIST		OMG
00878200	PMS-DIPHENHYDRAMINE		PMS
ST 25MG TABLET			
02176483	ALLER-AIDE		TEV
01949454	ALLERGY		TAN
02229492	ALLERGY FORMULA		VTH
02097583	ALLERNIX		TEV
02017849	BENADRYL		MCL
02257548	DIPHENHYDRAMINE		JMP
02239029	NADRYL		RIV
ST 50MG TABLET			
02230398	ALLERGY EXTRA STRENGTH		TAN
02097575	ALLERNIX EXTRA STRENGTH		TEV
02257556	DIPHENHYDRAMINE		JMP

04:04.20 ANTIHISTAMINE DRUGS

CHLORPHENIRAMINE MALEATE

ST 4MG TABLET			
00738972	CHLOR-TRIPOLON		BAY
00021288	NOVO-PHENIRAM		TEV
ST 12MG TABLET (EXTENDED RELEASE)			
00738964	CHLOR-TRIPOLON		BAY

04:08.00 ANTIHISTAMINE DRUGS

CETIRIZINE HYDROCHLORIDE

ST 1MG/ML SYRUP			
02238337	REACTINE		MCL
ST 10MG TABLET			
02315955	ALLERGY RELIEF		PMS
02231603	APO-CETIRIZINE		APX
02375095	CETIRIZINE		APX
02451778	JAMP-CETIRIZINE		JMP
02427133	MAR-CETIRIZINE		MAR
02223554	REACTINE		MCL
ST 20MG TABLET			
02453363	APO-CETIRIZINE		APX
02450526	CETIRIZINE		PDL
02427141	MAR-CETIRIZINE		MAR

04:08.00 ANTIHISTAMINE DRUGS

CETIRIZINE HYDROCHLORIDE

ST 20MG TABLET			
02315963	PMS-CETIRIZINE		PMS
02427192	PRIVA-CETIRIZINE		PHA
01900978	REACTINE		MCL

DESLOMATADINE

ST 0.5MG/ML SYRUP			
02247193	AERIUS KIDS		BAY
ST 5MG TABLET			
02243919	AERIUS		BAY
02369656	ALLERNIX MULTI SYMPTOM		TEV
02338424	DESLOMATADINE		APX
02298155	DESLOMATADINE ALLERGY CONTROL		PMS

FEXOFENADINE HYDROCHLORIDE

ST 60MG TABLET			
02231462	ALLEGRA 12 HOUR		SAC
ST 120MG TABLET			
02242819	ALLEGRA 24 HOUR		SAC

LORATADINE

ST 1MG/ML SYRUP			
02241523	CLARITIN KIDS		BAY
ST 10MG TABLET			
02280159	24 HOUR ALLERGY REMEDY		VTH
02375990	ALLERGY REMEDY		APX
02418959	ALLERTIN		APX
02243880	APO-LORATADINE		APX
00782696	CLARITIN		BAY
02366444	LORATADINE		APX

04:92.00 ANTIHISTAMINE DRUGS

KETOTIFEN FUMARATE

ST 0.2MG/ML SYRUP			
00600784	ZADITEN		TEV
ST 1MG TABLET			
00577308	ZADITEN		TEV

08:00 ANTI-INFECTIVE AGENTS

08:08.00 ANTHELMINTICS

MEBENDAZOLE

100MG TABLET

00556734 VERMOX JSO

PYRANTEL PAMOATE

50MG SUSPENSION

02412470 JAMP-PYRANTEL PAMOATE JMP

50MG/ML SUSPENSION

01944355 COMBANTRIN MCL

125MG TABLET

01944363 COMBANTRIN MCL

08:12.02 AMINOGLYCOSIDES

AMIKACIN SULFATE

Limited use benefit (prior approval required).

250MG LIQUID

02242971 AMIKACIN SULFATE SDZ

GENTAMICIN SULFATE

1MG/ML SOLUTION

02082136 GENTAMICIN IV BAX

1.6MG/ML SOLUTION

02082152 GENTAMICIN IV BAX

10MG/ML SOLUTION

02268531 GENTAMICIN SDZ

40MG/ML SOLUTION

02225131 CIDOMYCIN UNK

02242652 GENTAMICIN SDZ

PDIN FOR EXTEMPORANEOUS MIXTURE

99506004 GENTAMYCIN STERILE INFUSION UNK

TOBRAMYCIN

Limited use benefit (prior approval required).

28MG CAPSULE

02365154 TOBI PODHALER NVR

1.2G POWDER FOR SOLUTION

00533688 TOBRAMYCIN FKD

02285150 TOBRAMYCIN RAX

10MG/ML SOLUTION

02230639 TOBRAMYCIN FKD

02241209 TOBRAMYCIN SDZ

40MG/ML SOLUTION

02420287 JAMP-TOBRAMYCIN JMP

02230640 TOBRAMYCIN FKD

02241210 TOBRAMYCIN SDZ

02382814 TOBRAMYCIN MYL

99005069 TOBRAMYCINE UNK

60MG SOLUTION

02389622 TEVA-TOBRAMYCIN TEV

300MG SOLUTION

02443368 TOBRAMYCIN INHALATION SDZ

08:12.06 CEPHALOSPORINS

CEFACLOR

250MG CAPSULE

02230263 APO-CEFACLOR APX

08:12.06 CEPHALOSPORINS

CEFACLOR

500MG CAPSULE

02230264 APO-CEFACLOR APX

CEFADROXIL

500MG CAPSULE

02240774 APO-CEFADROXIL APX

02311062 PRO-CEFADROXIL PDL

02235134 TEVA-CEFADROXIL TEV

CEFAZOLIN SODIUM

500MG POWDER FOR SOLUTION

02108119 CEFAZOLIN TEV

02237137 CEFAZOLIN FKD

02308932 CEFAZOLIN SDZ

1G POWDER FOR SOLUTION

02108127 CEFAZOLIN TEV

02237138 CEFAZOLIN FKD

02297205 CEFAZOLIN PFI

02308959 CEFAZOLIN SDZ

02437112 CEFAZOLIN RAX

10G POWDER FOR SOLUTION

02108135 CEFAZOLIN TEV

02237140 CEFAZOLIN FKD

02297213 CEFAZOLIN PFI

02308967 CEFAZOLIN SDZ

02437120 CEFAZOLIN RAX

PDIN FOR EXTEMPORANEOUS MIXTURE

99506000 CEFAZOLIN STERILE INFUSION UNK

CEFIXIME

20MG/ML POWDER FOR SUSPENSION

00868965 SUPRAX ODN

100MG POWDER FOR SUSPENSION

02468689 AURO-CEFIXIME AUR

400MG TABLET

02432773 AURO-CEFIXIME AUR

00868981 SUPRAX ODN

CEFPROZIL

25MG/ML POWDER FOR SUSPENSION

02293943 APO-CEFPROZIL APX

02329204 RAN-CEFPROZIL RBY

50MG/ML POWDER FOR SUSPENSION

02293951 APO-CEFPROZIL APX

02293579 RAN-CEFPROZIL RBY

250MG TABLET

02292998 APO-CEFPROZIL APX

02347245 AURO-CEFPROZIL AUR

02293528 RAN-CEFPROZIL RBY

02302179 SANDOZ CEFPROZIL SDZ

500MG TABLET

02293005 APO-CEFPROZIL APX

02347253 AURO-CEFPROZIL AUR

02293536 RAN-CEFPROZIL RBY

02302187 SANDOZ CEFPROZIL SDZ

08:12.06 CEPHALOSPORINS

CEFTAZIDIME

Limited use benefit (prior approval required).

1G POWDER FOR SOLUTION

00886971	CEFTAZIDIME	FKD
02437848	CEFTAZIDIME	RAX
02212218	FORTAZ 1G	GSK

2G POWDER FOR SOLUTION

00886955	CEFTAZIDIME	FKD
02437856	CEFTAZIDIME	RAX
02212226	FORTAZ 2G	GSK

3G POWDER FOR SOLUTION

02439522	CEFTAZIDIME	RAX
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6G POWDER FOR SOLUTION

00886963	CEFTAZIDIME	FKD
02437864	CEFTAZIDIME	RAX
02212234	FORTAZ 6G	GSK

CEFTRIAXONE SODIUM

250MG POWDER FOR SOLUTION

02250276	CEFTRIAXONE	PFI
02289679	CEFTRIAXONE	FKD
02292262	CEFTRIAXONE	SDZ
02292866	CEFTRIAXONE	PFI
02325594	CEFTRIAXONE	RAX

1G POWDER FOR SOLUTION

02250292	CEFTRIAXONE	PFI
02287633	CEFTRIAXONE	TEV
02292270	CEFTRIAXONE	SDZ
02292874	CEFTRIAXONE	PFI
02325616	CEFTRIAXONE	RAX

2G POWDER FOR SOLUTION

02250306	CEFTRIAXONE	PFI
02292289	CEFTRIAXONE	SDZ
02292882	CEFTRIAXONE	PFI
02325624	CEFTRIAXONE	RAX

10G POWDER FOR SOLUTION

02325632	CEFTRIAXONE SODIUM FOR BP	RAX
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PDIN FOR EXTEMPORANEOUS MIXTURE

99506001	CEFTRIAXONE STERILE INFUSION	UNK
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CEFUROXIME AXETIL

25MG/ML POWDER FOR SOLUTION

02212307	CEFTIN	GSK
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250MG TABLET

02244393	APO-CEFUROXIME	APX
02344823	AURO-CEFUROXIME	APL
02212277	CEFTIN	GSK
02242656	RATIO-CEFUROXIME	TEV

500MG TABLET

02244394	APO-CEFUROXIME	APX
02344831	AURO-CEFUROXIME	APL
02212285	CEFTIN	GSK
02311453	PRO-CEFUROXIM	PDL
02242657	RATIO-CEFUROXIME	TEV

08:12.06 CEPHALOSPORINS

CEPHALEXIN

250MG CAPSULE

00342084	TEVA-CEPHALEXIN	TEV
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500MG CAPSULE

00342114	TEVA-CEPHALEXIN	TEV
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25MG/ML POWDER FOR SUSPENSION

02177862	DOM-CEPHALEXIN	DPC
00015547	KEFLEX	PED
00342106	TEVA-CEPHALEXIN	TEV

50MG/ML POWDER FOR SUSPENSION

02177870	DOM-CEPHALEXIN	DPC
00035645	KEFLEX	PED
00342092	TEVA-CEPHALEXIN	TEV

250MG TABLET

00768723	APO-CEPHALEX	APX
02470578	AURO-CEPHALEXIN	AUR
02177846	DOM-CEPHALEXIN	DPC
00403628	KEFLEX	PED
02177781	PMS-CEPHALEXIN	PMS
00583413	TEVA-CEPHALEXIN	TEV

500MG TABLET

00768715	APO-CEPHALEX	APX
02470586	AURO-CEPHALEXIN	AUR
00828866	CEPHALEXIN-500	PDL
02177854	DOM-CEPHALEXIN	DPC
00244392	KEFLEX	PED
02177803	PMS-CEPHALEXIN	PMS
00583421	TEVA-CEPHALEXIN	TEV

08:12.07 MISCELLANEOUS B-LACTAM ANTIBIOTICS

AZTREONAM

Limited use benefit (prior approval required).

For the management of cystic fibrosis (CF) in patients if the following criteria are met:

- Patient has CF with chronic pulmonary Pseudomonas aeruginosa infections; AND
- Prescribed by a clinician with experience in the diagnosis and treatment of CF.

75MG POWDER FOR SOLUTION

02329840	CAYSTON	GIL
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ERTAPENEM

Limited use benefit (prior approval required).

1G POWDER FOR SOLUTION

02247437	INVANZ	FRS
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MEROPENEM

Limited use benefit (prior approval required).

500MG POWDER FOR SOLUTION

02378787	MEROPENEM	SDZ
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1G POWDER FOR SOLUTION

02378795	MEROPENEM	SDZ
02436507	MEROPENEM	RAX

08:12.12 MACROLIDES

AZITHROMYCIN

20MG/ML POWDER FOR SUSPENSION

02274566	GD-AZITHROMYCIN	PFI
02418452	PMS-AZITHROMYCIN	PMS
02332388	SANDOZ AZITHROMYCIN	SDZ
02223716	ZITHROMAX	PFI

40MG/ML POWDER FOR SUSPENSION

02274574	GD-AZITHROMYCIN	PFI
02418460	PMS-AZITHROMYCIN	PMS
02332396	SANDOZ AZITHROMYCIN	SDZ
02223724	ZITHROMAX	PFI

250MG TABLET

02247423	APO-AZITHROMYCIN	APX
02415542	APO-AZITHROMYCIN	APX
02330881	AZITHROMYCIN	SAN
02442434	AZITHROMYCIN	SIV
02278499	DOM-AZITHROMYCIN	DPC
02274531	GD-AZITHROMYCIN	PFI
02452308	JAMP-AZITHROMYCIN	JMP
02452324	MAR-AZITHROMYCIN	MAR
02261634	PMS-AZITHROMYCIN	PMS
02310600	PRO-AZITHROMYCINE	PDL
02275309	RIVA-AZITHROMYCIN	RIV
02265826	SANDOZ AZITHROMYCIN	SDZ
02267845	TEVA-AZITHROMYCIN	TEV
02212021	ZITHROMAX	PFI

600MG TABLET

02256088	ACT AZITHROMYCIN	ACG
02261642	PMS-AZITHROMYCIN	PMS
02231143	ZITHROMAX	PFI

CLARITHROMYCIN

25MG/ML GRANULES FOR SUSPENSION

02146908	BIAXIN	BGP
02408988	CLARITHROMYCIN	SAN
02390442	TARO-CLARITHROMYCIN	TAR

50MG/ML GRANULES FOR SUSPENSION

02244641	BIAXIN	BGP
02408996	CLARITHROMYCIN	SAN
02390450	TARO-CLARITHROMYCIN	TAR

250MG TABLET

02274744	APO-CLARITHROMYCIN	APX
01984853	BIAXIN	BGP
02324482	CLARITHROMYCIN	PDL
02442469	CLARITHROMYCIN	SIV
02466120	CLARITHROMYCIN	SAN
02471388	M-CLARITHROMYCIN	MAN
02247573	PMS-CLARITHROMYCIN	PMS
02361426	RAN-CLARITHROMYCIN	RBV
02266539	SANDOZ CLARITHROMYCIN	SDZ
02248804	TEVA-CLARITHROMYCIN	TEV

500MG TABLET

02274752	APO-CLARITHROMYCIN	APX
02126710	BIAXIN	BGP
02324490	CLARITHROMYCIN	PDL
02442485	CLARITHROMYCIN	SIV

08:12.12 MACROLIDES

CLARITHROMYCIN

500MG TABLET

02351005	DOM-CLARITHROMYCIN	DPC
02471396	M-CLARITHROMYCIN	MAN
02247574	PMS-CLARITHROMYCIN	PMS
02361434	RAN-CLARITHROMYCIN	RBV
02346532	RIVA-CLARITHROMYCIN	RIV
02266547	SANDOZ CLARITHROMYCIN	SDZ
02248805	TEVA-CLARITHROMYCIN	TEV

500MG TABLET (EXTENDED RELEASE)

02403196	ACT CLARITHROMYCIN XL	ACG
02413345	APO-CLARITHROMYCIN XL	APX
02244756	BIAXIN XL	BGP

ERYTHROMYCIN

250MG CAPSULE (ENTERIC COATED)

00607142	ERYC	PFI
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333MG CAPSULE (ENTERIC COATED)

00873454	ERYC	PFI
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250MG TABLET

00682020	ERYTHRO BASE	AAP
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ERYTHROMYCIN STEARATE

250MG TABLET

00545678	ERYTHRO-S	AAP
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FIDAXOMICIN

Limited use benefit (prior approval required).

For the treatment of confirmed severe Clostridium Difficile Infection (CDI); AND
Fidaxomicin has been prescribed or recommended by an infectious disease specialist or gastroenterologist; AND
There is a documented allergy (immune-mediated reaction) or severe intolerance to oral vancomycin resulting in discontinuation of vancomycin.

OR

- After an unsuccessful but adequate trial of oral vancomycin; AND
- Retreatment with vancomycin is not an option; AND
- The patient is at a high risk of hospitalization due to severe complications; AND
- Fidaxomicin is being used as monotherapy.

Notes:

- a. Severe infection is defined as having any of the following symptoms: white blood cell count > 15,000 mm³ and fever; acute kidney injury with rising serum creatinine ≥ 1.5 times pre-morbid level or ≥ 175 micromoles/L; pseudomembranous colitis, hypotension, shock or megacolon.
- b. An adequate trial of oral vancomycin is considered to be at least 10 days of therapy with a dose of at least 125mg four times daily.
- c. Retreatment with fidaxomicin in recurrent CDI will be considered in symptomatic patients who require treatment of a previously resolved CDI episode. This is defined as a subsequent CDI episode occurring within 2 to 8 weeks of a previous episode from the date of diagnosis.

200MG TABLET

02387174	DIFICID	FRS
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08:12.16 PENICILLINS

AMOXICILLIN

250MG CAPSULE

02352710	AMOXICILLIN	SAN
00628115	APO-AMOXI	APX
02388073	AURO-AMOXICILLIN	AUR
02433060	JAMP-AMOXICILLIN	JMP
00406724	NOVAMOXIN	TEV
02230243	PMS-AMOXICILLIN	PMS

500MG CAPSULE

02352729	AMOXICILLIN	SAN
02401509	AMOXICILLIN	SIV
00628123	APO-AMOXI	APX
02388081	AURO-AMOXICILLIN	AUR
02433079	JAMP-AMOXICILLIN	JMP
02238172	MYLAN-AMOXICILLIN	MYL
00406716	NOVAMOXIN	TEV
02230244	PMS-AMOXICILLIN	PMS
00644315	PRO AMOX	PDL

25MG/ML GRANULES FOR SUSPENSION

00452149	NOVAMOXIN	TEV
01934171	NOVAMOXIN	TEV

50MG/ML GRANULES FOR SUSPENSION

02352753	AMOXICILLIN	SAN
02401541	AMOXICILLIN	SIV
02352788	AMOXICILLIN (SUGAR REDUCED)	SAN
00452130	NOVAMOXIN	TEV
01934163	NOVAMOXIN	TEV

25MG/ML POWDER FOR SUSPENSION

00628131	APO-AMOXI	APX
02230245	PMS-AMOXICILLIN	PMS

50MG/ML POWDER FOR SUSPENSION

00628158	APO-AMOXI	APX
02230880	APO-AMOXI SUGAR FREE	APX
02230246	PMS-AMOXICILLIN	PMS
00644331	PRO-AMOX	PDL

125MG TABLET (CHEWABLE)

02036347	NOVAMOXIN	TEV
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250MG TABLET (CHEWABLE)

02036355	NOVAMOXIN	TEV
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AMOXICILLIN, CLAVULANIC ACID

25MG & 6.25MG/ML POWDER FOR SUSPENSION

02243986	APO-AMOXI CLAV	APX
01916882	CLAVULIN 125 F	GSK

40MG & 5.7MG/ML POWDER FOR SUSPENSION

02288559	APO-AMOXI CLAV	APX
02238831	CLAVULIN 200	GSK

50MG & 12.5MG/ML POWDER FOR SUSPENSION

02243987	APO-AMOXI CLAV	APX
01916874	CLAVULIN 250 F	GSK

80MG & 11.4MG/ML POWDER FOR SUSPENSION

02238830	CLAVULIN 400	GSK
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250MG & 125MG TABLET

02243350	APO-AMOXI CLAV	APX
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500MG & 125MG TABLET

02243351	APO-AMOXI CLAV	APX
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08:12.16 PENICILLINS

AMOXICILLIN, CLAVULANIC ACID

500MG & 125MG TABLET

01916858	CLAVULIN 500 F	GSK
02482576	SANDOZ AMOXI-CLAV	SDZ

875MG & 125MG TABLET

02245623	APO-AMOXI CLAV	APX
02238829	CLAVULIN 875	GSK

AMPICILLIN

250MG CAPSULE

00020877	TEVA-AMPICILLIN	TEV
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500MG CAPSULE

00020885	TEVA-AMPICILLIN	TEV
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1G POWDER FOR SOLUTION

01933345	AMPICILLIN SODIUM	TEV
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2G POWDER FOR SOLUTION

02226995	AMPICILLIN	FKD
01933353	AMPICILLIN SODIUM	TEV
02462346	AMPICILLIN SODIUM FOR BP	AUR

50MG/ML SUSPENSION

00603287	APO AMPI	APX
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PDIN FOR EXTEMPORANEOUS MIXTURE

99506005	AMPICILLIN STERILE INFUSION	UNK
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CLOXACILLIN SODIUM

250MG CAPSULE

00337765	TEVA-CLOXACILLIN	TEV
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500MG CAPSULE

00337773	TEVA-CLOXACILLIN	TEV
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25MG/ML GRANULES FOR SOLUTION

00337757	TEVA-CLOXACILLIN	TEV
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PENICILLIN G BENZATHINE

600,000U/ML SUSPENSION

02291924	BICILLIN	PFI
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PENICILLIN G POTASSIUM

1MU INJECTION

00773727	NOVO-PENICILLIN G POTASSIUM	NOP
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PENICILLIN G SODIUM

10MU POWDER FOR SOLUTION

02220296	PENICILLIN G	FKD
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1000000U POWDER FOR SOLUTION

02220261	PENICILLIN G SODIUM	FKD
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5000000U POWDER FOR SOLUTION

02220288	PENICILLIN G SODIUM	FKD
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PDIN FOR EXTEMPORANEOUS MIXTURE

99506003	PENICILLIN G STERILE INFUSION	UNK
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PENICILLIN V POTASSIUM

25MG/ML POWDER FOR SOLUTION

00642223	APO PEN VK	APX
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60MG/ML POWDER FOR SOLUTION

00642231	APO PEN VK	APX
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300MG TABLET

00642215	PEN-VK	AAP
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08:12.16 PENICILLINS

PIPERACILLIN, TAZOBACTAM

Limited use benefit (prior approval required).

2G & 0.25G POWDER FOR SOLUTION

02299623	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	SDZ
02370158	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	TEV

3G & 0.375G POWDER FOR SOLUTION

02299631	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	SDZ
02308452	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	APX
02362627	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	RAX
02370166	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	TEV

4G & 0.5G POWDER FOR SOLUTION

02299658	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	SDZ
02308460	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	APX
02362635	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	RAX
02370174	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	TEV

12G & 1.5G POWDER FOR SOLUTION

02330547	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	SDZ
02377748	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	RAX

36G & 4.5G POWDER FOR SOLUTION

02439131	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	RAX
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08:12.18 QUINOLONES

CIPROFLOXACIN HYDROCHLORIDE

100MG/ML SUSPENSION

02237514	CIPRO	BAY
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250MG TABLET

02247339	ACT CIPROFLOXACIN	TEV
02229521	APO-CIPROFLOX	APX
02381907	AURO-CIPROFLOXACIN	AUR
02353318	CIPROFLOXACIN	SAN
02386119	CIPROFLOXACIN	SIV
02380358	JAMP-CIPROFLOXACIN	JMP
02379686	MAR-CIPROFLOXACIN	MAR
02423553	MINT-CIPROFLOX	MIN
02248437	PMS-CIPROFLOXACIN	PMS
02317796	PRO-CIPROFLOXACIN	PDL
02303728	RAN-CIPROFLOX	RBY
02251221	RIVA-CIPROFLOXACIN	RIV
02248756	SANDOZ CIPROFLOXACIN	SDZ
02379627	SEPTA-CIPROFLOXACIN	SPT
02266962	TARO-CIPROFLOXACIN	TAR

500MG TABLET

02247340	ACT CIPROFLOXACIN	TEV
02229522	APO-CIPROFLOX	APX
02381923	AURO-CIPROFLOXACIN	AUR
02444887	BIO-CIPROFLOXACIN	BMI

08:12.18 QUINOLONES

CIPROFLOXACIN HYDROCHLORIDE

500MG TABLET

02155966	CIPRO	BAY
02353326	CIPROFLOXACIN	SAN
02386127	CIPROFLOXACIN	SIV
02251280	DOM-CIPROFLOXACIN	DPC
02380366	JAMP-CIPROFLOXACIN	JMP
02379694	MAR-CIPROFLOXACIN	MAR
02423561	MINT-CIPROFLOX	MIN
02248438	PMS-CIPROFLOXACIN	PMS
02317818	PRO-CIPROFLOXACIN	PDL
02303736	RAN-CIPROFLOX	RBY
02251248	RIVA-CIPROFLOXACIN	RIV
02248757	SANDOZ CIPROFLOXACIN	SDZ
02379635	SEPTA-CIPROFLOXACIN	SPT
02266970	TARO-CIPROFLOXACIN	TAR

750MG TABLET

02247341	ACT CIPROFLOXACIN	TEV
02229523	APO-CIPROFLOX	APX
02380374	JAMP-CIPROFLOXACIN	JMP
02379708	MAR-CIPROFLOXACIN	MAR
02423588	MINT-CIPROFLOX	MIN
02248439	PMS-CIPROFLOXACIN	PMS
02303744	RAN-CIPROFLOX	RBY
02251256	RIVA-CIPROFLOXACIN	RIV
02248758	SANDOZ CIPROFLOXACIN	SDZ
02379643	SEPTA-CIPROFLOXACIN	SPT

LEVOFLOXACIN HEMIHYDRATE

Limited use benefit (prior approval not required).

Coverage will be limited to 14 tablets every 14 days, followed by a 14 day lockout.

240MG SOLUTION

02442302	QUINSAIR	UNK
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250MG TABLET

02315424	ACT LEVOFLOXACIN	TEV
02284707	APO-LEVOFLOXACIN	APX
02284677	PMS-LEVOFLOXACIN	PMS
02298635	SANDOZ LEVOFLOXACIN	SDZ
02248262	TEVA-LEVOFLOXACIN	TEV

500MG TABLET

02315432	ACT LEVOFLOXACIN	TEV
02284715	APO-LEVOFLOXACIN	APX
02415879	LEVOFLOXACIN	PDL
02284685	PMS-LEVOFLOXACIN	PMS
02298643	SANDOZ LEVOFLOXACIN	SDZ
02248263	TEVA-LEVOFLOXACIN	TEV

750MG TABLET

02315440	ACT LEVOFLOXACIN	TEV
02325942	APO-LEVOFLOXACIN	APX
02305585	PMS-LEVOFLOXACIN	PMS
02298651	SANDOZ LEVOFLOXACIN	SDZ
02285649	TEVA-LEVOFLOXACIN	TEV

08:12.18 QUINOLONES

MOXIFLOXACIN HYDROCHLORIDE

Limited use benefit (prior approval not required).

Coverage will be limited to 14 tablets every 14 days, followed by a 14 day lockout.

400MG TABLET

02404923	APO-MOXIFLOXACIN	APX
02432242	AURO-MOXIFLOXACIN	AUR
02242965	AVELOX	BAY
02447266	BIO-MOXIFLOXACIN	BMI
02443929	JAMP-MOXIFLOXACIN	JMP
02447061	JAMP-MOXIFLOXACIN	JMP
02447053	MAR-MOXIFLOXACIN	MAR
02457814	MED-MOXIFLOXACIN	GMP
02472791	M-MOXIFLOXACIN	MAN
02462974	MOXIFLOXACIN	PDL
02450976	RIVA-MOXIFLOXACIN	RIV
02383381	SANDOZ MOXIFLOXACIN	SDZ
02375702	TEVA-MOXIFLOXACIN	TEV

NORFLOXACIN

400MG TABLET

02229524	NORFLOXACIN	AAP
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08:12.20 SULFONAMIDES

SULFAMETHOXAZOLE, TRIMETHOPRIM

40MG & 8MG/ML SUSPENSION

00726540	TEVA-TRIMEL	TEV
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100MG & 20MG TABLET

00445266	SULFATRIM PEDIATRIC	APX
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400MG & 80MG TABLET

00445274	SULFATRIM	APX
00510637	TEVA-TRIMEL	TEV

800MG & 160MG TABLET

00512524	PROTRIN DF	PDL
00445282	SULFATRIM DS	APX
00510645	TEVA-TRIMEL DS	TEV

SULFASALAZINE

500MG TABLET

00598461	PMS-SULFASALAZINE	PMS
02064480	SALAZOPYRIN	PFI

500MG TABLET (ENTERIC COATED)

00598488	PMS-SULFASALAZINE	PMS
02064472	SALAZOPYRIN EN	PFI

08:12.24 TETRACYCLINES

DOXYCYCLINE HYCLATE

100MG CAPSULE

00740713	APO-DOXY	APX
00817120	DOXYCIN	RIV
02351234	DOXYCYCLINE	SAN
00725250	TEVA-DOXYCYCLINE	TEV

100MG TABLET

00874256	APO-DOXY	APX
00860751	DOXYCIN	RIV
02351242	DOXYCYCLINE	SAN
00887064	DOXYTAB	PDL

08:12.24 TETRACYCLINES

DOXYCYCLINE HYCLATE

100MG TABLET

02158574	TEVA-DOXYCYCLINE	TEV
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MINOCYCLINE HYDROCHLORIDE

Limited use benefit (prior approval required).

For patients who cannot tolerate other tetracyclines or doxycycline.

For patients with severe widespread acne who have failed on tetracycline or doxycycline.

50MG CAPSULE

02084090	APO-MINOCYCLINE	APX
02153394	MINOCYCLINE	PDL
02230735	MYLAN-MINOCYCLINE	MYL
02294419	PMS-MINOCYCLINE	PMS
02237313	SANDOZ MINOCYCLINE	SDZ
02108143	TEVA-MINOCYCLINE	TEV

100MG CAPSULE

02084104	APO-MINOCYCLINE	APX
02154366	MINOCYCLINE	PDL
02230736	MYLAN-MINOCYCLINE	MYL
02294427	PMS-MINOCYCLINE	PMS
02237314	SANDOZ MINOCYCLINE	SDZ
02108151	TEVA-MINOCYCLINE	TEV

TETRACYCLINE HYDROCHLORIDE

250MG CAPSULE

00580929	TETRACYCLINE	AAP
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08:12.28 MISCELLANEOUS ANTIBIOTICS

CLINDAMYCIN HYDROCHLORIDE

150MG CAPSULE

02245232	APO-CLINDAMYCIN	APX
02436906	AURO-CLINDAMYCIN	AUR
02400529	CLINDAMYCIN	SAN
02248525	CLINDAMYCINE	PDL
00030570	DALACIN C	PFI
02468476	RIVA-CLINDAMYCIN	RIV
02241709	TEVA-CLINDAMYCIN	TEV

300MG CAPSULE

02245233	APO-CLINDAMYCIN	APX
02436914	AURO-CLINDAMYCIN	AUR
02400537	CLINDAMYCIN	SAN
02248526	CLINDAMYCINE	PDL
02182866	DALACIN C	PFI
02241710	TEVA-CLINDAMYCIN	TEV

PDIN FOR EXTEMPORANEOUS MIXTURE

99506008	CLINDAMYCIN STERILE INFUSION	UNK
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CLINDAMYCIN PALMITATE HYDROCHLORIDE

15MG/ML POWDER FOR SOLUTION

00225851	DALACIN C	PFI
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CLINDAMYCIN PHOSPHATE

150MG/ML INJECTION

02139286	CLINDAMYCIN	FKD
02230535	CLINDAMYCIN	SDZ
02230540	CLINDAMYCIN	SDZ

08:12.28 MISCELLANEOUS ANTIBIOTICS

CLINDAMYCIN PHOSPHATE

150MG/ML INJECTION

00260436	DALACIN C PHOSPHATE	PFI
02215683	NOVO-CLINDAMYCIN	NOP

12MG SOLUTION

02408511	CLINDAMYCIN IV INFUSION	SDZ
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18MG SOLUTION

02408538	CLINDAMYCIN IV INFUSION	SDZ
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COLISTIN

Limited use benefit (prior approval required).

For the management of cystic fibrosis (CF) in patients if the following criteria are met:

- Patient has CF with chronic pulmonary Pseudomonas aeruginosa infections; AND
- Prescribed by a clinician with experience in the diagnosis and treatment of CF.

150MG POWDER FOR SOLUTION

02244849	COLISTIMETHATE FOR U.S.P	RAX
00476420	COLY-MYCIN M PARENTERAL	ERF

LINEZOLID

Limited use benefit (prior approval required).

Tablets:

For treatment of proven vancomycin-resistant enterococci (VRE) infections.

For the treatment of proven methicillin-resistant staphylococcus aureus (MRSA) infections in patients who cannot tolerate vancomycin.

I.V. Solution:

When linezolid cannot be administered orally in the above mentioned situations.

Oral Liquid:

When linezolid cannot be administered orally in the above mentioned situations;

Plus at least one of the following:

- For treatment of proven vancomycin-resistant enterococci (VRE) infections
- For the treatment of proven methicillin-resistant staphylococcus aureus (MRSA) infections in patients who cannot tolerate vancomycin.

100MG POWDER FOR SUSPENSION

02243686	ZYVOXAM	PFI
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2MG/ML SOLUTION

02243685	ZYVOXAM	PFI
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600MG TABLET

02426552	APO-LINEZOLID	APX
02422689	SANDOZ LINEZOLID	SDZ
02243684	ZYVOXAM	PFI

08:12.28 MISCELLANEOUS ANTIBIOTICS

RIFAXIMIN

Limited use benefit (prior approval required).

For reducing the risk of overt hepatic encephalopathy (HE) recurrence in patients:

- Who are unable to achieve adequate control of HE recurrence with a maximal tolerated dose of lactulose alone; AND
- When used in combination with a maximal tolerated dose of lactulose.

ST **500MG TABLET**

02410702	ZAXINE	SLX
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VANCOMYCIN HYDROCHLORIDE

Limited use benefit (prior approval required).

Used for the treatment of patients diagnosed with symptomatic Clostridium difficile infection.

Note: Oral vancomycin is not appropriate for systemic infections due to poor absorption from the GI tract.

125MG CAPSULE

02407744	JAMP-VANCOMYCIN	JMP
02430185	PMS-VANCOMYCIN	PMS
00800430	VANOCIN	SEA
02377470	VANCOMYCIN	FKD
02380544	VANCOMYCIN	UNK

250MG CAPSULE

02407752	JAMP-VANCOMYCIN	JMP
00788716	VANOCIN	SEA
02377489	VANCOMYCIN	FKD
02380552	VANCOMYCIN	UNK

VANCOMYCIN HYDROCHLORIDE (INJECTION)

Limited use benefit (prior approval required).

POWDER

99100176	VANCOMYCIN	MDS
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500MG POWDER FOR SOLUTION

02420295	JAMP-VANCOMYCIN	JMP
02406535	MYLAN-VANCOMYCIN	MYL
02139375	VANCOMYCIN	FKD
02230191	VANCOMYCIN	PFI
02394626	VANCOMYCIN	SDZ
02411032	VANCOMYCIN	RAX
02435713	VANCOMYCIN	GMP
02342855	VANCOMYCIN HYDROCHLORIDE	RAX

1,000MG POWDER FOR SOLUTION

02230192	VANCOMYCIN	PFI
02396386	VANCOMYCIN	RAX
02435721	VANCOMYCIN	GMP

1G POWDER FOR SOLUTION

02420309	JAMP-VANCOMYCIN	JMP
02406543	MYLAN-VANCOMYCIN	MYL
02241821	PMS-VANCOMYCIN 1 G	PMS
02139383	VANCOMYCIN	FKD
02394634	VANCOMYCIN	SDZ
02342863	VANCOMYCIN HYDROCHLORIDE	RAX

5G POWDER FOR SOLUTION

02420317	JAMP-VANCOMYCIN	JMP
02406551	MYLAN-VANCOMYCIN	MYL

08:12.28 MISCELLANEOUS ANTIBIOTICS

VANCOMYCIN HYDROCHLORIDE (INJECTION)

Limited use benefit (prior approval required).

5G POWDER FOR SOLUTION

02139243	VANCOMYCIN	FKD
02378337	VANCOMYCIN	PFI
02394642	VANCOMYCIN	SDZ

10G POWDER FOR SOLUTION

02420325	JAMP-VANCOMYCIN	JMP
02406578	MYLAN-VANCOMYCIN	MYL
02241807	VANCOMYCIN	FKD
02378345	VANCOMYCIN	PFI
02394650	VANCOMYCIN	SDZ
02411040	VANCOMYCIN	RAX
02405830	VANCOMYCIN HYDROCHLORIDE	RAX

08:14.04 ALLYLAMINES

TERBINAFINE HYDROCHLORIDE

250MG TABLET

02254727	ACT TERBINAFINE	TEV
02239893	APO-TERBINAFINE	APX
02320134	AURO-TERBINAFINE	AUR
02299275	DOM-TERBINAFINE	DPC
02357070	JAMP-TERBINAFINE	JMP
02031116	LAMISIL	NVR
02240807	PMS-TERBINAFINE	PMS
02294273	PMS-TERBINAFINE	PMS
02262924	RIVA-TERBINAFINE	RIV
02242735	TERBINAFINE	PDL
02353121	TERBINAFINE	SAN
02385279	TERBINAFINE	SIV
02240346	TEVA-TERBINAFINE	TEV

08:14.08 AZOLES

FLUCONAZOLE

150MG CAPSULE

02241895	APO-FLUCONAZOLE	APX
02311690	CANESORAL	BAY
02141442	DIFLUCAN	PFI
02432471	JAMP-FLUCONAZOLE	JMP
02428792	MAR-FLUCONAZOLE	MAR
02243645	NOVO-FLUCONAZOLE	NOP
02246620	PMS-FLUCONAZOLE	PMS
02433702	PRIVA-FLUCONAZOLE	PHA
02255510	RIVA-FLUCONAZOLE	RIV

10MG/ML POWDER FOR SOLUTION

02024152	DIFLUCAN	PFI
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50MG TABLET

02281260	ACT FLUCONAZOLE	ACG
02237370	APO-FLUCONAZOLE	APX
00891800	DIFLUCAN	PFI
02245292	MYLAN-FLUCONAZOLE	MYL
02245643	PMS-FLUCONAZOLE	PMS
02249294	TARO-FLUCONAZOLE	TAR
02236978	TEVA-FLUCONAZOLE	TEV

100MG TABLET

02281279	ACT FLUCONAZOLE	ACG
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08:14.08 AZOLES

FLUCONAZOLE

100MG TABLET

02237371	APO-FLUCONAZOLE	APX
02246109	DOM-FLUCONAZOLE	DPC
02245293	MYLAN-FLUCONAZOLE	MYL
02245644	PMS-FLUCONAZOLE	PMS
02310686	PRO-FLUCONAZOLE	PDL
02249308	TARO-FLUCONAZOLE	TAR
02236979	TEVA-FLUCONAZOLE	TEV

ITRACONAZOLE

100MG CAPSULE

02462559	MINT-ITRACONAZOLE	MIN
02047454	SPORANOX	JSO

POWDER

09991094	ITRACONAZOLE PDR	MDS
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10MG/ML SOLUTION

02231347	SPORANOX	JSO
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KETOCONAZOLE

200MG TABLET

02237235	APO-KETOCONAZOLE	APX
02231061	TEVA-KETOCONAZOLE	TEV

VORICONAZOLE

Limited use benefit (prior approval required).

For the treatment of patients with invasive aspergillosis; OR
For the treatment of culture proven invasive candidiasis with documented resistance to fluconazole.

50MG TABLET

02409674	APO-VORICONAZOLE	APX
02399245	SANDOZ VORICONAZOLE	SDZ
02396866	TEVA-VORICONAZOLE	TEV
02256460	VFEND	PFI

200MG TABLET

02409682	APO-VORICONAZOLE	APX
02399253	SANDOZ VORICONAZOLE	SDZ
02396874	TEVA-VORICONAZOLE	TEV
02256479	VFEND	PFI

08:14.28 POLYENES

NYSTATIN

100,000U/ML SUSPENSION

02125145	DOM-NYSTATIN	DPC
02433443	JAMP-NYSTATIN	JMP
00792667	PMS-NYSTATIN	PMS
02194201	TEVA-NYSTATIN	TEV

08:16.04 ANTITUBERCULOSIS AGENTS

ETHAMBUTOL HYDROCHLORIDE

100MG TABLET

00247960	ETIBI	VAE
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400MG TABLET

00247979	ETIBI	VAE
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ISONIAZID

10MG/ML SOLUTION

00265500	ISOTAMINE	VAE
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08:16.04 ANTITUBERCULOSIS AGENTS

ISONIAZID

10MG/ML SOLUTION

00577812 PDP-ISONIAZID PED

100MG TABLET

00261270 ISOTAMINE VAE

00577790 PDP-ISONIAZID PED

300MG TABLET

00272655 ISOTAMINE VAE

00577804 PDP-ISONIAZID PED

PDIN FOR EXTEMPORANEOUS MIXTURE

99503031 ISONIAZID ORAL LIQUID UNK

PYRAZINAMIDE

500MG TABLET

00618810 PDP-PYRAZINAMIDE PED

00283991 TEBRAZID VAE

RIFABUTIN

150MG CAPSULE

02063786 MYCOBUTIN PFI

RIFAMPIN

150MG CAPSULE

02091887 RIFADIN SAC

00393444 ROFACT VAE

300MG CAPSULE

02092808 RIFADIN SAC

00343617 ROFACT VAE

PDIN FOR EXTEMPORANEOUS MIXTURE

99503022 RIFAMPIN ORAL LIQUID UNK

**08:16.92 MISCELLANEOUS
ANTIMYCOBACTERIALS**

DAPSONE

100MG TABLET

02041510 DAPSONE JAC

02481227 MAR-DAPSONE MAR

08:18.04 ADAMANTANES

AMANTADINE HYDROCHLORIDE

100MG CAPSULE

01990403 PMS-AMANTADINE PED

10MG/ML SYRUP

02022826 PMS-AMANTADINE PED

08:18.08 ANTIRETROVIRALS

ABACAVIR SUFLATE, LAMIVUDINE

600MG & 300MG TABLET

02458381 PMS-ABACAVIR/LAMIVUDINE PMS

ABACAVIR SULFATE

20MG/ML SOLUTION

02240358 ZIAGEN VII

300MG TABLET

02396769 APO-ABACAVIR APX

02480956 MINT-ABACAVIR MIN

02240357 ZIAGEN VII

08:18.08 ANTIRETROVIRALS

ABACAVIR SULFATE, LAMIVUDINE

600MG & 300MG TABLET

02399539 APO-ABACAVIR-LAMIVUDINE APX

02454513 AURO-ABACAVIR/LAMIVUDINE AUR

02269341 KIVEXA VII

02450682 MYLAN-ABACAVIR/LAMIVUDINE MYL

02416662 TEVA-ABACAVIR/LAMIVUDINE TEV

**ABACAVIR SULFATE, LAMIVUDINE,
DOLUTEGRAVIR SODIUM**

600MG & 300MG & 50MG TABLET

02430932 TRIUMEQ VII

ABACAVIR SULFATE, LAMIVUDINE, ZIDOVUDINE

300MG & 150MG & 300MG TABLET

02416255 APO-ABACAVIR-LAMIVUDINE- APX
ZIDOVUDINE

02244757 TRIZIVIR VII

ATAZANAVIR SULFATE

150MG CAPSULE

02456877 MYLAN-ATAZANAVIR MYL

02248610 REYATAZ BMS

02443791 TEVA-ATAZANAVIR TEV

200MG CAPSULE

02456885 MYLAN-ATAZANAVIR MYL

02248611 REYATAZ BMS

02443813 TEVA-ATAZANAVIR TEV

300MG CAPSULE

02456893 MYLAN-ATAZANAVIR MYL

02294176 REYATAZ BMS

02443821 TEVA-ATAZANAVIR TEV

DARUNAVIR ETHANOLATE

75MG TABLET

02338432 PREZISTA JSO

150MG TABLET

02369753 PREZISTA JSO

400MG TABLET

02324016 PREZISTA JSO

600MG TABLET

02324024 PREZISTA JSO

800MG TABLET

02393050 PREZISTA JSO

DARUNAVIR ETHANOLATE, COBICISTAT

150MG & 800MG TABLET

02426501 PREZCOBIX JSO

DIDANOSINE

125MG CAPSULE (ENTERIC COATED)

02244596 VIDEX EC BMS

200MG CAPSULE (ENTERIC COATED)

02244597 VIDEX EC BMS

250MG CAPSULE (ENTERIC COATED)

02244598 VIDEX EC BMS

400MG CAPSULE (ENTERIC COATED)

02244599 VIDEX EC BMS

08:18.08 ANTIRETROVIRALS

DOLUTEGRAVIR SODIUM

50MG TABLET

02414945 TIVICAY VII

DOLUTEGRAVIR SODIUM, RILPIVIRINE HYDROCHLORIDE

50MG & 25MG TABLET

02475774 JULUCA VII

EFAVIRENZ

50MG CAPSULE

02239886 SUSTIVA BMS

200MG CAPSULE

02239888 SUSTIVA BMS

600MG TABLET

02418428 AURO-EFAVIRENZ AUR

02381524 MYLAN-EFAVIRENZ MYL

02246045 SUSTIVA BMS

02389762 TEVA-EFAVIRENZ TEV

EFAVIRENZ, EMTRICITABINE, TENOFOVIR DISOPROXIL FUMARATE

600MG & 200MG & 300MG TABLET

02468247 APO-EFAVIRENZ-EMTRICITABINE-TENOFOVIR APX

02300699 ATRIPLA GIL

02461412 MYLAN-EFAVIRENZ/EMTRICITABINE/TENOFOVIR DISOPROXIL FUMARATE MYL

02393549 TEVA-EFAVIRENZ/EMTRICITABINE/TENOFOVIR TEV

EMTRICITABINE, COBICISTAT, ELVITEGRAVIR, TENOFOVIR ALAFENAMIDE

200MG & 150MG & 150MG & 10MG TABLET

02449498 GENVOYA GIL

EMTRICITABINE, RILPIVIRINE HYDROCHLORIDE, TENOFOVIR ALAFENAMIDE

200MG & 25MG & 25MG TABLET

02461463 ODEFSEY GIL

ETRAVIRINE

100MG TABLET

02306778 INTELENCE JSO

200MG TABLET

02375931 INTELENCE JSO

FOSAMPRENAVIR CALCIUM

50MG/ML SUSPENSION

02261553 TELZIR VII

700MG TABLET

02261545 TELZIR VII

LAMIVUDINE

5MG SOLUTION

02239194 HEPTO VIR GSK

10MG/ML SOLUTION

02192691 3TC VII

08:18.08 ANTIRETROVIRALS

LAMIVUDINE

100MG TABLET

02393239 APO-LAMIVUDINE HBV APX

02239193 HEPTO VIR GSK

150MG TABLET

02192683 3TC VII

02369052 APO-LAMIVUDINE APX

300MG TABLET

02247825 3TC VII

02369060 APO-LAMIVUDINE APX

LAMIVUDINE, ZIDOVUDINE

150MG & 300MG TABLET

02375540 APO-LAMIVUDINE-ZIDOVUDINE APX

02414414 AURO-LAMIVUDINE/ZIDOVUDINE AUR

02239213 COMBIVIR VII

02387247 TEVA-LAMIVUDINE/ZIDOVUDINE TEV

LOPINAVIR, RITONAVIR

80MG & 20MG/ML SOLUTION

02243644 KALETRA ABV

100MG & 25MG TABLET

02312301 KALETRA ABV

200MG & 50MG TABLET

02285533 KALETRA ABV

MARAVIROC

150MG TABLET

02299844 CELSENTRI VII

300MG TABLET

02299852 CELSENTRI VII

NELFINAVIR MESYLATE

50MG/G POWDER

02238618 VIRACEPT PFI

250MG TABLET

02238617 VIRACEPT PFI

625MG TABLET

02248761 VIRACEPT PFI

NEVIRAPINE

200MG TABLET

02318601 AURO-NEVIRAPINE APL

02405776 JAMP NEVIRAPINE JMP

02387727 MYLAN-NEVIRAPINE MYL

02238748 VIRAMUNE BOE

400MG TABLET (EXTENDED RELEASE)

02427931 APO-NEVIRAPINE XR APX

02367289 VIRAMUNE XR BOE

RALTEGRAVIR POTASSIUM

400MG TABLET

02301881 ISENTRESS FRS

RILPIVIRINE HYDROCHLORIDE

25MG TABLET

02370603 EDURANT JSO

08:18.08 ANTIRETROVIRALS

RITONAVIR

100MG TABLET

02357593 NORVIR ABV

SAQUINAVIR MESYLATE

500MG TABLET

02279320 INVIRASE HLR

STAVUDINE

15MG CAPSULE

02216086 ZERIT BMS

20MG CAPSULE

02216094 ZERIT BMS

30MG CAPSULE

02216108 ZERIT BMS

40MG CAPSULE

02216116 ZERIT BMS

TENOFOVIR DISOPROXIL FUMARATE

Limited use benefit (prior approval required).

For the treatment of patients with HIV-1 infection who have failed or have experienced adverse events to an alternative agent.

For the treatment of patients with chronic hepatitis B infection who have cirrhosis documented on radiologic or histologic grounds and a HBV concentration above 2,000 IU/mL.

245MG TABLET

02247128 VIREAD GIL

300MG TABLET

02451980 APO-TENOFOVIR APX

02460173 AURO-TENOFOVIR AUR

02479087 JAMP-TENOFOVIR JMP

02452634 MYLAN-TENOFOVIR DISOPROXIL MYL

02472511 NAT-TENOFOVIR NPH

02453940 PMS-TENOFOVIR PMS

02403889 TEVA-TENOFOVIR TEV

TENOFOVIR DISOPROXIL FUMARATE, EMTRICITABINE

200MG & 300MG TABLET

02274906 TRUVADA GIL

300MG & 200MG TABLET

02452006 APO-EMTRICITABINE-TENOFOVIR APX

02443902 MYLAN-EMTRICITABINE/TENOFOVIR MYL

DISOPROXIL

02461110 PMS-EMTRICITABINE-TENOFOVIR PMS

02399059 TEVA-EMTRICITABINE/TENOFOVIR TEV

TENOFOVIR DISOPROXIL FUMARATE, EMTRICITABINE, COBICISTAT, ELVITEGRAVIR

150MG & 200MG & 150MG & 300MG TABLET

02397137 STRIBILD GIL

TENOFOVIR DISOPROXIL FUMARATE, EMTRICITABINE, RILPIVIRINE HYDROCHLORIDE

200MG & 25MG & 300MG TABLET

02374129 COMPLERA GIL

08:18.08 ANTIRETROVIRALS

TIPRANA VIR

250MG CAPSULE

02273322 APTIVUS BOE

ZIDOVUDINE

100MG CAPSULE

01946323 APO-ZIDOVUDINE APX

01902660 RETROVIR VII

10MG/ML SYRUP

01902652 RETROVIR VII

08:18.20 INTERFERONS

INTERFERON ALFA-2B

6,000,000IU/ML SOLUTION

02238674 INTRON A FRS

10,000,000IU/ML SOLUTION

02238675 INTRON A FRS

10,000,000IU/VIAL SOLUTION

02223406 INTRON A FRS

PEGINTERFERON ALFA-2A

Limited use benefit (prior approval required).

For the treatment of patients with chronic hepatitis B infection who have a HBV DNA concentration above 2,000 IU/mL without decompensated cirrhosis, upon the written request of a hepatologist or other specialist in this area.

180MCG/0.5ML SOLUTION

02248077 PEGASYS HLR

PEGINTERFERON ALFA-2B, RIBAVIRIN

Limited use benefit (prior approval required).

For the treatment of chronic hepatitis C in patients who are treatment naive, upon the written request of a hepatologist or other specialist in this area.

- For genotypes 1, 4, 5 and 6, an initial 24 week supply will be approved. A further 24 week supply may be approved if patient has a viral reduction of at least 2 logs or HCV is undetectable at 12 weeks (48 weeks total).
- For genotypes 2 or 3, initial coverage for a maximum of 24 weeks will be approved. Renewals will not be covered.

50MCG/0.5ML & 200MG KIT

02254573 PEGETRON KIT FRS

PEGINTERFERON BETA-1A

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

94MCG INJECTION

02444402 PLEGRIDY UNK

125MCG LIQUID

02444399 PLEGRIDY UNK

08:18.28 NEURAMINIDASE INHIBITORS

OSELTAMIVIR

30MG CAPSULE

02472635 NAT-OSELTAMIVIR NPH

45MG CAPSULE

02472643 NAT-OSELTAMIVIR NPH

08:18.32 NUCLEOSIDES AND NUCLEOTIDES

ACYCLOVIR

40MG/ML SUSPENSION

00886157 ZOVIRAX GSK

200MG TABLET

02207621 APO-ACYCLOVIR APX

02242784 MYLAN-ACYCLOVIR MYL

02285959 TEVA-ACYCLOVIR TEV

400MG TABLET

02207648 APO-ACYCLOVIR APX

02242463 MYLAN-ACYCLOVIR MYL

02285967 TEVA-ACYCLOVIR TEV

800MG TABLET

02207656 APO-ACYCLOVIR APX

02242464 MYLAN-ACYCLOVIR MYL

02285975 TEVA-ACYCLOVIR TEV

ADEFOVIR DIPIVOXIL

Limited use benefit (prior approval required).

For the treatment of chronic hepatitis B infection when used in combination with lamivudine in patients who have developed failure to lamivudine, as defined by an increase in HBV DNA of $\geq 1 \log_{10}$ IU/mL above the nadir, measured on two separate occasions within an interval of at least one month, after the first three months of lamivudine therapy, and when failure to lamivudine is not due to poor adherence to therapy.

10MG TABLET

02420333 APO-ADEFOVIR APX

02247823 HEPSERA GIL

ENTECAVIR MONOHYDRATE

Limited use benefit (prior approval required).

For the treatment of chronic hepatitis B infection in patients with cirrhosis documented on radiologic or histologic grounds and a HBV DNA concentration above 2000IU/mL.

0.5MG TABLET

02396955 APO-ENTECAVIR APX

02448777 AURO-ENTECAVIR AUR

02282224 BARACLUDE BMS

02467232 JAMP ENTECAVIR JMP

02430576 PMS-ENTECAVIR PMS

FAMCICLOVIR

125MG TABLET

02305682 ACT FAMCICLOVIR ACG

02292025 APO-FAMCICLOVIR APX

02229110 FAMVIR NVR

02278081 PMS-FAMCICLOVIR PMS

02278634 SANDOZ FAMCICLOVIR SDZ

250MG TABLET

02305690 ACT FAMCICLOVIR ACG

08:18.32 NUCLEOSIDES AND NUCLEOTIDES

FAMCICLOVIR

250MG TABLET

02292041 APO-FAMCICLOVIR APX

02229129 FAMVIR NVR

02278103 PMS-FAMCICLOVIR PMS

02278642 SANDOZ FAMCICLOVIR SDZ

500MG TABLET

02305704 ACT FAMCICLOVIR ACG

02292068 APO-FAMCICLOVIR APX

02177102 FAMVIR NVR

02278111 PMS-FAMCICLOVIR PMS

02278650 SANDOZ FAMCICLOVIR SDZ

GANCICLOVIR SODIUM

500MG POWDER FOR SOLUTION

02162695 CYTOVENE CHE

VALACYCLOVIR HYDROCHLORIDE

500MG TABLET

02295822 APO-VALACYCLOVIR APX

02405040 AURO-VALACYCLOVIR AUR

02444860 BIO-VALACYCLOVIR BMI

02307936 DOM-VALACYCLOVIR DPC

02441454 JAMP-VALACYCLOVIR JMP

02441586 MAR-VALACYCLOVIR MAR

02351579 MYLAN-VALACYCLOVIR MYL

02298457 PMS-VALACYCLOVIR PMS

02441861 PRIVA-VALACYCLOVIR PHA

02315173 PRO-VALACYCLOVIR PDL

02316447 RIVA-VALACYCLOVIR RIV

02347091 SANDOZ VALACYCLOVIR SDZ

02357534 TEVA-VALACYCLOVIR TEV

02442000 VALACYCLOVIR SIV

02454645 VALACYCLOVIR SAN

02219492 VALTREX GSK

VALGANCICLOVIR HYDROCHLORIDE

50MG POWDER FOR SOLUTION

02306085 VALCYTE HLR

450MG TABLET

02393824 APO-VALGANCICLOVIR APX

02435179 AURO-VALGANCICLOVIR AUR

02413825 TEVA-VALGANCICLOVIR TEV

02245777 VALCYTE HLR

08:18.40 HCV ANTIVIRALS

DACLATASVIR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria: Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

30MG TABLET

02444747 DAKLINZA BMS

60MG TABLET

02444755 DAKLINZA BMS

ELBASVIR, GRAZOPREVR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria: Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

50MG & 100MG TABLET

02451131 ZEPATIER FRS

GLECAPREVIR, PIBRENTASVIR

Limited use benefit (prior approval required).

For treatment-naïve or treatment-experienced adult patients with genotypes 1, 2, 3, 4, 5, 6 with; OR For the treatment of direct acting antivirals (DAA)-experienced2 adult patients with genotype 1 with:
 • Chronic hepatitis C at any fibrosis stage (F0-F4); AND
 • Detectable levels of HCV RNA in the last 12 months;

For genotypes 1, 2, 3, 4, 5 or 6, treatment-experienced is defined as a patient who has been previously treated with interferon, peginterferon (P), ribavirin (R) and/or sofosbuvir (SOF) (PR, SOF + PR, SOF + RBV), but no prior treatment experience with an NS3/4A protease inhibitor or NS5A inhibitor.

For genotype 1, DAA treatment-experienced is defined as a patient who has been previously treated with DAA regimens containing NS5A inhibitor [daclatasvir (DCV) + SOF or DCV + PR or ledipasvir/sofosbuvir, but no prior treatment experience with NS3/4A protease inhibitors] or containing NS3/4A protease inhibitors [simeprevir+SOF or simeprevir+PR or boceprevir+PR or telaprevir+PR, but no prior treatment experience with an NS5Ainhibitor].

100MG & 40MG TABLET

02467550 MAVIRET ABV

08:18.40 HCV ANTIVIRALS

RIBAVIRIN

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria: Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

200MG TABLET

02439212 IBAVYR PED

400MG TABLET

02425890 IBAVYR PED

600MG TABLET

02425904 IBAVYR PED

SOFOSBUVIR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria: Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

400MG TABLET

02418355 SOVALDI GIL

SOFOSBUVIR, LEDIPASVIR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria: Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

400MG & 90MG TABLET

02432226 HARVONI GIL

08:18.40 HCV ANTIVIRALS

SOFOSBUVIR, VELPATASVIR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria: Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

400MG & 100MG TABLET

02456370 EPCLUSA GIL

SOFOSBUVIR, VELPATASVIR, VOXILAPREVIR

Limited use benefit (prior approval required).

For treatment-experienced adult patients with:
 • Chronic hepatitis C at any fibrosis stage (F0-F4); AND
 • Detectable levels of HCV RNA in the last 12 months; AND
 Treatment-experienced having failed a prior therapy with an HCV regimen containing:
 • NS5A inhibitor: daclatasvir (Daklinza), elbasvir (part of Zepatier), ledipasvir (part of Harvoni), ombitasvir (part of Holkira Pak), velpatasvir (part of Epclusa) for genotype 1, 2, 3, 4, 5 or 6; OR
 • sofosbuvir (Sovaldi) without an NS5A inhibitor for genotype 1, 2, 3 or 4.

400MG & 100MG & 100MG TABLET

02467542 VOSEVI GIL

08:30.04 AMEBICIDES

PAROMOMYCIN SULFATE

250MG CAPSULE

02078759 HUMATIN ERF

08:30.08 ANTIMALARIALS

CHLOROQUINE PHOSPHATE

250MG TABLET

00021261 TEVA-CHLOROQUINE TEV

HYDROXYCHLOROQUINE SULFATE

200MG TABLET

02246691 APO-HYDROXYQUINE APX

02424991 MINT-HYDROXYCHLOROQUINE MIN

02017709 PLAQUENIL SAC

02311011 PRO-HYDROXYQUINE PDL

PRIMAQUINE PHOSPHATE

26.3MG TABLET

02017776 PRIMAQUINE SAC

08:30.92 MISCELLANEOUS ANTIPROTOZOALS

ATOVAQUONE

150MG/ML SUSPENSION

02217422 MEPRON GSK

08:30.92 MISCELLANEOUS ANTIPROTOZOALS

METRONIDAZOLE

500MG CAPSULE

02248562 APO-METRONIDAZOLE APX

02470284 AURO-METRONIDAZOLE AUR

01926853 FLAGYL ODN

250MG TABLET

00545066 METRONIDAZOLE AAP

PDIN FOR EXTEMPORANEOUS MIXTURE

99503012 METRONIDAZOLE ORAL LIQUID UNK

08:36.00 URINARY ANTI-INFECTIVES

FOSFOMYCIN TROMETHAMINE

Limited use benefit (prior approval required).

For the treatment of women (>12 years old) with:
 • Urinary tract infections with organisms resistant to first line therapy; OR
 • Urinary tract infections in pregnancy when first-line agents are contraindicated.

3G/PK POWDER FOR SOLUTION

02240335 MONUROL PAL

3G POWDER FOR SOLUTION

02473801 JAMP-FOSFOMYCIN JMP

NITROFURANTOIN

100MG CAPSULE

02063662 MACROBID ALL

02455676 PMS-NITROFURANTOIN PMS

50MG CAPSULE (DELAYED RELEASE)

02231015 TEVA-NITROFURANTOIN TEV

100MG CAPSULE (DELAYED RELEASE)

02231016 TEVA-NITROFURANTOIN TEV

50MG TABLET

00319511 NITROFURANTOIN AAP

100MG TABLET

00312738 NITROFURANTOIN AAP

PDIN FOR EXTEMPORANEOUS MIXTURE

99503004 NITRO-FURANTOIN ORAL LIQUID UNK

TRIMETHOPRIM

100MG TABLET

02243116 TRIMETHOPRIM AAP

200MG TABLET

02243117 TRIMETHOPRIM AAP

PDIN FOR EXTEMPORANEOUS MIXTURE

99503017 TRIMETHOPRIM ORAL LIQUID UNK

10:00 ANTINEOPLASTIC AGENTS

10:00.00 ANTINEOPLASTIC AGENTS

ABIRATERONE ACETATE

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of metastatic castration resistant prostate cancer patients (mCRPC) who are asymptomatic or mildly symptomatic after failure of androgen deprivation therapy (ADT) and who have not received prior chemotherapy if they meet the following criteria:

- Used in combination with prednisone; AND
- Patient has an ECOG performance status of 0 or 1.

For the treatment of metastatic castration resistant prostate cancer patients (mCRPC) who progressed on docetaxel-based chemotherapy if they meet the following criteria:

- Used in combination with prednisone; AND
- Patient has an ECOG performance status ≤ 2 ; AND
- Abiraterone is not used as an add-on therapy to enzalutamide (Xtandi); AND
- Abiraterone has not been used in the pre-docetaxel setting.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression

250MG TABLET

02371065 ZYTIGA JSO

500MG TABLET

02457113 ZYTIGA JSO

AFATINIB DIMALEATE

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the treatment of patients with advanced Non-Small Cell Lung Cancer (NSCLC) who meet ALL of the following criteria:

- First line treatment of patients; AND
- EGFR mutation positive; AND
- Advanced or metastatic adenocarcinoma of the lung; AND
- An ECOG performance status of 0 or 1.

Criteria for renewal every 6 months:

- There is no objective evidence of disease progression.

Use of afatinib precludes the use of any other EGFR inhibitor as a subsequent line of therapy.

20MG TABLET

02415666 GIOTRIF BOE

30MG TABLET

02415674 GIOTRIF BOE

40MG TABLET

02415682 GIOTRIF BOE

10:00.00 ANTINEOPLASTIC AGENTS

ALECTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

First-line treatment of patients with locally advanced or metastatic non-small cell lung cancer (NSCLC); OR
Second-line treatment of patients with locally advanced not amenable to curative therapy or metastatic NSCLC who have disease progression on or intolerance to crizotinib.

AND

To be used as monotherapy; AND

Disease is anaplastic lymphoma kinase (ALK)-positive; AND

Patient has a good performance status.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

150MG CAPSULE

02458136 ALECEASARO HLR

ANASTROZOLE

1MG TABLET

02351218 ACH-ANASTROZOLE ACC

02394898 ACT ANASTROZOLE TEV

02395649 ANASTROZOLE PDL

02442736 ANASTROZOLE SAN

02374420 APO-ANASTROZOLE APX

02224135 ARIMIDEX AZC

02392488 BIO-ANASTROZOLE BMI

02339080 JAMP-ANASTROZOLE JMP

02379562 MAR-ANASTROZOLE MAR

02379104 MED-ANASTROZOLE GMP

02393573 MINT-ANASTROZOLE MIN

02417855 NAT-ANASTROZOLE NPH

02320738 PMS-ANASTROZOLE PMS

02328690 RAN-ANASTROZOLE RBY

02392259 RIVA-ANASTROZOLE RIV

02338467 SANDOZ ANASTROZOLE SDZ

02365650 TARO-ANASTROZOLE TAR

02427818 VAN-ANASTROZOLE VAN

AXITINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the second-line treatment of patients with advanced or metastatic clear cell renal carcinoma after failure of prior therapy with a first-line agent.

Patients are only eligible for either everolimus or axitinib in the second-line setting, but not sequential use of both agents except in cases of intolerance.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

1MG TABLET

02389630 INLYTA PFI

5MG TABLET

02389649 INLYTA PFI

BICALUTAMIDE

50MG TABLET

02325985 ACH-BICALUTAMIDE ACC

10:00.00 ANTINEOPLASTIC AGENTS

BICALUTAMIDE

50MG TABLET

02296063	APO-BICALUTAMIDE	APX
02382423	BICALUTAMIDE	SIV
02184478	CASODEX	AZC
02357216	JAMP-BICALUTAMIDE	JMP
02275589	PMS-BICALUTAMIDE	PMS
02311038	PRO-BICALUTAMIDE	PDL
02371324	RAN-BICALUTAMIDE	RBY
02276089	SANDOZ BICALUTAMIDE	SDZ
02270226	TEVA-BICALUTAMIDE	TEV
02428709	VAN-BICALUTAMIDE	VAN

BOSUTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

Patients has Philadelphia chromosome positive (Ph+) chronic myelogenous leukemia (CML); AND

Patient has an ECOG performance status of 0 to 2; AND

- Documented resistance/disease progression to at least one prior oral tyrosine kinase inhibitor [TKI] (imatinib, dasatinib or nilotinib); OR

- Documented intolerance to one prior oral TKI (imatinib, dasatinib or nilotinib) where subsequent treatment with an alternative oral TKI is not clinically appropriate.

Criteria for renewal every 12 months:

Confirmation from the clinician that the patient has experienced hematologic and/or cytogenetic response and is expected to continue to do so AND has not developed unacceptable toxicities.

100MG TABLET

02419149	BOSULIF	PFI
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500MG TABLET

02419157	BOSULIF	PFI
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BUSERELIN ACETATE

6.3MG/IMPLANT IMPLANT

02228955	SUPREFACT DEPOT 2 MONTHS	SAC
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9.45MG/IMPLANT IMPLANT

02240749	SUPREFACT DEPOT 3 MONTHS	SAC
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1MG/ML SOLUTION

02225166	SUPREFACT	SAC
02225158	SUPREFACT (NASAL)	SAC

BUSULFAN

2MG TABLET

00004618	MYLERAN	ASP
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CAPECITABINE

150MG TABLET

02426757	ACH-CAPECITABINE	ACC
02421917	SANDOZ CAPECITABINE	SDZ
02457490	TARO-CAPECITABINE	TAR
02400022	TEVA-CAPECITABINE	TEV
02238453	XELODA	HLR

500MG TABLET

02426765	ACH-CAPECITABINE	ACC
02421925	SANDOZ CAPECITABINE	SDZ

10:00.00 ANTINEOPLASTIC AGENTS

CAPECITABINE

500MG TABLET

02457504	TARO-CAPECITABINE	TAR
02400030	TEVA-CAPECITABINE	TEV
02238454	XELODA	HLR

CERITINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- Second-line treatment of patients with locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer (NSCLC) who have disease progression on or intolerance to crizotinib; AND

- To be used as monotherapy; AND

- Disease is anaplastic lymphoma kinase (ALK)-positive; AND
- Patient has an ECOG performance status of 0 to 2.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

150MG CAPSULE

02436779	ZYKADIA	NVR
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CHLORAMBUCIL

2MG TABLET

00004626	LEUKERAN	ASP
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COBIMETINIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the first-line treatment of patients with metastatic or unresectable melanoma in combination with vemurafenib (Zelboraf).

AND for patients who meet the following criteria:

- Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND

- Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND

- Patient has an ECOG performance status of 0 to 1.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

20MG TABLET

02452340	COTELLIC	HLR
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10:00.00 ANTINEOPLASTIC AGENTS

CRIZOTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

First-line treatment of patients with advanced non-small cell lung cancer (NSCLC); OR
 Second-line treatment of patients with advanced NSCLC who have received one prior chemotherapy regimen.*
 AND
 • Patient is anaplastic lymphoma kinase (ALK)-positive; AND
 • Patient has an ECOG performance status of 0 to 2.

*Patients who have progressed during or following first-line therapy with crizotinib are not eligible to receive crizotinib as a second-line therapy.

Criteria for renewal every 12 months:

The patient has experienced a hematologic and/or cytogenic response to crizotinib and is expected to continue to do so.

200MG CAPSULE

02384256 XALKORI PFI

CYCLOPHOSPHAMIDE

25MG TABLET

02241795 PROCYTOX BAX

50MG TABLET

02241796 PROCYTOX BAX

DABRAFENIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

• For the first-line treatment of patients with metastatic or unresectable melanoma as monotherapy; OR
 • For the first-line treatment of patients with metastatic or unresectable melanoma in combination with trametinib (Mekinist)

AND for patients who meet the following criteria:

• Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND
 • Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND
 • Patient has an ECOG performance status of 0 to 1; AND
 • Patient is previously untreated.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

50MG CAPSULE

02409607 TAFINLAR NVR

75MG CAPSULE

02409615 TAFINLAR NVR

10:00.00 ANTINEOPLASTIC AGENTS

ENZALUTAMIDE

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of patients with metastatic castration-resistant prostate cancer (mCRPC) who are/have:
 • Asymptomatic or mildly symptomatic after failure of androgen deprivation therapy (ADT) who have not received prior chemotherapy; AND
 • Have an ECOG performance status of 0 or 1 with no risk factors for seizures; OR
 • Progressed on docetaxel-based chemotherapy with an ECOG performance status ≤ 2 and no risk factors for seizures; AND
 • Would be an alternative to abiraterone for patients in the post-docetaxel setting but would not be an add-on therapy to abiraterone treatment.

Patients previously treated with abiraterone would not be eligible for enzalutamide unless unable to tolerate abiraterone.

Use of enzalutamide in the post-docetaxel setting is not permitted if previously used in the pre-chemotherapy setting.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

40MG CAPSULE

02407329 XTANDI AST

ERLOTINIB HYDROCHLORIDE

Limited use benefit (prior approval required).

Treatment of non-small cell lung cancer (NSCLC) after failure of at least one prior chemotherapy regimen, and whose EGFR expression status is positive or unknown.

25MG TABLET

02461862 APO-ERLOTINIB APX

02483912 NAT-ERLOTINIB NPH

02269007 TARCEVA HLR

02377691 TEVA-ERLOTINIB TEV

100MG TABLET

02461870 APO-ERLOTINIB APX

02454386 PMS-ERLOTINIB PMS

02269015 TARCEVA HLR

02377705 TEVA-ERLOTINIB TEV

150MG TABLET

02461889 APO-ERLOTINIB APX

02454394 PMS-ERLOTINIB PMS

02269023 TARCEVA HLR

02377713 TEVA-ERLOTINIB TEV

ETOPOSIDE

50MG CAPSULE

00616192 VEPESID BMS

10:00.00 ANTINEOPLASTIC AGENTS

EVEROLIMUS

Limited use benefit (prior approval required).

For the treatment of:

- Advanced breast cancer according to established criteria.
- Advanced or metastatic renal cell carcinoma (mRCC) according to established criteria.
- Progressive, unresectable, well or moderately differentiated, locally advanced or metastatic pancreatic neuroendocrine tumors (pNET) according to established criteria.
- Non-functional neuroendocrine tumors (NETs) of gastrointestinal or lung origin (GIL) according to established criteria.

(Please refer to Appendix A).

2.5MG TABLET

02369257 AFINITOR NVR

5MG TABLET

02339501 AFINITOR NVR

10MG TABLET

02339528 AFINITOR NVR

2MG TABLET FOR SUSPENSION

02425645 AFINITOR DISPERZ NVR

3MG TABLET FOR SUSPENSION

02425653 AFINITOR DISPERZ NVR

5MG TABLET FOR SUSPENSION

02425661 AFINITOR DISPERZ NVR

EXEMESTANE

25MG TABLET

02390183 ACT EXEMESTANE ACG

02419726 APO-EXEMESTANE APX

02242705 AROMASIN PFI

02407841 MED-EXEMESTANE GMP

02408473 TEVA-EXEMESTANE TEV

FLUDARABINE PHOSPHATE

10MG TABLET

02246226 FLUDARA SAC

FLUTAMIDE

250MG TABLET

02238560 FLUTAMIDE AAP

02230104 PMS-FLUTAMIDE PMS

GEFITINIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the treatment of patients with locally advanced or metastatic non-small cell lung cancer (NSCLC) who meet ALL of the following criteria:

- First-line treatment; AND
- EGFR mutation positive; AND
- Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

250MG TABLET

02468050 APO-GEFITINIB APX

02248676 IRESSA AZC

10:00.00 ANTINEOPLASTIC AGENTS

HYDROXYUREA

500MG CAPSULE

02247937 APO-HYDROXYUREA APX

00465283 HYDREA BMS

02242920 MYLAN-HYDROXYUREA MYL

IBRUTINIB

Limited use benefit (prior approval required).

For the treatment of:

- Previously untreated chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL) (first-line) according to established criteria.
- Chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL) (second-line) according to established criteria.
- Relapsed/refractory mantle cell lymphoma (MCL) according to established criteria.

(Please refer to Appendix A).

140MG CAPSULE

02434407 IMBRUVICA JSO

IDELALISIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

- For the treatment of patients with relapsed chronic lymphocytic leukemia (CLL) in combination with rituximab. Treatment should continue until unacceptable toxicity or disease progression.

Criteria for renewal every 6 months:

- There is no objective evidence of disease progression.

100MG TABLET

02438798 ZYDELIG GIL

150MG TABLET

02438801 ZYDELIG GIL

IMATINIB MESYLATE

Limited use benefit (prior approval required).

- For the treatment of patients with chronic myeloid leukemia (CML) in blast crisis, accelerated phase, or in chronic phase.
- For the treatment of patients with gastrointestinal stromal tumour.
- For newly diagnosed adult patients with Philadelphia chromosome-positive (CML).
- For the treatment of adult patients with Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ ALL).

100MG TABLET

02355337 APO-IMATINIB APX

02253275 GLEEVEC NVR

02397285 NAT-IMATINIB NPH

02431114 PMS-IMATINIB PMS

02399806 TEVA-IMATINIB TEV

400MG TABLET

02355345 APO-IMATINIB APX

02253283 GLEEVEC NVR

02397293 NAT-IMATINIB NPH

02431122 PMS-IMATINIB PMS

02399814 TEVA-IMATINIB TEV

10:00.00 ANTINEOPLASTIC AGENTS

LENALIDOMIDE

Limited use benefit (prior approval required).

For the treatment of:

- Myelodysplastic syndrome (MDS)
- Refractory/relapsed Multiple Myeloma after one prior therapy (MM-AOPT)
- Newly diagnosed Multiple Myeloma for patients who are not eligible for autologous stem cell transplant - (MM-TNE)
- Maintenance treatment for newly diagnosed Multiple Myeloma post-autologous stem cell transplant – (NDMM post-ASCT)

(Please refer to Appendix A).

2.5MG CAPSULE

02459418 REVLIMID UNK

5MG CAPSULE

02304899 REVLIMID UNK

10MG CAPSULE

02304902 REVLIMID UNK

15MG CAPSULE

02317699 REVLIMID UNK

20MG CAPSULE

02440601 REVLIMID UNK

25MG CAPSULE

02317710 REVLIMID UNK

LENVATINIB

Limited use benefit (prior approval required).

Criteria for initial 4-month coverage:

- Used as monotherapy for treatment of patients with locally recurrent or metastatic, progressive differentiated thyroid cancer (DTC); AND
- DTC is refractory to radioactive iodine treatment; AND
- Have an ECOG performance status of ≤ 2 ; AND

Patient meets the eligibility criteria of the SELECT trial as follows:

- Pathologically confirmed differentiated thyroid cancer (patients with anaplastic or medullary thyroid cancer are not eligible)
- Evidence of iodine-131 refractory disease according to at least one of the following criteria:
 - At least one measurable lesion without iodine uptake on any iodine-131 scan
 - At least one measurable lesion that had progressed according to RECIST criteria within 12 months after iodine-131 therapy despite iodine-131 avidity at the time of treatment
 - Total lifetime radioactive iodine dose greater than 600 mCi (millicurie)
- Radiologic evidence of progression within the previous 13 months
- No prior therapy with a tyrosine kinase inhibitor or have received one prior treatment regimen with a tyrosine kinase inhibitor

Criteria for renewal every 4 months:

There is no objective evidence of disease progression.

10MG CAPSULE

02450321 LENVIMA EIS

14MG CAPSULE

02450313 LENVIMA EIS

10:00.00 ANTINEOPLASTIC AGENTS

LENVATINIB

Limited use benefit (prior approval required).

Criteria for initial 4-month coverage:

- Used as monotherapy for treatment of patients with locally recurrent or metastatic, progressive differentiated thyroid cancer (DTC); AND
- DTC is refractory to radioactive iodine treatment; AND
- Have an ECOG performance status of ≤ 2 ; AND

Patient meets the eligibility criteria of the SELECT trial as follows:

- Pathologically confirmed differentiated thyroid cancer (patients with anaplastic or medullary thyroid cancer are not eligible)
- Evidence of iodine-131 refractory disease according to at least one of the following criteria:
 - At least one measurable lesion without iodine uptake on any iodine-131 scan
 - At least one measurable lesion that had progressed according to RECIST criteria within 12 months after iodine-131 therapy despite iodine-131 avidity at the time of treatment
 - Total lifetime radioactive iodine dose greater than 600 mCi (millicurie)
- Radiologic evidence of progression within the previous 13 months
- No prior therapy with a tyrosine kinase inhibitor or have received one prior treatment regimen with a tyrosine kinase inhibitor

Criteria for renewal every 4 months:

There is no objective evidence of disease progression.

20MG CAPSULE

02450305 LENVIMA EIS

24MG CAPSULE

02450291 LENVIMA EIS

LETROZOLE

ST **2.5MG TABLET**

02338459	ACH-LETROZOLE	ACC
02358514	APO-LETROZOLE	APX
02392496	BIO-LETROZOLE	BMI
02231384	FEMARA	NVR
02373009	JAMP-LETROZOLE	JMP
02402025	LETROZOLE	PDL
02373424	MAR-LETROZOLE	MAR
02322315	MED-LETROZOLE	GMP
02421585	NAT-LETROZOLE	NPH
02309114	PMS-LETROZOLE	PMS
02372282	RAN-LETROZOLE	RBY
02398656	RIVA-LETROZOLE	RIV
02344815	SANDOZ LETROZOLE	SDZ
02343657	TEVA-LETROZOLE	TEV
02378213	ZINDA-LETROZOLE	UNK

LEUPROLIDE ACETATE

10.5MG/VIAL POWDER FOR SUSPENSION

02248239 ELIGARD SAC

22.5MG/VIAL POWDER FOR SUSPENSION

02248240 ELIGARD SAC

30MG/VIAL POWDER FOR SUSPENSION

02248999 ELIGARD SAC

10:00.00 ANTINEOPLASTIC AGENTS

LEUPROLIDE ACETATE

45MG/VIAL POWDER FOR SUSPENSION

02268892 ELIGARD SAC

LOMUSTINE

10MG CAPSULE

00360430 CEENU BMS

40MG CAPSULE

00360422 CEENU BMS

100MG CAPSULE

00360414 CEENU BMS

MEGESTROL ACETATE

40MG TABLET

02195917 MEGESTROL AAP

160MG TABLET

02195925 MEGESTROL AAP

MELPHALAN

2MG TABLET

00004715 ALKERAN ASP

MERCAPTOPURINE

50MG TABLET

02415275 MERCAPTOPURINE RAX

00004723 PURINETHOL TEV

METHOTREXATE SODIUM

7.5MG SOLUTION

02320029 METOJECT UNK

02454823 METOJECT SUBCUTANEOUS UNK

10MG SOLUTION

02320037 METOJECT UNK

02454831 METOJECT SUBCUTANEOUS UNK

10MG/0.4ML SOLUTION

02422174 METHOTREXATE PMS

10MG/ML SOLUTION

02182947 METHOTREXATE PFI

12.5MG SOLUTION

02454750 METOJECT SUBCUTANEOUS UNK

15MG SOLUTION

02320045 METOJECT UNK

02454858 METOJECT SUBCUTANEOUS UNK

15MG/0.6ML SOLUTION

02422182 METHOTREXATE PMS

17.5MG SOLUTION

02454769 METOJECT SUBCUTANEOUS UNK

20MG SOLUTION

02454866 METOJECT SUBCUTANEOUS UNK

20MG/0.8ML SOLUTION

02422190 METHOTREXATE PMS

22.5MG SOLUTION

02454777 METOJECT SUBCUTANEOUS UNK

25MG SOLUTION

02454874 METOJECT SUBCUTANEOUS UNK

25MG/ML SOLUTION

02419173 JAMP-METHOTREXATE JMP

02099705 METHOTREXATE TEV

10:00.00 ANTINEOPLASTIC AGENTS

METHOTREXATE SODIUM

25MG/ML SOLUTION

02182777 METHOTREXATE PFI

02182955 METHOTREXATE PFI

02398427 METHOTREXATE SDZ

02417626 METHOTREXATE MYL

02422166 METHOTREXATE PMS

02422204 METHOTREXATE PMS

2.5MG TABLET

02182963 APO-METHOTREXATE PFI

02170698 PMS-METHOTREXATE PMS

10MG TABLET

02182750 METHOTREXATE PFI

MIDOSTAURIN

Limited use benefit (prior approval required).

Criteria for 12-month coverage:

- Patient has newly diagnosed FMS-like tyrosine kinase 3 (FLT3)-mutated acute myeloid leukemia (AML); AND
- Patient's FLT3-mutation status has been confirmed; AND
- Midostaurin is being used in combination with standard cytarabine and daunorubicin (or idarubicin) induction and cytarabine consolidation chemotherapy; AND
- Patient has an ECOG performance status of 0 to 2.

25MG CAPSULE

02466236 RYDAPT NVR

MITOTANE

500MG TABLET

00463221 LYSODREN LAP

NILOTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- Patients has newly diagnosed Philadelphia chromosome positive (Ph+) chronic myelogenous leukemia in chronic phase; OR
- Patient has chronic phase or accelerated phase Philadelphia chromosome positive (Ph+) chronic myelogenous leukemia; AND
- Patient has disease progression/resistance to imatinib; OR
- Documented intolerance to a prior oral TKI (imatinib, dasatinib or bosutinib).

Criteria for renewal every 12 months:

- Confirmation from the clinician that the patient has experienced hematologic and/or cytogenetic response and is expected to continue to do so AND has not developed unacceptable toxicities.

150MG CAPSULE

02368250 TASIGNA NVR

200MG CAPSULE

02315874 TASIGNA NVR

NILUTAMIDE

50MG TABLET

02221861 ANANDRON SAC

10:00.00 ANTINEOPLASTIC AGENTS

OLAPARIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- Maintenance treatment of adult patients with high grade serous epithelial ovarian fallopian tube cancer; OR
- Primary peritoneal cancer;
- AND
- Platinum-sensitive disease; AND
- Relapsed BRCA-mutated disease (germline or somatic as detected by approved testing)
- Have completed at least two previous lines of platinum-based chemotherapy; AND
- Radiologic response (complete or partial response) to their most recent platinum-based chemotherapy regimen as per the SOLO-2 trial; AND
- Patient has an ECOG performance status of 0 to 2;
- AND
- Olaparib is used as monotherapy

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

50MG CAPSULE

02454408 LYNPARZA AZC

100MG TABLET

02475200 LYNPARZA AZC

150MG TABLET

02475219 LYNPARZA AZC

OSIMERTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- Patient with locally advanced or metastatic non-small cell lung cancer (NSCLC) who has progressed on epidermal growth factor receptor (EGFR) tyrosine kinase inhibitor therapy;
- AND
- Patient is EGFR T790M mutation- positive; AND
- Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

40MG TABLET

02456214 TAGRISSO AZC

80MG TABLET

02456222 TAGRISSO AZC

10:00.00 ANTINEOPLASTIC AGENTS

PALBOCICLIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- For the treatment of post-menopausal women with estrogen receptor (ER)-positive, human epidermal growth factor receptor 2 (HER2)-negative advanced breast cancer; AND
- The patient has not received any prior treatment for metastatic disease (first-line treatment); AND
 - Palbociclib will be used in combination with an aromatase inhibitor; AND
 - Patient has an ECOG performance status of 0 to 2; AND
 - Patient is not resistant to prior (neo)adjuvant aromatase inhibitor therapy; AND
 - Patient does not have active or uncontrolled metastases to the central nervous system.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

75MG CAPSULE

02453150 IBRANCE PFI

100MG CAPSULE

02453169 IBRANCE PFI

125MG CAPSULE

02453177 IBRANCE PFI

PAZOPANIB

Limited use benefit (prior approval required).

Initial coverage criteria (12 months)

- For the first-line treatment of patients with advanced or metastatic clear cell renal carcinoma; AND
- Patient has an ECOG performance status of 0 to 2.

Renewal coverage criteria (12 months)

There is no objective evidence of disease progression.

200MG TABLET

02352303 VOTRIENT NVR

POMALIDOMIDE

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- For the treatment of relapsed or refractory multiple myeloma who meet all of the following criteria:
- Used in combination with dexamethasone; AND
 - Patient has relapsed or is refractory to at least two treatment regimens, including both bortezomib and lenalidomide; AND
 - Patient has demonstrated disease progression on the last regimen.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression or development of unacceptable toxicity to pomalidomide requiring discontinuation of therapy.

1MG CAPSULE

02419580 POMALYST UNK

2MG CAPSULE

02419599 POMALYST UNK

3MG CAPSULE

02419602 POMALYST UNK

4MG CAPSULE

02419610 POMALYST UNK

10:00.00 ANTINEOPLASTIC AGENTS

PONATINIB HYDROCHLORIDE

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

- For the treatment of patients who have confirmed T315i mutation positive disease, independent of previous TKI therapy; OR
- Treatment of last resort for patients with intolerances or contraindications to imatinib and all other second generation TKI's (dasatinib, nilotinib, bosutinib); OR
- For the treatment of patients with chronic phase chronic myeloid leukemia (CML) who have resistance/disease progression after at least two prior lines of TKI therapy where lclusig would be available as third-line TKI option; OR
- For the treatment of patients with accelerated phase or blast phase CML or Ph+ ALL who have resistance or disease progression after at least one second generation TKI therapy; AND
- An ECOG performance status of 0 to 2.

Note: Second generation TKI's (dasatinib, nilotinib, bosutinib) are not covered as options after ponatinib.

Criteria for renewal every 6 months:

- There is no objective evidence of disease progression.

- 15MG TABLET**
02437333 ICLUSIG ARI
- 45MG TABLET**
02437341 ICLUSIG ARI

PROCARBAZINE HYDROCHLORIDE

- 50MG CAPSULE**
00012750 MATULANE UNK

REGORAFENIB

Limited use benefit (prior approval required).

1. For the treatment of Gastrointestinal Stromal Tumors (GIST)
Criteria for initial six-month coverage:

- For patients with gastrointestinal stromal tumors (GIST) who have failed or are unable to tolerate imatinib and sunitinib therapy; AND
- Patient has an ECOG performance status of 0 or 1;

Note: Regorafenib will not be funded concomitantly with imatinib or sunitinib.

Criteria for assessment every 12 months:

- There is no objective evidence of disease progression.

2. For the treatment of Hepatocellular Carcinoma (HCC)
Criteria for initial six-month coverage:

- Patient diagnosed with unresectable HCC; AND
- Patient has been previously treated with sorafenib; AND
- Patient was able to tolerate sorafenib as defined in the RESORCE trial criteria (≥400mg/day for ≥20 days of the last 28 days of treatment); AND
- Patient has a Child-Pugh class status of A; AND
- Patient has an ECOG* performance status of 0 to 1

Criteria for assessment every 12 months:

- There is no objective evidence of disease progression.

- 40MG TABLET**
02403390 STIVARGA BAY

10:00.00 ANTINEOPLASTIC AGENTS

RITUXIMAB

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Granulomatosis polyangiitis according to established criteria.
- Microscopic polyangiitis according to established criteria.

(Please refer to Appendix A).

- 10MG/ML SOLUTION**
02241927 RITUXAN HLR

10:00.00 ANTINEOPLASTIC AGENTS

RUXOLITINIB

Limited use benefit (prior approval required).

1. For the treatment of Myelofibrosis:

Criteria for initial 6-month coverage:

- Intermediate to high risk symptomatic myelofibrosis as assessed using the Dynamic International Prognostic Scoring System (DIPSS) Plus; OR
- Patient has symptomatic splenomegaly;
- AND
- Patient has an ECOG performance status of 0 to 3; AND
- Patient previously untreated OR refractory to other treatment.

Criteria for renewal every 12 months:

- Reduction in spleen size; OR
- Improvement in disease symptoms.

2. For the treatment of patients with polycythemia vera:

Criteria for initial 6-month coverage:

Disease is resistant to hydroxyurea (HU) according to the modified European LeukemiaNet Criteria defined as below: After 3 months of at least 2g/day of HU or at the maximally tolerated HU dose, patient showed:

- Need for phlebotomy to keep hematocrit < 45%; OR
- Uncontrolled myeloproliferation (platelet > 400x109/L and WBC > 10x109/L); OR
- Failure to reduce massive splenomegaly > 50% as measured by palpation.

OR

Patient is intolerant to HU according to the modified European LeukemiaNet Criteria defined below:

After any dose of HU, patient showed:

- Absolute neutrophil count < 1.0 x 109/L , or platelet < 100x109/L or hemoglobin < 100 g/L at the lowest dose of HU required to achieve a response (response defined as hematocrit < 45% without phlebotomy, and/or all of the following : platelet ≤ 400x109/L , WBC ≤ 10 x 109/L , and non-palpable spleen); OR
- Presence of leg ulcers or other unacceptable HU-related non-hematological toxicities (such as mucocutaneous manifestations, gastrointestinal symptoms, pneumonitis or fever, defined as Common Terminology Criteria for Adverse Events (CTCAE) version 3.0 grade 3 or 4, or more than one week of CTCAE version 3.0 grade 2, or permanent discontinuation of HU, or interruption of HU until toxicity resolved, or hospitalization due to HU toxicity).

AND

- Patient has an ECOG performance status of 0 to 3.

Criteria for renewal every 12 months:

- Reduction in spleen size; OR
- Improvement in disease symptoms.

5MG TABLET

02388006 JAKAVI NVR

10MG TABLET

02434814 JAKAVI NVR

15MG TABLET

02388014 JAKAVI NVR

20MG TABLET

02388022 JAKAVI NVR

10:00.00 ANTINEOPLASTIC AGENTS

SUNITINIB MALATE

Limited use benefit (Prior approval required).

Criteria for initial 6-month coverage:

- For patients with histologically proven unresectable or recurrent/metastatic GIST who have failed or are unable to tolerate imatinib therapy.

Sunitinib will not be funded concomitantly with imatinib. OR

Criteria for initial 12-month coverage:

- Documented, progressive, unresectable, well or moderately differentiated, locally advanced or metastatic pancreatic neuroendocrine tumors; AND
- Patient has an ECOG performance status of 0 to 2.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

12.5MG CAPSULE

02280795 SUTENT PFI

25MG CAPSULE

02280809 SUTENT PFI

50MG CAPSULE

02280817 SUTENT PFI

TAMOXIFEN CITRATE

10MG TABLET

00812404 APO-TAMOX APX

00851965 TEVA-TAMOXIFEN TEV

20MG TABLET

00812390 APO-TAMOX APX

02048485 NOLVADEX-D AZC

00851973 TEVA-TAMOXIFEN TEV

TEMOZOLOMIDE

5MG CAPSULE

02441160 ACT TEMOZOLOMIDE ACG

02443473 TARO-TEMOZOLOMIDE TAR

02241093 TEMODAL FRS

20MG CAPSULE

02395274 ACT TEMOZOLOMIDE ACG

02443481 TARO-TEMOZOLOMIDE TAR

02241094 TEMODAL FRS

100MG CAPSULE

02395282 ACT TEMOZOLOMIDE ACG

02443511 TARO-TEMOZOLOMIDE TAR

02241095 TEMODAL FRS

140MG CAPSULE

02395290 ACT TEMOZOLOMIDE ACG

02413116 APO-TEMOZOLOMIDE APX

02443538 TARO-TEMOZOLOMIDE TAR

02312794 TEMODAL FRS

250MG CAPSULE

02395312 ACT TEMOZOLOMIDE ACG

02443554 TARO-TEMOZOLOMIDE TAR

02241096 TEMODAL FRS

THIOGUANINE

40MG TABLET

00282081 LANVIS ASP

10:00.00 ANTINEOPLASTIC AGENTS

TRAMETINIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:
 For the first-line treatment of patients with metastatic or unresectable melanoma as monotherapy; OR
 For the first-line treatment of patients with metastatic or unresectable melanoma in combination with dabrafenib(Tafinlar)

AND for patients who meet the following criteria:
 • Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND
 • Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND
 • Patient has an ECOG performance status of 0 to 1; AND
 • Patient is previously untreated.

Criteria for renewal every 6 months:
 There is no objective evidence of disease progression.

0.5MG TABLET
 02409623 MEKINIST NVR

2MG TABLET
 02409658 MEKINIST NVR

TRETINOIN

10MG CAPSULE
 02145839 VESANOID CHE

TRIPTORELIN PAMOATE

3.75MG/VIAL POWDER FOR SUSPENSION
 02240000 TRELSTAR ALL

11.25MG/VIAL POWDER FOR SUSPENSION
 02243856 TRELSTAR ALL

22.5MG POWDER FOR SUSPENSION
 02412322 TRELSTAR ALL

VANDETANIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:
 For patients with symptomatic and/or progressive medullary thyroid cancer with unresectable locally advanced or metastatic disease; AND
 An ECOG* performance status of 0 to 2.

Criteria for renewal every 12 months:
 There is no objective evidence of disease progression.

100MG TABLET
 02378582 CAPRELSA SAC

300MG TABLET
 02378590 CAPRELSA SAC

10:00.00 ANTINEOPLASTIC AGENTS

VEMURAFENIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:
 For the first-line treatment of patients with metastatic or unresectable melanoma as monotherapy; OR
 For the first-line treatment of patients with metastatic or unresectable melanoma in combination with cobimetinib (Cotellic).

AND for patients who meet the following criteria:
 • Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND
 • Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND
 • Patient has an ECOG performance status of 0 to 1.

Criteria for renewal every 6 months:
 There is no objective evidence of disease progression.

ST **240MG TABLET**
 02380242 ZELBORAF HLR

VENETOCLAX

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:
 For the treatment of chronic lymphocytic leukemia (CLL) who meet all of the following criteria:
 • Venclexta will be used as monotherapy; AND
 • Patient has received at least one prior therapy; AND
 • Patient has failed a B-cell receptor inhibitor (BCRi) or is intolerant to prior ibrutinib therapy; AND
 • Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 12 months:
 There is no objective evidence of disease progression or unacceptable toxicity.
 Coverage is for a maximum duration of two years.

10MG TABLET
 02458039 VENCLEXTA ABV

50MG TABLET
 02458047 VENCLEXTA ABV

100MG TABLET
 02458055 VENCLEXTA ABV
 02458063 VENCLEXTA ABV

12:00 AUTONOMIC DRUGS

12:04.00 PARASYMPATHOMIMETIC AGENTS

BETHANECHOL CHLORIDE

10MG TABLET

01947958 DUVOID PAL

25MG TABLET

01947931 DUVOID PAL

50MG TABLET

01947923 DUVOID PAL

DONEPEZIL HYDROCHLORIDE

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:
 • Diagnosis of mild to moderate Alzheimer's disease; AND
 • Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 • Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 • Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
 Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:
 • Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
 • Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST **5MG TABLET**

02419866	ACCEL-DONEPEZIL	ACP
02362260	APO-DONEPEZIL	APX
02232043	ARICEPT	PFI
02400561	AURO-DONEPEZIL	AUR
02412853	BIO-DONEPEZIL	BMI
02402645	DONEPEZIL	ACC
02416417	DONEPEZIL	PDL
02420597	DONEPEZIL	SIV
02426846	DONEPEZIL	SAN
02475278	DONEPEZIL	RIV
02416948	JAMP-DONEPEZIL	JMP
02402092	MAR-DONEPEZIL	MAR
02467453	M-DONEPEZIL	MAN
02439557	NAT-DONEPEZIL	NPH
02322331	PMS-DONEPEZIL	PMS
02381508	RAN-DONEPEZIL	RBV
02412918	RIVA-DONEPEZIL	RIV
02328666	SANDOZ DONEPEZIL	SDZ
02428482	SEPTA DONEPEZIL	SPT
02340607	TEVA-DONEPEZIL	TEV

ST **10MG TABLET**

02419874	ACCEL-DONEPEZIL	ACP
02362279	APO-DONEPEZIL	APX
02232044	ARICEPT	PFI
02400588	AURO-DONEPEZIL	AUR
02412861	BIO-DONEPEZIL	BMI
02402653	DONEPEZIL	ACC
02416425	DONEPEZIL	PDL

12:04.00 PARASYMPATHOMIMETIC AGENTS

DONEPEZIL HYDROCHLORIDE

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:
 • Diagnosis of mild to moderate Alzheimer's disease; AND
 • Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 • Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 • Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
 Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:
 • Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
 • Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST **10MG TABLET**

02420600	DONEPEZIL	SIV
02426854	DONEPEZIL	SAN
02475286	DONEPEZIL	RIV
02416956	JAMP-DONEPEZIL	JMP
02402106	MAR-DONEPEZIL	MAR
02467461	M-DONEPEZIL	MAN
02439565	NAT-DONEPEZIL	NPH
02322358	PMS-DONEPEZIL	PMS
02381516	RAN-DONEPEZIL	RBV
02412934	RIVA-DONEPEZIL	RIV
02328682	SANDOZ DONEPEZIL	SDZ
02428490	SEPTA DONEPEZIL	SPT
02340615	TEVA-DONEPEZIL	TEV

GALANTAMINE HYDROBROMIDE

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:
 • Diagnosis of mild to moderate Alzheimer's disease; AND
 • Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 • Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 • Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
 Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:
 • Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
 • Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST **8MG CAPSULE (EXTENDED RELEASE)**

02425157	AURO-GALANTAMINE ER	AUR
02443015	GALANTAMINE	SAN
02416573	GALANTAMINE ER	PDL
02420821	MAR-GALANTAMINE ER	MAR
02339439	MYLAN-GALANTAMINE ER	MYL
02316943	PAT-GALANTAMINE ER	KLA

12:04.00 PARASYMPATHOMIMETIC AGENTS

GALANTAMINE HYDROBROMIDE

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:
 • Diagnosis of mild to moderate Alzheimer's disease; AND
 • Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 • Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 • Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
 Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:
 • Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
 • Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 8MG CAPSULE (EXTENDED RELEASE)			
02398370	PMS-GALANTAMINE ER		PMS
ST 16MG CAPSULE (EXTENDED RELEASE)			
02425165	AURO-GALANTAMINE ER		AUR
02443023	GALANTAMINE		SAN
02416581	GALANTAMINE ER		PDL
02420848	MAR-GALANTAMINE ER		MAR
02339447	MYLAN-GALANTAMINE ER		MYL
02316951	PAT-GALANTAMINE ER		KLA
02398389	PMS-GALANTAMINE ER		PMS
ST 24MG CAPSULE (EXTENDED RELEASE)			
02425173	AURO-GALANTAMINE ER		AUR
02443031	GALANTAMINE		SAN
02416603	GALANTAMINE ER		PDL
02420856	MAR-GALANTAMINE ER		MAR
02339455	MYLAN-GALANTAMINE ER		MYL
02316978	PAT-GALANTAMINE ER		KLA
02398397	PMS-GALANTAMINE ER		PMS

NEOSTIGMINE BROMIDE

ST 15MG TABLET			
00869945	PROSTIGMIN		VAE

PILOCARPINE HYDROCHLORIDE

ST 5MG TABLET			
02402483	PILOCARPINE HYDROCHLORIDE		RAX
02216345	SALAGEN		PFI

PYRIDOSTIGMINE BROMIDE

ST 60MG TABLET			
00869961	MESTINON		VAE
ST 180MG TABLET (EXTENDED RELEASE)			
00869953	MESTINON-SR		VAE

12:04.00 PARASYMPATHOMIMETIC AGENTS

RIVASTIGMINE HYDROGEN TARTRATE

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:
 • Diagnosis of mild to moderate Alzheimer's disease; AND
 • Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 • Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 • Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
 Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:
 • Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
 • Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 1.5MG CAPSULE			
02336715	APO-RIVASTIGMINE		APX
02242115	EXELON		NVR
02401614	MED-RIVASTIGMINE		GMP
02306034	PMS-RIVASTIGMINE		PMS
02416999	RIVASTIGMINE		PDL
02324563	SANDOZ RIVASTIGMINE		SDZ
ST 3MG CAPSULE			
02336723	APO-RIVASTIGMINE		APX
02242116	EXELON		NVR
02401622	MED-RIVASTIGMINE		GMP
02306042	PMS-RIVASTIGMINE		PMS
02417006	RIVASTIGMINE		PDL
02324571	SANDOZ RIVASTIGMINE		SDZ
ST 4.5MG CAPSULE			
02336731	APO-RIVASTIGMINE		APX
02242117	EXELON		NVR
02401630	MED-RIVASTIGMINE		GMP
02306050	PMS-RIVASTIGMINE		PMS
02417014	RIVASTIGMINE		PDL
02324598	SANDOZ RIVASTIGMINE		SDZ
ST 6MG CAPSULE			
02336758	APO-RIVASTIGMINE		APX
02242118	EXELON		NVR
02401649	MED-RIVASTIGMINE		GMP
02306069	PMS-RIVASTIGMINE		PMS
02417022	RIVASTIGMINE		PDL
02324601	SANDOZ RIVASTIGMINE		SDZ
ST 2MG/ML SOLUTION			
02245240	EXELON		NVR
12:08.08 ANTIMUSCARINICS / ANTISPASMODICS			
ACLIDINIUM BROMIDE			
400MCG POWDER			
02409720	TUDORZA GENUAIR		AZC

**12:08.08 ANTIMUSCARINICS /
ANTISPASMODICS**

GLYCOPYRRONIUM BROMIDE

50MCG CAPSULE

02394936 SEEBRI BREEZHALER NVR

HYOSCINE BUTYLBROMIDE

ST **10MG TABLET**

00363812 BUSCOPAN SAC

INDACATEROL MALEATE, GLYCOPYRRONIUM BROMIDE

Open benefit with therapeutic notes (prior approval is not required).

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry or standardized scale*; AND
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA)

*As defined by the Canadian Thoracic Society COPD classification. Moderate: shortness of breath from COPD causing the patient to stop after walking approximately 100 meters (or after a few minutes) on the level. Severe: shortness of breath from COPD resulting in the patient being too breathless to leave the house or breathless after undressing, or the presence of chronic respiratory failure or clinical signs of right heart failure.

110MCG & 50MCG CAPSULE

02418282 ULTIBRO BREEZHALER NVR

IPRATROPIUM BROMIDE

20MCG/INHALATION AEROSOL

02247686 ATROVENT HFA BOE

0.03% NASAL SPRAY

02240508 DOM-IPRATROPIUM DPC

02239627 PMS-IPRATROPIUM PMS

21MCG NASAL SPRAY

02246083 IPRAVENT AAP

42MCG NASAL SPRAY

02246084 IPRAVENT AAP

125MCG/ML SOLUTION

02231135 PMS-IPRATROPIUM PMS

250MCG/ML SOLUTION

02126222 APO-IPRAVENT APX

02231136 PMS-IPRATROPIUM PMS

02231244 PMS-IPRATROPIUM PMS

02231245 PMS-IPRATROPIUM PMS

99001446 RATIO-IPRATROPIUM RPH

02216221 TEVA-IPRATROPIUM STERINEBS TEV

IPRATROPIUM BROMIDE, SALBUTAMOL SULFATE

0.2MG & 1MG/ML SOLUTION

02231675 COMBIVENT BOE

02243789 RATIO-IPRA SAL TEV

02272695 TEVA-COMBO STERINEBS TEV

100MCG & 20MCG SOLUTION

02419106 COMBIVENT RESPIMAT BOE

**12:08.08 ANTIMUSCARINICS /
ANTISPASMODICS**

TIOTROPIUM BROMIDE MONOHYDRATE

18MCG CAPSULE

02246793 SPIRIVA BOE

2.5MCG SOLUTION

02435381 SPIRIVA RESPIMAT BOE

TRIMEBUTINE MALEATE

Limited use benefit (prior approval required).

For the treatment and relief of symptoms associated with functional bowel disorders including Irritable Bowel Syndrome (IBS), spastic colon, spastic colitis and mucous colitis; OR In postoperative paralytic ileus in order to accelerate the resumption of the intestinal transit following abdominal surgery.

100MG TABLET

02349027 AA-TRIMEBUTINE AAP

02245663 TRIMEBUTINE AAP

200MG TABLET

02349035 AA-TRIMEBUTINE AAP

02245664 TRIMEBUTINE AAP

UMECLIDIUM BROMIDE

62.5MCG POWDER

02423596 INCRUSE ELLIPTA GSK

UMECLIDIUM BROMIDE, VILANTEROL TRIFENATATE

Open benefit with therapeutic notes (prior approval is not required).

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry or standardized scale*; AND
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA)

*As defined by the Canadian Thoracic Society COPD classification. Moderate: shortness of breath from COPD causing the patient to stop after walking approximately 100 meters (or after a few minutes) on the level. Severe: shortness of breath from COPD resulting in the patient being too breathless to leave the house or breathless after undressing, or the presence of chronic respiratory failure or clinical signs of right heart failure.

62.5MCG/25MCG POWDER

02418401 ANORO ELLIPTA GSK

12:12.04 ALPHA ADRENERGIC AGONISTS

MIDODRINE HYDROCHLORIDE

2.5MG TABLET

02278677 APO-MIDODRINE APX

02473984 MAR-MIDODRINE MAR

5MG TABLET

02278685 APO-MIDODRINE APX

02473992 MAR-MIDODRINE MAR

12:12.08 BETA ADRENERGIC AGONISTS

ACLIDINIUM BROMIDE, FORMOTEROL FUMARATE DIHYDRATE

Open benefit with therapeutic notes (prior approval is not required).

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry or standardized scale*; AND
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA)

*As defined by the Canadian Thoracic Society COPD classification. Moderate: shortness of breath from COPD causing the patient to stop after walking approximately 100 meters (or after a few minutes) on the level. Severe: shortness of breath from COPD resulting in the patient being too breathless to leave the house or breathless after undressing, or the presence of chronic respiratory failure or clinical signs of right heart failure.

400MCG & 12MCG POWDER

02439530 DUAKLIR GENUAIR AZC

FLUTICASONE FUROATE, VILANTEROL TRIFENATATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

OR

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry; OR
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA).

100MCG & 25MCG POWDER

02408872 BREO ELLIPTA GSK

FLUTICASONE FUROATE, VILANTEROL TRIFENATATE (ASTHMA)

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

200MCG & 25MCG POWDER

02444186 BREO ELLIPTA GSK

12:12.08 BETA ADRENERGIC AGONISTS

FORMOTEROL FUMARATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of a rapid-onset, short-duration bronchodilator.

OR

For the treatment of Chronic Obstructive Pulmonary Disease (COPD) in patients not adequately controlled with either ipratropium, tiotropium or a short acting beta-agonist.

12MCG/CAPSULE CAPSULE

02230898 FORADIL NVR

FORMOTEROL FUMARATE DIHYDRATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of rapid onset, short duration bronchodilator.

6MCG/DOSE POWDER

02237225 OXEZE TURBUHALER AZC

12MCG/DOSE POWDER

02237224 OXEZE TURBUHALER AZC

FORMOTEROL FUMARATE DIHYDRATE, BUDESONIDE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

OR

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry; OR
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA).

6MCG & 100MCG/INHALATION POWDER

02245385 SYMBICORT 100 TURBUHALER AZC

6MCG & 200MCG/INHALATION POWDER

02245386 SYMBICORT 200 TURBUHALER AZC

FORMOTEROL FUMARATE DIHYDRATE, MOMETASONE FUROATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of rapid onset, short duration bronchodilator.

5MCG & 100MCG/INHALATION AEROSOL

02361752 ZENHALE FRS

5MCG & 200MCG/INHALATION AEROSOL

02361760 ZENHALE FRS

5MCG & 50MCG/INHALATION AEROSOL

02361744 ZENHALE FRS

12:12.08 BETA ADRENERGIC AGONISTS

INDACATEROL MALEATE

Limited use benefit (prior approval required).

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who:

- are not adequately controlled with either ipratropium, tiotropium or a short acting beta-agonist; OR
- have moderate to severe COPD, as defined by spirometry.

75MCG CAPSULE

02376938 ONBREZ BREEZHALER NVR

OLODATEROL HYDROCHLORIDE, TIOTROPIUM BROMIDE MONOHYDRATE

2.5MCG & 2.5MCG SOLUTION

02441888 INSPIOLTO RESPIMAT BOE

ORCIPRENALINE SULFATE

2MG/ML SYRUP

02236783 ORCIPRENALINE AAP

SALBUTAMOL SULFATE

100MCG/INHALATION AEROSOL

02232570 AIROMIR VAE

02245669 APO-SALVENT CFC FREE APX

02419858 SALBUTAMOL HFA SAN

02326450 TEVA-SALBUTAMOL HFA TEV

02241497 VENTOLIN HFA GSK

200MCG POWDER

02243115 VENTOLIN DISKUS GSK

0.5MG/ML SOLUTION

02208245 PMS-SALBUTAMOL PMS

02239365 RATIO-SALBUTAMOL TEV

1MG/ML SOLUTION

02216949 DOM-SALBUTAMOL DPC

02208229 PMS-SALBUTAMOL PMS

01926934 TEVA-SALBUTAMOL TEV

02213419 VENTOLIN P.F GSK

2MG/ML SOLUTION

02208237 PMS-SALBUTAMOL PMS

02173360 TEVA-SALBUTAMOL TEV

02213427 VENTOLIN P.F GSK

5MG/ML SOLUTION

02139324 DOM-SALBUTAMOL DPC

02213486 VENTOLIN RESPIRATOR GSK

SALMETEROL XINAFOATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of a rapid-onset, short-duration bronchodilator.

OR

For the treatment of Chronic Obstructive Pulmonary Disease (COPD) in patients not adequately controlled with either ipratropium, tiotropium or a short acting beta-agonist.

50MCG/INHALATION POWDER

02231129 SEREVENT DISKUS GSK

12:12.08 BETA ADRENERGIC AGONISTS

SALMETEROL XINAFOATE, FLUTICASONE PROPIONATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

OR

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry; OR
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA).

25MCG & 125MCG AEROSOL

02245126 ADVAIR 125 GSK

25MCG & 250MCG AEROSOL

02245127 ADVAIR 250 GSK

50MCG & 100MCG POWDER

02240835 ADVAIR 100 DISKUS GSK

50MCG & 250MCG POWDER

02240836 ADVAIR 250 DISKUS GSK

50MCG & 500MCG POWDER

02240837 ADVAIR 500 DISKUS GSK

TERBUTALINE SULFATE

500MCG/INHALATION POWDER

00786616 BRICANYL TURBUHALER AZC

12:12.12 ALPHA AND BETA ADRENERGIC AGONISTS

EPINEPHRINE

0.15MG SOLUTION

02382059 ALLERJECT KAL

0.3MG SOLUTION

02382067 ALLERJECT KAL

0.5MG/ML SOLUTION

00578657 EPIPEN JR MYL

1MG/ML SOLUTION

00155357 ADRENALIN ERF

00721891 EPINEPHRINE PFI

00509558 EPIPEN MYL

12:16.00 SYMPATHOLYTIC AGENTS

DIHYDROERGOTAMINE MESYLATE

1MG/ML LIQUID

00027243 DIHYDROERGOTAMINE RAX

4MG/ML LIQUID

02228947 MIGRANAL RAX

12:16.04 ALPHA-ADRENERGIC BLOCKING AGENTS

ALFUZOSIN HYDROCHLORIDE

ST **10MG TABLET (EXTENDED RELEASE)**

02447576 ALFUZOSIN SIV

02315866 APO-ALFUZOSIN APX

12:16.04 ALPHA-ADRENERGIC BLOCKING AGENTS

ALFUZOSIN HYDROCHLORIDE

ST **10MG TABLET (EXTENDED RELEASE)**

02443201	AURO-ALFUZOSIN	AUR
02304678	SANDOZ ALFUZOSIN	SDZ
02245565	XATRAL	SAC

TAMSULOSIN HYDROCHLORIDE

ST **0.4MG CAPSULE (SUSTAINED RELEASE)**

02294265	RATIO-TAMSULOSIN	TEV
09857334	RATIO-TAMSULOSIN	RPH
02319217	SANDOZ TAMSULOSIN	SDZ
02281392	TEVA-TAMSULOSIN	TEV

ST **0.4MG TABLET (EXTENDED RELEASE)**

02362406	APO-TAMSULOSIN	APX
02270102	FLOMAX	BOE
02340208	SANDOZ TAMSULOSIN	SDZ
02413612	TAMSULOSIN	PDL
02427117	TAMSULOSIN	SAN
02429667	TAMSULOSIN	SIV
02368242	TEVA-TAMSULOSIN	TEV

12:20.04 CENTRALL ACTING SKELETAL MUSCLE RELAXANTS

CYCLOBENZAPRINE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

For relief of muscle spasm associated with acute, painful musculoskeletal conditions. Coverage is limited to 60mg per day for three (3) weeks renewable every two (2) months.

ST **10MG TABLET**

02177145	APO-CYCLOBENZAPRINE	APX
02348853	AURO-CYCLOBENZAPRINE	AUR
02220644	CYCLOBENZAPRINE	PDL
02287064	CYCLOBENZAPRINE	SAN
02424584	CYCLOBENZAPRINE	SIV
02238633	DOM-CYCLOBENZAPRINE	DPC
02357127	JAMP-CYCLOBENZAPRINE	JMP
02212048	PMS-CYCLOBENZAPRINE	PMS
02242079	RIVA-CYCLOBENZAPRINE	RIV
02080052	TEVA-CYCLOBENZAPRINE	TEV

TIZANIDINE HYDROCHLORIDE

Limited use benefit (prior approval required).

For treatment of spasticity in patients with multiple sclerosis, who have failed therapy with or are intolerant to baclofen.

4MG TABLET

02239170	PAL-TIZANIDINE	PAL
02259893	TIZANIDINE	AAP

12:20.08 DIRECT-ACTING SKELETAL MUSCLE RELAXANTS

DANTROLENE SODIUM

25MG CAPSULE

01997602	DANTRIUM	PPH
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12:20.12 GABA-DERIVATIVE SKELETAL MUSCLE RELAXANTS

BACLOFEN

ST **10MG TABLET**

02139332	APO-BACLOFEN	APX
02152584	BACLOFEN	PDL
02287021	BACLOFEN	SAN
02138271	DOM-BACLOFEN	DPC
00455881	LIORESAL	NVR
02088398	MYLAN-BACLOFEN	MYL
02063735	PMS-BACLOFEN	PMS
02242150	RIVA-BACLOFEN	RIV

ST **20MG TABLET**

02139391	APO-BACLOFEN	APX
02152592	BACLOFEN	PDL
02287048	BACLOFEN	SAN
02138298	DOM-BACLOFEN	DPC
00636576	LIORESAL	NVR
02088401	MYLAN-BACLOFEN	MYL
02063743	PMS-BACLOFEN	PMS
02236508	RATIO-BACLOFEN	TEV
02242151	RIVA-BACLOFEN	RIV

PDIN FOR EXTEMPORANEOUS MIXTURE

99503011	BACLOFEN ORAL LIQUID	UNK
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12:92.00 MISCELLANEOUS AUTONOMIC DRUGS

NICOTINE (GUM)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 945 pieces during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine gum or lozenges when one year has elapsed from the day the initial prescription was filled.

ST **2MG GUM**

02091933	NICORETTE GUM	KIM
80015240	RUGBY NICOTINE POLACRILEX GUM	ACG
80000396	THRIVE NICOTINELL GUM	GSK

ST **4MG GUM**

02091941	NICORETTE GUM	KIM
80000118	NICOTINE GUM	PER
80000402	THRIVE NICOTINELL GUM	NVC

NICOTINE (INHALER)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 945 doses during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine gum or lozenges when one year has elapsed from the day the initial prescription was filled.

ST **10MG SPRAY**

02241742	NICORETTE INHALER	KIM
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12:92.00 MISCELLANEOUS AUTONOMIC DRUGS

NICOTINE (LOZENGE)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:
Coverage is limited to 945 pieces during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine gum or lozenges when one year has elapsed from the day the initial prescription was filled.

ST	1MG LOZENGE		
	80007461	THRIVE NICOTINE LOZENGES	NVC
ST	2MG LOZENGE		
	02247347	NICORETTE LOZENGE	KIM
	80007464	THRIVE NICOTINE LOZENGES	NVC
ST	4MG LOZENGE		
	02247348	NICORETTE LOZENGE	KIM

NICOTINE (PATCH)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:
Coverage will be provided for up to the allowable number of patches for one of the following products, during a one-year period. The year starts on the date the first prescription is filled.

NIHB clients are eligible to receive:
- up to 252 nicotine patches of any listed brand in a 12-month period; AND
- ONE course of an as-needed nicotine replacement therapy (NRT) product (i.e. gum, lozenge or inhaler) in a 12-month period; AND
- up to 180 tablets of Zyban in a 12-month period; AND
- up to 165 tablets of Champix in a 12-month period.

Once this quantity has been reached, the client is eligible again for coverage for nicotine patches when one year has elapsed from the day the initial prescription was filled.

ST	2MG GUM		
	80025660	CHU NICOTINE ANTI SMOKING AID	UNK
	94799974	THRIVE GUM (NS)	NVC
ST	1MG LOZENGE		
	80061161	NICHIT	EUR
ST	2MG LOZENGE		
	80059877	NICHIT	EUR
ST	7MG PATCH		
	01943057	HABITROL	NVC
	80051602	NICOTINE TRANSDERMAL	APX
	80044393	TRANSDERMAL NICOTINE	ACG
ST	14MG PATCH		
	01943065	HABITROL	NVC
	80051600	NICOTINE TRANSDERMAL	APX
	80013549	NICOTINE TRANSDERMAL SYSTEM	ADD
	80044392	TRANSDERMAL NICOTINE	ACG
ST	16MG PATCH		
	80014321	NICOTINE TRANSDERMAL SYSTEM	ADD
ST	18MG PATCH		
	02241227	TRANSDERMAL NICOTINE PATCHDAY	NVC

12:92.00 MISCELLANEOUS AUTONOMIC DRUGS

NICOTINE (PATCH)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:
Coverage will be provided for up to the allowable number of patches for one of the following products, during a one-year period. The year starts on the date the first prescription is filled.

NIHB clients are eligible to receive:
- up to 252 nicotine patches of any listed brand in a 12-month period; AND
- ONE course of an as-needed nicotine replacement therapy (NRT) product (i.e. gum, lozenge or inhaler) in a 12-month period; AND
- up to 180 tablets of Zyban in a 12-month period; AND
- up to 165 tablets of Champix in a 12-month period.

Once this quantity has been reached, the client is eligible again for coverage for nicotine patches when one year has elapsed from the day the initial prescription was filled.

ST	21MG PATCH		
	01943073	HABITROL	NVC
	80051603	NICOTINE TRANSDERMAL	APX
	80014250	NICOTINE TRANSDERMAL SYSTEM	ADD
	80044389	TRANSDERMAL NICOTINE	ACG
ST	36MG PATCH		
	02093111	NICODERM	KIM
ST	53MG PATCH		
	02241228	TRANSDERMAL NICOTINE PATCHDAY	NVC
ST	78MG PATCH		
	02093138	NICODERM	KIM
ST	114MG PATCH		
	02093146	NICODERM	KIM

NICOTINE (SPRAY)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:
Coverage is limited to 3450 sprays during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine spray when one year has elapsed from the day the initial prescription was filled.

ST	1MG ORAL SPRAY		
	80038858	NICORETTE QUICKMIST	KIM

VARENICLINE TARTRATE

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage will be limited to 165 tablets during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for varenicline (Champix®) when one year has elapsed from the day the initial prescription was filled.

ST	0.5MG TABLET		
	02419882	APO-VARENICLINE	APX
	02291177	CHAMPIX	PFI

**12:92.00 MISCELLANEOUS AUTONOMIC
DRUGS**

VARENICLINE TARTRATE

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage will be limited to 165 tablets during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for varenicline (Champix®) when one year has elapsed from the day the initial prescription was filled.

ST **0.5MG & 1MG TABLET**

02298309	CHAMPIX STARTER PACK	PFI
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ST **1MG TABLET**

02419890	APO-VARENICLINE	APX
02435675	APO-VARENICLINE	APX
02291185	CHAMPIX	PFI

**20:00 BLOOD FORMATION
COAGULATION AND
THROMBOSIS**

20:04.04 IRON PREPARATIONS

FERROUS FUMARATE

100MG CAPSULE

80061196 MFER FUMARATE MAN

ST 300MG CAPSULE

02237556 EUROFER EUR

00482064 NEO-FER NEB

01923420 PALAFER VAE

ST 20MG SUSPENSION

80029822 JAMP-FERROUS FUMARATE JMP

ST 60MG/ML SUSPENSION

01923439 PALAFER VAE

ST 300MG/5ML SUSPENSION

02246590 FERRATE EUR

ST 100MG TABLET

80024544 JAMP FERROUS FUMARATE JMP

ST 300MG TABLET

00031089 FERROUS FUMARATE WAM

FERROUS GLUCONATE

ST 300MG TABLET

00545031 APO-FERROUS GLUCONATE APX

00031097 FERROUS GLUCONATE JMP

00041157 FERROUS GLUCONATE ADA

02244532 FERROUS GLUCONATE PMT

80000435 FERROUS GLUCONATE NUR

80002426 FERROUS GLUCONATE WNP

80006316 FERROUS GLUCONATE UNK

80009681 WAMPOLE FERROUS GLUCONATE WAM

ST 324MG TABLET

00582727 IRON FERROUS GLUCONATE VTH

FERROUS SULFATE

ST 30MG/ML LIQUID

80008295 JAMP FERROUS SULFATE LIQUID5 JMP

ST 75MG/ML LIQUID

00762954 ENFAMIL FERINSOL MJO

80008309 JAMP FERROUS SULFATE JMP

ST 6MG/ML SOLUTION

00017884 ENFAMIL FERINSOL MJO

02242863 PEDIAFER EUR

ST 15MG/ML SOLUTION

02237385 FERODAN INFANT DROPS ODN

02232202 PEDIAFER EUR

02222574 PMS-FERROUS SULFATE PMS

ST 30MG/ML SOLUTION

00758469 FERODAN ODN

00792675 PMS-FERROUS SULFATE PMS

ST 125MG/ML SOLUTION

00816035 PMS-FERROUS SULFATE PMS

ST 60MG TABLET

80012039 IRON WNP

20:04.04 IRON PREPARATIONS

FERROUS SULFATE

ST 300MG TABLET

02246733 EURO-FERROUS SULFATE EUR

02248699 FERODAN ODN

00346918 FERROUS SULFATE PMT

00782114 FERROUS SULFATE VTH

00031100 FERROUS SULPHATE JMP

80057416 M-SULFATE FERREUX MAN

00586323 PMS-FERROUS SULFATE PMS

IRON

ST 100MG CAPSULE

80024232 JAMP-FER JMP

12.5MG/ML LIQUID

02243333 FERRLECIT SAC

IRON DEXTRAN

50MG/ML LIQUID

02221780 INFUFER SDZ

50MG/ML SOLUTION

02205963 DEXIRON UNK

IRON SUCROSE

20MG/ML SOLUTION

02243716 VENOFER UNK

PDIN FOR EXTEMPORANEOUS MIXTURE

99506015 IRON SUCROSE STERILE INFUSION UNK

POLYSACCHARIDE IRON COMPLEX

Limited use benefit (prior approval not required).

For children 12 years of age or under.

15MG POWDER

80033717 FERAMAX POWDER WATER SOLUBLE POLYSACCHARIDE IRON COMPLEX BSY

20:12.04 ANTICOAGULANTS

ACENOCOUMAROL

ST 1MG TABLET

00010383 SINTROM PAL

ST 4MG TABLET

00010391 SINTROM PAL

20:12.04 ANTICOAGULANTS

APIXABAN

Limited use benefit (prior approval required).

For at-risk patients (CHADS2 score ≥ 1) with non-valvular atrial fibrillation who require apixaban for the prevention of stroke and systemic embolism AND in whom:

- Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
- Anticoagulation with warfarin is contraindicated; OR
- Anticoagulation with warfarin is not possible due to inability to regularly monitor via INR testing (i.e., no access to INR testing services at a laboratory, clinic, pharmacy and at home).

OR

For the treatment of venous thromboembolism: Deep Vein Thrombosis (DVT) or Pulmonary Embolism (PE)

ST **2.5MG TABLET**

02377233 ELIQUIS BMS

ST **5MG TABLET**

02397714 ELIQUIS BMS

DABIGATRAN ETEXILATE MESILATE

Limited use benefit (prior approval required).

For at-risk patients (CHADS2 score ≥ 1) with non-valvular atrial fibrillation who require dabigatran etexilate for the prevention of stroke and systemic embolism AND in whom:

- Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
- Anticoagulation with warfarin is contraindicated; OR
- Anticoagulation with warfarin is not possible due to inability to regularly monitor via INR testing (i.e., no access to INR testing services at a laboratory, clinic, pharmacy and at home).

ST **110MG CAPSULE**

02312441 PRADAXA BOE

ST **150MG CAPSULE**

02358808 PRADAXA BOE

DALTEPARIN SODIUM

2,500IU/0.2ML SOLUTION

02132621 FRAGMIN PFI

3,500IU/0.28ML SOLUTION

02430789 FRAGMIN PFI

5,000IU/0.2ML SOLUTION

02132648 FRAGMIN PFI

7,500IU/0.3ML SOLUTION

02352648 FRAGMIN PFI

10,000IU/0.4ML SOLUTION

02352656 FRAGMIN PFI

10,000IU/ML SOLUTION

02132664 FRAGMIN PFI

12,500IU/0.5ML SOLUTION

02352664 FRAGMIN PFI

15,000IU/0.6ML SOLUTION

02352672 FRAGMIN PFI

18,000IU/0.72ML SOLUTION

02352680 FRAGMIN PFI

25,000IU/ML SOLUTION

02231171 FRAGMIN PFI

20:12.04 ANTICOAGULANTS

EDOxabAN

Limited use benefit (prior approval required).

For at-risk patients (CHADS2 score ≥ 1) with non-valvular atrial fibrillation who require apixaban for the prevention of stroke and systemic embolism AND in whom:

- Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
- Anticoagulation with warfarin is contraindicated; OR
- Anticoagulation with warfarin is not possible due to inability to regularly monitor via INR testing (i.e., no access to INR testing services at a laboratory, clinic, pharmacy and at home).

OR

For the treatment of venous thromboembolism: Deep Vein Thrombosis (DVT) or Pulmonary Embolism (PE)

15MG TABLET

02458640 LIXIANA SEV

30MG TABLET

02458659 LIXIANA SEV

60MG TABLET

02458667 LIXIANA SEV

ENOXAPARIN SODIUM

30MG/0.3ML SOLUTION

02012472 LOVENOX SAC

40MG/0.4ML SOLUTION

02236883 LOVENOX SAC

60MG/0.6ML SOLUTION

02378426 LOVENOX SAC

80MG/0.8ML SOLUTION

02378434 LOVENOX SAC

100MG/1ML SOLUTION

02378442 LOVENOX SAC

150MG/1.0ML SOLUTION

02242692 LOVENOX HP SAC

150MG/ML SOLUTION

02378469 LOVENOX HP SAC

300MG/3ML SOLUTION

02236564 LOVENOX SAC

HEPARIN SODIUM

100U/ML LIQUID

00727520 HEPARIN LEO LEO

1,000U/ML LIQUID

00453811 HEPARIN LEO LEO

1,000 U/ML SOLUTION

02303086 HEPARIN SODIUM (MULTIDOSE VIAL-WITH PRESERVATIVE) SDZ

10,000 U/ML SOLUTION

02303108 HEPARIN SODIUM (MULTIDOSE VIAL-WITH PRESERVATIVE) SDZ

02303094 HEPARIN SODIUM (SINGLE USE VIAL-PRESERVATIVE FREE) SDZ

5000U SOLUTION

02456958 HEPARIN SODIUM UNK

10,000U SOLUTION

02392453 HEPARIN SODIUM FKD

20:12.04 ANTICOAGULANTS

NADROPARIN CALCIUM

9,500IU/ML SOLUTION		
02236913	FRAXIPARINE	ASP
19,000IU/ML SOLUTION		
02240114	FRAXIPARINE FORTE	ASP

RIVAROXABAN

Limited use benefit (prior approval required).

Criteria for rivaroxaban 15 mg, 20mg tablets (Xarelto) for Stroke Prevention in Atrial Fibrillation (SPAF)
 For at-risk patients (CHADS2 score ≥1) with non-valvular atrial fibrillation who require rivaroxaban for the prevention of stroke and systemic embolism AND in whom:
 • Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
 • Anticoagulation with warfarin is contraindicated; OR
 • Anticoagulation is not possible due to inability to regularly monitor via International Normalized Ratio (INR) testing (i.e., no access to INR testing service at a laboratory, clinic, pharmacy, and at home)

Criteria for rivaroxaban 15 mg, 20mg tablets (Xarelto) For the treatment of venous thromboembolism: Deep Vein Thrombosis (DVT) or Pulmonary Embolism (PE).

ST	15MG TABLET		
	02378604	XARELTO	BAY
ST	20MG TABLET		
	02378612	XARELTO	BAY

RIVAROXABAN (10)

Limited use benefit (prior approval not required).

For the prevention of venous thromboembolism following total knee replacement or total hip replacement surgery, for up to 35 days.

ST	10MG TABLET		
	02316986	XARELTO	BAY

TINZAPARIN SODIUM

2,500IU/0.25ML SOLUTION			
	02229755	INNOHEP	LEO
3,500IU/0.35ML SOLUTION			
	02358158	INNOHEP	LEO
4,500IU/0.45ML SOLUTION			
	02358166	INNOHEP	LEO
8,000IU/0.4ML SOLUTION			
	02429462	INNOHEP	LEO
10,000IU/0.5ML SOLUTION			
	02231478	INNOHEP	LEO
10,000IU/ML SOLUTION			
	02167840	INNOHEP	LEO
12,000IU/0.6ML SOLUTION			
	02429470	INNOHEP	LEO
14,000IU/0.7ML SOLUTION			
	02358174	INNOHEP	LEO
16,000IU/0.8ML SOLUTION			
	02429489	INNOHEP	LEO
18,000IU/0.9ML SOLUTION			
	02358182	INNOHEP	LEO

20:12.04 ANTICOAGULANTS

TINZAPARIN SODIUM

20,000IU/ML SOLUTION			
	02229515	INNOHEP	LEO

WARFARIN SODIUM

ST	1MG TABLET		
	02242924	APO-WARFARIN	APX
	01918311	COUMADIN	BMS
	02242680	TARO-WARFARIN	TAR
ST	2MG TABLET		
	02242925	APO-WARFARIN	APX
	01918338	COUMADIN	BMS
	02242681	TARO-WARFARIN	TAR
ST	2.5MG TABLET		
	02242926	APO-WARFARIN	APX
	01918346	COUMADIN	BMS
	02242682	TARO-WARFARIN	TAR
ST	3MG TABLET		
	02245618	APO-WARFARIN	APX
	02240205	COUMADIN	BMS
	02242683	TARO-WARFARIN	TAR
ST	4MG TABLET		
	02242927	APO-WARFARIN	APX
	02007959	COUMADIN	BMS
	02242684	TARO-WARFARIN	TAR
ST	5MG TABLET		
	02242928	APO-WARFARIN	APX
	01918354	COUMADIN	BMS
	02242685	TARO-WARFARIN	TAR
6MG TABLET			
	02240206	COUMADIN	BMS
	02242686	TARO-WARFARIN	TAR
ST	7.5MG TABLET		
	02242697	TARO-WARFARIN	TAR
ST	10MG TABLET		
	02242929	APO-WARFARIN	APX
	01918362	COUMADIN	BMS
	02242687	TARO-WARFARIN	TAR

20:12.14 PLATELET AGGREGATION INHIBITORS

ANAGRELIDE HYDROCHLORIDE

ST	0.5MG CAPSULE		
	02236859	AGRYLIN	SHI
	02274949	PMS-ANAGRELIDE	PMS
	02260107	SANDOZ ANAGRELIDE	SDZ

20:12.18 PLATELET AGGREGATION INHIBITORS

CLOPIDOGREL BISULFATE

ST	75MG TABLET		
	02303027	ACT CLOPIDOGREL	ACG
	02252767	APO-CLOPIDOGREL	APX
	02416387	AURO-CLOPIDOGREL	AUR
	02385813	CLOPIDOGREL	SIV
	02394820	CLOPIDOGREL	PDL

20:12.18 PLATELET AGGREGATION INHIBITORS

CLOPIDOGREL BISULFATE

ST **75MG TABLET**

02400553	CLOPIDOGREL	SAN
02378507	DOM-CLOPIDOGREL	DPC
02415550	JAMP-CLOPIDOGREL	JMP
02422255	MAR-CLOPIDOGREL	MAR
02238682	PLAVIX	SAC
02348004	PMS-CLOPIDOGREL	PMS
02379813	RAN-CLOPIDOGREL	RBV
02388529	RIVA-CLOPIDOGREL	RIV
02359316	SANDOZ CLOPIDOGREL	SDZ
02293161	TEVA-CLOPIDOGREL	TEV

TICAGRELOR

Limited use benefit (prior approval not required).

For the treatment of Acute Coronary Syndrome, defined as unstable angina or myocardial infarction, when initiated in hospital in consultation with a Specialist in Cardiology, Cardiac Surgery, Cardiovascular & Thoracic Surgery, Internal Medicine or General Surgery. Treatment must be in combination with low dose ASA. Special authorization may be granted for 12 months.

ST **90MG TABLET**

02368544	BRILINTA	AZC
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TICLOPIDINE HYDROCHLORIDE

ST **250MG TABLET**

02237701	TICLOPIDINE	AAP
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20:16.00 HEMATOPOIETIC AGENTS

FILGRASTIM

300MCG/ML INJECTION

09853464	NEUPOGEN (ON)	AMG
99001454	NEUPOGEN (QC)	AMG

300MCG SOLUTION

02441489	GRASTOFIL	APX
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300MCG/ML SOLUTION

01968017	NEUPOGEN	AMG
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480MCG SOLUTION

02454548	GRASTOFIL	APX
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20:16.00 HEMATOPOIETIC AGENTS

PEGFILGRASTIM

Limited use benefit (prior approval required).

CHEMOTHERAPY SUPPORT

Primary Prophylaxis

For use in previously untreated patients receiving a moderate to severely myelosuppressive chemotherapy regimen (i.e. ≥40% incidence of febrile neutropenia). Febrile neutropenia is defined as a temperature ≥38.5°C or >38.0°C three times in a 24 hour period and neutropenia with an absolute neutrophil count (ANC) <0.5 x 10⁹/L.

Secondary Prophylaxis

For use in patients receiving myelosuppressive chemotherapy who have experienced an episode of febrile neutropenic sepsis or profound neutropenia in a previous cycle of chemotherapy; OR
For use in patients who have experienced a dose reduction or treatment delay longer than one week, due to neutropenia.

The recommended dosage of pegfilgrastim is a single subcutaneous injection of 6 mg, administered once per cycle of chemotherapy. Pegfilgrastim should be administered no sooner than 24 hours after the administration of cytotoxic chemotherapy.

10MG/ML SOLUTION

02249790	NEULASTA	AMG
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PEGFILGRASTIM (LAPELGA)

6MG SOLUTION

02474565	LAPELGA	APX
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PLERIXAFOR

Limited use benefit (prior approval required).

For use in combination with filgrastim to mobilize hematopoietic stem cells for subsequent autologous transplantation in patients with:

- Non-Hodgkin's lymphoma (NHL); OR
- Multiple myeloma (MM);

AND

- Prescribed by an oncologist or hematologist.

AND if one of the following are met

- A PBCD34+ count of < 10 cells/uL after 4 days of filgrastim; OR
- Less than 50% of the target CD34 yield is achieved on the 1st day of apheresis (after being mobilized with filgrastim alone or following chemotherapy); OR
- If a patient has failed a previous stem cell mobilization with filgrastim alone or following chemotherapy.

Reimbursement is limited to a maximum of 4 doses (0.24mg/kg given daily) for a single mobilization attempt. The dose of Mozobil is limited to a maximum of 40mg per day

20MG SOLUTION

02377225	MOZOBIL	SAC
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20:24.00 HEMORRHEOLOGIC AGENTS

PENTOXIFYLLINE

ST **400MG TABLET (EXTENDED RELEASE)**

02230090	PENTOXIFYLLINE	AAP
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20:28.16 HEMOSTATICS

TRANEXAMIC ACID

500MG TABLET

02064405	CYKLOKAPRON	PFI
02409097	GD-TRANEXAMIC ACID	PFI
02401231	TRANEXAMIC ACID	RAX

PDIN FOR EXTEMPORANEOUS MIXTURE

99503006	TRANEXAMIC DENTAL MOUTHWASH	UNK
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24:00 CARDIOVASCULAR DRUGS

24:04.04 ANTIARRHYTHMIC AGENTS

AMIODARONE HYDROCHLORIDE

ST 100MG TABLET		
02292173	PMS-AMIODARONE	PMS
ST 200MG TABLET		
02364336	AMIODARONE	SAN
02385465	AMIODARONE	SIV
02246194	APO-AMIODARONE	APX
02246331	DOM-AMIODARONE	DPC
02242472	PMS-AMIODARONE	PMS
02309661	PRO-AMIODARONE	PDL
02247217	RIVA-AMIODARONE	RIV
02243836	SANDOZ AMIODARONE	SDZ
02239835	TEVA-AMIODARONE	TEV
ST PDIN FOR EXTEMPORANEOUS MIXTURE		
99503016	AMIODARONE ORAL LIQUID	UNK

DISOPYRAMIDE

ST 100MG CAPSULE		
02224801	RYTHMODAN	SAC

FLECAINIDE ACETATE

ST 50MG TABLET		
02275538	APO-FLECAINIDE	APX
02459957	AURO-FLECAINIDE	AUR
01966197	TAMBOCOR	VAE
ST 100MG TABLET		
02275546	APO-FLECAINIDE	APX
02459965	AURO-FLECAINIDE	AUR
01966200	TAMBOCOR	VAE

MEXILETINE HYDROCHLORIDE

ST 100MG CAPSULE		
02230359	TEVA-MEXILETINE	TEV
ST 200MG CAPSULE		
02230360	TEVA-MEXILETINE	TEV

PROCAINAMIDE HYDROCHLORIDE

ST 250MG CAPSULE		
00713325	APO-PROCAINAMIDE	APX
ST 250MG TABLET (EXTENDED RELEASE)		
00638692	PROCAN SR	ERF

PROPAFENONE HYDROCHLORIDE

ST 150MG TABLET		
02243324	APO-PROPAFENONE	APX
02457172	MYLAN-PROPAFENONE	MYL
02294559	PMS-PROPAFENONE	PMS
02343053	PROPAFENONE	SAN
00603708	RYTHMOL	BGP
ST 300MG TABLET		
02243325	APO-PROPAFENONE	APX
02457164	MYLAN-PROPAFENONE	MYL
02294575	PMS-PROPAFENONE	PMS
02343061	PROPAFENONE	SAN
00603716	RYTHMOL	BGP

24:04.08 CARDIOTONIC AGENTS

DIGOXIN

ST 0.05MG/ML SOLUTION		
02242320	TOLOXIN	PED
ST 0.0625MG TABLET		
02335700	TOLOXIN	PED
ST 0.125MG TABLET		
02335719	TOLOXIN	PED
ST 0.250MG TABLET		
02335727	TOLOXIN	PED

24:04.92 MISCELLANEOUS CARDIAC DRUGS

IVABRADINE (IVABRADINE HYDROCHLORIDE)

Limited use benefit (prior approval required).

For the treatment of stable chronic heart failure with New York Heart Association (NYHA) class II or III symptoms in adult patients if the following criteria are met:

- Left ventricular ejection fraction \leq 35%; AND
- 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; AND
- Patient has had at least one hospitalization due to heart failure in the last year; AND
- NYHA class II to III symptoms despite at least four weeks of treatment with an angiotensin converting enzyme inhibitor (ACEI) or an angiotensin II receptor antagonist (ARB) in combination with a beta blocker and, if tolerated, a mineralocorticoid receptor antagonist (MRA).

5MG TABLET		
02459973	LANCORA	SEV
7.5MG TABLET		
02459981	LANCORA	SEV

24:06.04 BILE ACID SEQUESTRANTS

CHOLESTYRAMINE RESIN

ST 4G POWDER FOR SUSPENSION		
02455609	CHOLESTYRAMINE-ODAN	ODN
00890960	OLESTYR	PMS
02210320	OLESTYR	PMS

COLESEVELAM HYDROCHLORIDE

ST 3.75G POWDER FOR SUSPENSION		
02432463	LODALIS	VAE
ST 625MG TABLET		
02373955	LODALIS	VAE

COLESTIPOL HYDROCHLORIDE

ST 5G GRANULES		
00642975	COLESTID	PFI
02132699	COLESTID ORANGE	PFI
ST 1G TABLET		
02132680	COLESTID	PFI

24:06.05 CHOLESTEROL ABSORPTION INHIBITORS

EZETIMIBE

ST 10MG TABLET		
02425610	ACH-EZETIMIBE	ACC
02475898	AG-EZETIMIBE	ANG

24:06.05 CHOLESTEROL ABSORPTION INHIBITORS

EZETIMIBE

ST **10MG TABLET**

02427826	APO-EZETIMIBE	APX
02469286	AURO-EZETIMIBE	AUR
02422549	EZETIMIBE	PDL
02429659	EZETIMIBE	SIV
02431300	EZETIMIBE	SAN
02478544	EZETIMIBE	RIV
02247521	EZETROL	FRS
02423235	JAMP-EZETIMIBE	JMP
02422662	MAR-EZETIMIBE	MAR
02467437	M-EZETIMIBE	MAN
02423243	MINT-EZETIMIBE	MIN
02481669	NRA-EZETIMIBE	UNK
02416409	PMS-EZETIMIBE	PMS
02425238	PRIVA-EZETIMIBE	PHA
02419548	RAN-EZETIMIBE	RBV
02424436	RIVA-EZETIMIBE	RIV
02416778	SANDOZ EZETIMIBE	SDZ
02354101	TEVA-EZETIMIBE	TEV

24:06.06 FIBRIC ACID DERIVATIVES

BEZAFIBRATE

ST **200MG TABLET**

02240331	PMS-BEZAFIBRATE	PMS
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ST **400MG TABLET (EXTENDED RELEASE)**

02083523	BEZALIP SR	ACG
02453312	JAMP-BEZAFIBRATE	JMP

FENOFIBRATE

ST **67MG CAPSULE**

02243180	APO-FENO-MICRO	APX
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ST **100MG CAPSULE**

02225980	APO-FENOFIBRATE	APX
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ST **160MG CAPSULE**

02250004	FENOMAX	CIP
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ST **200MG CAPSULE**

02239864	APO-FENO-MICRO	APX
02240360	FENO-MICRO	PDL
02250039	RATIO-FENOFIBRATE	TEV

ST **48MG TABLET**

02269074	LIPIDIL EZ	BGP
02390698	SANDOZ FENOFIBRATE E	SDZ

ST **100MG TABLET**

02246859	APO-FENO-SUPER	APX
02310228	PRO-FENO-SUPER	PDL
02288044	SANDOZ FENOFIBRATE S	SDZ

ST **145MG TABLET**

02269082	LIPIDIL EZ	BGP
02465167	MINT-FENOFIBRATE E	MIN
02390701	SANDOZ FENOFIBRATE E	SDZ

ST **160MG TABLET**

02246860	APO-FENO-SUPER	APX
02241602	LIPIDIL SUPRA	BGP
02310236	PRO-FENO-SUPER	PDL

24:06.06 FIBRIC ACID DERIVATIVES

FENOFIBRATE

ST **160MG TABLET**

02288052	SANDOZ FENOFIBRATE S	SDZ
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GEMFIBROZIL

ST **300MG CAPSULE**

01979574	APO-GEMFIBROZIL	APX
02241608	DOM-GEMFIBROZIL	DPC
02239951	PMS-GEMFIBROZIL	PMS
02241704	TEVA-GEMFIBROZIL	TEV

ST **600MG TABLET**

01979582	APO-GEMFIBROZIL	APX
02142074	TEVA-GEMFIBROZIL	TEV

24:06.08 HMG-COA REDUCTASE INHIBITORS

ATORVASTATIN CALCIUM

ST **10MG TABLET**

02295261	APO-ATORVASTATIN	APX
02346486	ATORVASTATIN	PDL
02348705	ATORVASTATIN	SAN
02396424	ATORVASTATIN	APX
02475022	ATORVASTATIN	RIV
02411350	ATORVASTATIN-10	SIV
02407256	AURO-ATORVASTATIN	AUR
02355612	DOM-ATORVASTATIN	DPC
02399482	DOM-ATORVASTATIN	DPC
02391058	JAMP-ATORVASTATIN	JMP
02230711	LIPITOR	PFI
02454017	MAR-ATORVASTATIN	MAR
02471167	M-ATORVASTATIN	MAN
02392933	MYLAN-ATORVASTATIN	MYL
02313448	PMS-ATORVASTATIN	PMS
02399377	PMS-ATORVASTATIN	PMS
02313707	RAN-ATORVASTATIN	RBV
02417936	REDDY-ATORVASTATIN	REC
02422751	RIVA-ATORVASTATIN	RIV
02324946	SANDOZ ATORVASTATIN	SDZ
02310899	TEVA-ATORVASTATIN	TEV

ST **20MG TABLET**

02295288	APO-ATORVASTATIN	APX
02346494	ATORVASTATIN	PDL
02348713	ATORVASTATIN	SAN
02396432	ATORVASTATIN	APX
02475030	ATORVASTATIN	RIV
02411369	ATORVASTATIN-20	SIV
02407264	AURO-ATORVASTATIN	AUR
02355620	DOM-ATORVASTATIN	DPC
02399490	DOM-ATORVASTATIN	DPC
02391066	JAMP-ATORVASTATIN	JMP
02230713	LIPITOR	PFI
02454025	MAR-ATORVASTATIN	MAR
02471175	M-ATORVASTATIN	MAN
02392941	MYLAN-ATORVASTATIN	MYL
02313456	PMS-ATORVASTATIN	PMS
02399385	PMS-ATORVASTATIN	PMS
02313715	RAN-ATORVASTATIN	RBV

**24:06.08 HMG-COA REDUCTASE
INHIBITORS**

ATORVASTATIN CALCIUM

ST **20MG TABLET**

02417944	REDDY-ATORVASTATIN	REC
02422778	RIVA-ATORVASTATIN	RIV
02324954	SANDOZ ATORVASTATIN	SDZ
02310902	TEVA-ATORVASTATIN	TEV

ST **40MG TABLET**

02295296	APO-ATORVASTATIN	APX
02346508	ATORVASTATIN	PDL
02348721	ATORVASTATIN	SAN
02396440	ATORVASTATIN	APX
02411377	ATORVASTATIN-40	SIV
02407272	AURO-ATORVASTATIN	AUR
02355639	DOM-ATORVASTATIN	DPC
02399504	DOM-ATORVASTATIN	DPC
02391074	JAMP-ATORVASTATIN	JMP
02230714	LIPITOR	PFI
02454033	MAR-ATORVASTATIN	MAR
02471183	M-ATORVASTATIN	MAN
02392968	MYLAN-ATORVASTATIN	MYL
02313464	PMS-ATORVASTATIN	PMS
02399393	PMS-ATORVASTATIN	PMS
02313723	RAN-ATORVASTATIN	RBV
02417952	REDDY-ATORVASTATIN	REC
02422786	RIVA-ATORVASTATIN	RIV
02324962	SANDOZ ATORVASTATIN	SDZ
02310910	TEVA-ATORVASTATIN	TEV

ST **80MG TABLET**

02295318	APO-ATORVASTATIN	APX
02346516	ATORVASTATIN	PDL
02348748	ATORVASTATIN	SAN
02396459	ATORVASTATIN	APX
02475057	ATORVASTATIN	RIV
02411385	ATORVASTATIN-80	SIV
02407280	AURO-ATORVASTATIN	AUR
02391082	JAMP-ATORVASTATIN	JMP
02243097	LIPITOR	PFI
02454041	MAR-ATORVASTATIN	MAR
02471191	M-ATORVASTATIN	MAN
02392976	MYLAN-ATORVASTATIN	MYL
02313472	PMS-ATORVASTATIN	PMS
02399407	PMS-ATORVASTATIN	PMS
02313758	RAN-ATORVASTATIN	RBV
02417960	REDDY-ATORVASTATIN	REC
02422794	RIVA-ATORVASTATIN	RIV
02324970	SANDOZ ATORVASTATIN	SDZ
02310929	TEVA-ATORVASTATIN	TEV

FLUVASTATIN SODIUM

ST **20MG CAPSULE**

02299224	TEVA-FLUVASTATIN	TEV
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ST **40MG CAPSULE**

02299232	TEVA-FLUVASTATIN	TEV
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ST **80MG TABLET (EXTENDED RELEASE)**

02250527	LESCOL XL	NVR
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**24:06.08 HMG-COA REDUCTASE
INHIBITORS**

LOVASTATIN

ST **20MG TABLET**

02248572	ACT LOVASTATIN	ACG
02220172	APO-LOVASTATIN	APX
02353229	LOVASTATIN	SAN
02246013	PMS-LOVASTATIN	PMS
02247056	SANDOZ LOVASTATIN	SDZ

ST **40MG TABLET**

02248573	ACT LOVASTATIN	ACG
02220180	APO-LOVASTATIN	APX
02353237	LOVASTATIN	SAN
02246014	PMS-LOVASTATIN	PMS
02247057	SANDOZ LOVASTATIN	SDZ

PRAVASTATIN SODIUM

ST **10MG TABLET**

02243506	APO-PRAVASTATIN	APX
02458977	AURO-PRAVASTATIN	AUR
02446251	BIO-PRAVASTATIN	BMI
02249723	DOM-PRAVASTATIN	DPC
02330954	JAMP-PRAVASTATIN	JMP
02317451	MINT-PRAVASTATIN	MIN
02476274	M-PRAVASTATIN	MAN
02247655	PMS-PRAVASTATIN	PMS
02356546	PRAVASTATIN	SAN
02389703	PRAVASTATIN	SIV
02243824	PRAVASTATIN-10	PDL
02284421	RAN-PRAVASTATIN	RBV
02468700	SANDOZ PRAVASTATIN	SDZ
02247008	TEVA-PRAVASTATIN	TEV

ST **20MG TABLET**

02243507	APO-PRAVASTATIN	APX
02458985	AURO-PRAVASTATIN	AUR
02446278	BIO-PRAVASTATIN	BMI
02249731	DOM-PRAVASTATIN	DPC
02330962	JAMP-PRAVASTATIN	JMP
02317478	MINT-PRAVASTATIN	MIN
02476282	M-PRAVASTATIN	MAN
02247656	PMS-PRAVASTATIN	PMS
00893757	PRAVACHOL	BMS
02356554	PRAVASTATIN	SAN
02389738	PRAVASTATIN	SIV
02243825	PRAVASTATIN-20	PDL
02284448	RAN-PRAVASTATIN	RBV
02468719	SANDOZ PRAVASTATIN	SDZ
02247009	TEVA-PRAVASTATIN	TEV

ST **40MG TABLET**

02243508	APO-PRAVASTATIN	APX
02458993	AURO-PRAVASTATIN	AUR
02446286	BIO-PRAVASTATIN	BMI
02249758	DOM-PRAVASTATIN	DPC
02330970	JAMP-PRAVASTATIN	JMP
02317486	MINT-PRAVASTATIN	MIN
02476290	M-PRAVASTATIN	MAN
02247657	PMS-PRAVASTATIN	PMS

**24:06.08 HMG-COA REDUCTASE
INHIBITORS**

PRAVASTATIN SODIUM

ST **40MG TABLET**

02222051	PRAVACHOL	BMS
02356562	PRAVASTATIN	SAN
02389746	PRAVASTATIN	SIV
02243826	PRAVASTATIN-40	PDL
02284456	RAN-PRAVASTATIN	RBV
02468727	SANDOZ PRAVASTATIN	SDZ
02247010	TEVA-PRAVASTATIN	TEV

ROSUVASTATIN CALCIUM

ST **5MG TABLET**

02339765	ACT ROSUVASTATIN	ACG
02337975	APO-ROSUVASTATIN	APX
02442574	AURO-ROSUVASTATIN	AUR
02444968	BIO-ROSUVASTATIN	BMI
02265540	CRESTOR	AZC
02386704	DOM-ROSUVASTATIN	DPC
02391252	JAMP-ROSUVASTATIN	JMP
02413051	MAR-ROSUVASTATIN	MAR
02399164	MED-ROSUVASTATIN	GMP
02397781	MINT-ROSUVASTATIN	MIN
02378523	PMS-ROSUVASTATIN	PMS
02380013	RIVA-ROSUVASTATIN	RIV
02381176	ROSUVASTATIN	PDL
02405628	ROSUVASTATIN	SAN
02411628	ROSUVASTATIN	SIV
02338726	SANDOZ ROSUVASTATIN	SDZ
02382644	SUNPHARMA ROSUVASTATIN	SUN
02354608	TEVA-ROSUVASTATIN	TEV

ST **10MG TABLET**

02339773	ACT ROSUVASTATIN	ACG
02337983	APO-ROSUVASTATIN	APX
02442582	AURO-ROSUVASTATIN	AUR
02444976	BIO-ROSUVASTATIN	BMI
02247162	CRESTOR	AZC
02386712	DOM-ROSUVASTATIN	DPC
02391260	JAMP-ROSUVASTATIN	JMP
02413078	MAR-ROSUVASTATIN	MAR
02399172	MED-ROSUVASTATIN	GMP
02397803	MINT-ROSUVASTATIN	MIN
02378531	PMS-ROSUVASTATIN	PMS
02380056	RIVA-ROSUVASTATIN	RIV
02381184	ROSUVASTATIN	PDL
02405636	ROSUVASTATIN	SAN
02411636	ROSUVASTATIN	SIV
02338734	SANDOZ ROSUVASTATIN	SDZ
02382652	SUNPHARMA ROSUVASTATIN	SUN
02354616	TEVA-ROSUVASTATIN	TEV

ST **20MG TABLET**

02339781	ACT ROSUVASTATIN	ACG
02337991	APO-ROSUVASTATIN	APX
02442590	AURO-ROSUVASTATIN	AUR
02444984	BIO-ROSUVASTATIN	BMI
02247163	CRESTOR	AZC

**24:06.08 HMG-COA REDUCTASE
INHIBITORS**

ROSUVASTATIN CALCIUM

ST **20MG TABLET**

02386720	DOM-ROSUVASTATIN	DPC
02391279	JAMP-ROSUVASTATIN	JMP
02413086	MAR-ROSUVASTATIN	MAR
02399180	MED-ROSUVASTATIN	GMP
02397811	MINT-ROSUVASTATIN	MIN
02378558	PMS-ROSUVASTATIN	PMS
02380064	RIVA-ROSUVASTATIN	RIV
02381192	ROSUVASTATIN	PDL
02405644	ROSUVASTATIN	SAN
02411644	ROSUVASTATIN	SIV
02338742	SANDOZ ROSUVASTATIN	SDZ
02382660	SUNPHARMA ROSUVASTATIN	SUN
02354624	TEVA-ROSUVASTATIN	TEV

ST **40MG TABLET**

02339803	ACT ROSUVASTATIN	ACG
02338009	APO-ROSUVASTATIN	APX
02442604	AURO-ROSUVASTATIN	AUR
02444992	BIO-ROSUVASTATIN	BMI
02247164	CRESTOR	AZC
02391287	JAMP-ROSUVASTATIN	JMP
02413108	MAR-ROSUVASTATIN	MAR
02399199	MED-ROSUVASTATIN	GMP
02397838	MINT-ROSUVASTATIN	MIN
02378566	PMS-ROSUVASTATIN	PMS
02380102	RIVA-ROSUVASTATIN	RIV
02381206	ROSUVASTATIN	PDL
02405652	ROSUVASTATIN	SAN
02411652	ROSUVASTATIN	SIV
02338750	SANDOZ ROSUVASTATIN	SDZ
02382679	SUNPHARMA ROSUVASTATIN	SUN
02354632	TEVA-ROSUVASTATIN	TEV

SIMVASTATIN

ST **5MG TABLET**

02247011	APO-SIMVASTATIN	APX
02405148	AURO-SIMVASTATIN	AUR
02253747	DOM-SIMVASTATIN	DPC
02281619	DOM-SIMVASTATIN	DPC
02375591	JAMP-SIMVASTATIN	JMP
02375036	MAR-SIMVASTATIN	MAR
02372932	MINT-SIMVASTATIN	MIN
02246582	MYLAN-SIMVASTATIN	MYL
02469979	PHARMA-SIMVASTATIN	PMS
02269252	PMS-SIMVASTATIN	PMS
02329131	RAN-SIMVASTATIN	RBV
02247827	SANDOZ SIMVASTATIN	SDZ
02284723	SIMVASTATIN	SAN
02386291	SIMVASTATIN	SIV
02250144	TEVA-SIMVASTATIN	TEV

ST **10MG TABLET**

02247012	APO-SIMVASTATIN	APX
02405156	AURO-SIMVASTATIN	AUR
02253755	DOM-SIMVASTATIN	DPC

**24:06.08 HMG-COA REDUCTASE
INHIBITORS**

SIMVASTATIN

ST **10MG TABLET**

02281627	DOM-SIMVASTATIN	DPC
02375605	JAMP-SIMVASTATIN	JMP
02375044	MAR-SIMVASTATIN	MAR
02372940	MINT-SIMVASTATIN	MIN
02246583	MYLAN-SIMVASTATIN	MYL
02469987	PHARMA-SIMVASTATIN	PMS
02269260	PMS-SIMVASTATIN	PMS
02329158	RAN-SIMVASTATIN	RBY
02247828	SANDOZ SIMVASTATIN	SDZ
02284731	SIMVASTATIN	SAN
02386305	SIMVASTATIN	SIV
02247221	SIMVASTATIN-10	PDL
02250152	TEVA-SIMVASTATIN	TEV
00884332	ZOCOR	FRS

ST **20MG TABLET**

02247013	APO-SIMVASTATIN	APX
02405164	AURO-SIMVASTATIN	AUR
02253763	DOM-SIMVASTATIN	DPC
02281635	DOM-SIMVASTATIN	DPC
02375613	JAMP-SIMVASTATIN	JMP
02375052	MAR-SIMVASTATIN	MAR
02372959	MINT-SIMVASTATIN	MIN
02246737	MYLAN-SIMVASTATIN	MYL
02469995	PHARMA-SIMVASTATIN	PMS
02269279	PMS-SIMVASTATIN	PMS
02329166	RAN-SIMVASTATIN	RBY
02247830	SANDOZ SIMVASTATIN	SDZ
02284758	SIMVASTATIN	SAN
02386313	SIMVASTATIN	SIV
02247222	SIMVASTATIN-20	PDL
02250160	TEVA-SIMVASTATIN	TEV
00884340	ZOCOR	FRS

ST **40MG TABLET**

02247014	APO-SIMVASTATIN	APX
02405172	AURO-SIMVASTATIN	AUR
02253771	DOM-SIMVASTATIN	DPC
02281643	DOM-SIMVASTATIN	DPC
02375621	JAMP-SIMVASTATIN	JMP
02375060	MAR-SIMVASTATIN	MAR
02372967	MINT-SIMVASTATIN	MIN
02246584	MYLAN-SIMVASTATIN	MYL
02470004	PHARMA-SIMVASTATIN	PMS
02269287	PMS-SIMVASTATIN	PMS
02329174	RAN-SIMVASTATIN	RBY
02247831	SANDOZ SIMVASTATIN	SDZ
02284766	SIMVASTATIN	SAN
02386321	SIMVASTATIN	SIV
02247223	SIMVASTATIN-40	PDL
02250179	TEVA-SIMVASTATIN	TEV
00884359	ZOCOR	FRS

ST **80MG TABLET**

02247015	APO-SIMVASTATIN	APX
02405180	AURO-SIMVASTATIN	AUR

**24:06.08 HMG-COA REDUCTASE
INHIBITORS**

SIMVASTATIN

ST **80MG TABLET**

02253798	DOM-SIMVASTATIN	DPC
02281651	DOM-SIMVASTATIN	DPC
02375648	JAMP-SIMVASTATIN	JMP
02375079	MAR-SIMVASTATIN	MAR
02372975	MINT-SIMVASTATIN	MIN
02246585	MYLAN-SIMVASTATIN	MYL
02470012	PHARMA-SIMVASTATIN	PMS
02269295	PMS-SIMVASTATIN	PMS
02329182	RAN-SIMVASTATIN	RBY
02247833	SANDOZ SIMVASTATIN	SDZ
02284774	SIMVASTATIN	SAN
02386348	SIMVASTATIN	SIV
02247224	SIMVASTATIN-80	PDL
02250187	TEVA-SIMVASTATIN	TEV

24:06.24

ALIROCUMAB

Limited use benefit (prior approval required).

Initial Coverage (12 weeks):

For adult patients with heterozygous familial hypercholesterolemia (HeFH) who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following clinical 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 an LDL-C target of < 2.0 mmol/L for secondary CVD prevention, or at least a 50% reduction in LDL-C from untreated baseline for primary prevention despite:
 - Confirmed adherence to a high-dose statin (e.g., atorvastatin 80 mg or rosuvastatin 40 mg) in combination with ezetimibe for a total of at least 3 months of continuous treatment;
 - OR
 - Patient is unable to tolerate at least two statins, with at least one statin initiated at the lowest daily starting dose; AND
 - For each statin (two statins in total), dose reduction was attempted to resolve intolerable myopathy or creatine kinase > 5 times the upper limit of normal rather than statin discontinuation; AND
 - For each statin (two statins in total), intolerable myopathy or creatine kinase > 5 times the upper limit of normal was reversed upon statin discontinuation, but reoccurred with statin rechallenge where clinically appropriate; AND
 - Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment; AND
 - Other known determinants of intolerable symptoms or abnormal biomarkers have been ruled out;
 - OR
 - Patient developed confirmed and documented rhabdomyolysis;
 - OR
 - Patient has a contraindication to statins; AND
 - Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment.

Continued coverage (6 months):

- Patient is adherent to therapy; AND
- Patient has achieved a reduction in LDL-C of at least 40% from baseline.

Note: Annual coverage is limited to 26 prefilled syringes or

75MG SOLUTION

02453754	PRALUENT	SAC
02453819	PRALUENT	SAC

150MG SOLUTION

02453762	PRALUENT	SAC
02453835	PRALUENT	SAC

24:06.24

EVOLOCUMAB

Limited use benefit (prior approval required).

Initial coverage criteria (Initial approval for 12 weeks):

For adult patients with heterozygous familial hypercholesterolemia (HeFH) who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following clinical 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 an LDL-C target of < 2.0 mmol/L for secondary CVD prevention, or at least a 50% reduction in LDL-C from untreated baseline for primary prevention despite:
 - Confirmed adherence to a high-dose statin (e.g., atorvastatin 80 mg or rosuvastatin 40 mg) in combination with ezetimibe for a total of at least 3 months of continuous treatment;
 - OR
 - Patient is unable to tolerate at least two statins, with at least one statin initiated at the lowest daily starting dose; AND
 - For each statin (two statins in total), dose reduction was attempted to resolve intolerable myopathy or creatine kinase > 5 times the upper limit of normal rather than statin discontinuation; AND
 - For each statin (two statins in total), intolerable myopathy or creatine kinase > 5 times the upper limit of normal was reversed upon statin discontinuation, but reoccurred with statin rechallenge where clinically appropriate; AND
 - Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment; AND
 - Other known determinants of intolerable symptoms or abnormal biomarkers have been ruled out;
 - OR
 - Patient developed confirmed and documented rhabdomyolysis;
 - OR
 - Patient has a contraindication to statins; AND
 - Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment.

Note: Annual coverage is limited to 26 prefilled autoinjectors (140 mg every 2 weeks) or 12 automated Mini-Dosers with prefilled cartridges (420 mg once a month).

Renewal coverage criteria (Renewal for 6 months):

- Patient is adherent to therapy; AND
- Patient has achieved a reduction in LDL-C of at least 40% from baseline.

Note: Annual coverage is limited to 26 prefilled Autoinjectors (140 mg every 2 weeks) or 12 automated Mini-Dosers with

120MG SOLUTION

02459779	REPATHA	AMG
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140MG SOLUTION

02446057	REPATHA	AMG
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24:08.16 CENTRAL ALPHA-AGONISTS

CLONIDINE HYDROCHLORIDE

ST **0.025MG TABLET**

02304163	TEVA-CLONIDINE	TEV
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ST **0.1MG TABLET**

02462192	MINT-CLONIDINE	MIN
02046121	TEVA-CLONIDINE	TEV

24:08.16 CENTRAL ALPHA-AGONISTS

CLONIDINE HYDROCHLORIDE

ST **0.2MG TABLET**

00868957	APO-CLONIDINE	APX
02462206	MINT-CLONIDINE	MIN
02046148	TEVA-CLONIDINE	TEV

ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503021	CLONIDINE ORAL LIQUID	UNK
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METHYLDOPA

ST **125MG TABLET**

00360252	METHYLDOPA	AAP
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ST **250MG TABLET**

00360260	METHYLDOPA	AAP
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ST **500MG TABLET**

00426830	METHYLDOPA	AAP
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METHYLDOPA, HYDROCHLOROTHIAZIDE

ST **250MG & 15MG TABLET**

00441708	APO METHAZIDE	APX
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ST **250MG & 25MG TABLET**

00441716	APO METHAZIDE	APX
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24:08.20 DIRECT VASODILATORS

DIAZOXIDE

ST **100MG CAPSULE**

00503347	PROGLYCEM	FRS
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HYDRALAZINE HYDROCHLORIDE

ST **10MG TABLET**

00441619	APO-HYDRALAZINE	APX
02457865	JAMP-HYDRALAZINE	JMP
02468778	MINT-HYDRALAZINE	MIN

ST **25MG TABLET**

00441627	APO-HYDRALAZINE	APX
02457873	JAMP-HYDRALAZINE	JMP
02468786	MINT-HYDRALAZINE	MIN

ST **50MG TABLET**

00441635	APO-HYDRALAZINE	APX
02457881	JAMP-HYDRALAZINE	JMP
02468794	MINT-HYDRALAZINE	MIN

MINOXIDIL

ST **2.5MG TABLET**

00514497	LONITEN	PFI
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ST **10MG TABLET**

00514500	LONITEN	PFI
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24:12.08 NITRATES AND NITRITES

ISOSORBIDE DINITRATE

ST **5MG TABLET**

00670944	ISDN	AAP
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ST **10MG TABLET**

00441686	ISDN	AAP
00786667	PMS-ISOSORBIDE	PMS

ST **30MG TABLET**

00441694	ISDN	AAP
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24:12.08 NITRATES AND NITRITES

ISOSORBIDE-5-MONONITRATE

ST **60MG TABLET (EXTENDED RELEASE)**

02272830	APO-ISMN	APX
02126559	IMDUR	UNK
02301288	PMS-ISMN	PMS
02311321	PRO-ISMN	PDL

NITROGLYCERIN

ST **0.2MG PATCH**

02162806	MINITRAN	VAE
02407442	MYLAN-NITRO	MYL
01911910	NITRO-DUR	FRS
00584223	TRANSDERM-NITRO	NVR
02230732	TRINIPATCH	PAL

ST **0.4MG PATCH**

02163527	MINITRAN	VAE
02407450	MYLAN-NITRO	MYL
01911902	NITRO-DUR	FRS
00852384	TRANSDERM-NITRO	NVR
02230733	TRINIPATCH	PAL

ST **0.6MG PATCH**

02163535	MINITRAN	VAE
02407469	MYLAN-NITRO	MYL
01911929	NITRO-DUR	FRS
02046156	TRANSDERM-NITRO	NVR
02230734	TRINIPATCH	PAL

ST **0.8MG PATCH**

02407477	MYLAN-NITRO	MYL
02011271	NITRO-DUR	FRS

0.4MG PUMP

02393433	APO-NITROGLYCERIN	APX
02243588	MYLAN-NITRO	MYL
02231441	NITROLINGUAL PUMPSPRAY	SAC
02238998	RHO-NITRO PUMPSPRAY	SDZ

ST **0.3MG TABLET**

00037613	NITROSTAT	PFI
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ST **0.6MG TABLET**

00037621	NITROSTAT	PFI
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24:12.12 PHOSPHODIESTERASE INHIBITORS

SILDENAFIL CITRATE

Limited use benefit (prior approval required).

Must be initiated by a Pulmonary Hypertension specialist

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization.

ST **20MG TABLET**

02418118	APO-SILDENAFIL R	APX
02412179	PMS-SILDENAFIL R	PMS
02279401	REVATIO	PFI
02319500	TEVA-SILDENAFIL R	TEV

24:12.12 PHOSPHODIESTERASE INHIBITORS

TADALAFIL

Limited use benefit (prior approval required).

Must be initiated by a Pulmonary Hypertension specialist

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization.

ST 20MG TABLET

02338327	ADCIRCA	LIL
02421933	APO-TADALAFIL PAH	APX

24:12.92 MISCELLANEOUS VASODILATING AGENTS

AMBRISENTAN

Limited use benefit (prior approval required).

Maximum dose covered is 10 mg once daily.

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST 5MG TABLET

02307065	VOLIBRIS	GSK
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ST 10MG TABLET

02307073	VOLIBRIS	GSK
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BOSENTAN MONOHYDRATE

Limited use benefit (prior approval required).

Maximum dose covered is 125 mg twice daily

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST 62.5MG TABLET

02399202	APO-BOSENTAN	APX
02383012	PMS-BOSENTAN	PMS
02386275	SANDOZ BOSENTAN	SDZ
02398400	TEVA-BOSENTAN	TEV
02244981	TRACLEER	JSO

ST 125MG TABLET

02383020	PMS-BOSENTAN	PMS
02386283	SANDOZ BOSENTAN	SDZ
02244982	TRACLEER	JSO

DIPYRIDAMOLE

ST 25MG TABLET

00895644	APO-DIPYRIDAMOLE	APX
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ST 50MG TABLET

00571245	APO-DIPYRIDAMOLE	APX
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24:12.92 MISCELLANEOUS VASODILATING AGENTS

DIPYRIDAMOLE

ST 50MG TABLET

00895652	APO-DIPYRIDAMOLE	APX
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ST 75MG TABLET

00601845	APO-DIPYRIDAMOLE	APX
00895660	APO-DIPYRIDAMOLE	APX

DIPYRIDAMOLE, ACETYSALICYLIC ACID

ST 200MG & 25MG CAPSULE (IMMEDIATE AND EXTENDED RELEASE)

02242119	AGGRENOX	BOE
02471051	TARO-DIPYRIDAMOLE/ ASA	TAR

24:20.00 ALPHA ADRENERGIC BLOCKING AGENTS

DOXAZOSIN MESYLATE

ST 1MG TABLET

02240588	APO-DOXAZOSIN	APX
02244527	PMS-DOXAZOSIN	PMS
02242728	TEVA-DOXAZOSIN	TEV

ST 2MG TABLET

02240589	APO-DOXAZOSIN	APX
02244528	PMS-DOXAZOSIN	PMS
02242729	TEVA-DOXAZOSIN	TEV

ST 4MG TABLET

02240590	APO-DOXAZOSIN	APX
02244529	PMS-DOXAZOSIN	PMS
02242730	TEVA-DOXAZOSIN	TEV

PRAZOSIN HYDROCHLORIDE

ST 1MG TABLET

00882801	APO-PRAZO	APX
00560952	MINIPRESS	ERF
01934198	TEVA-PRAZOSIN	TEV

ST 2MG TABLET

00882828	APO-PRAZO	APX
00560960	MINIPRESS	ERF
01934201	TEVA-PRAZOSIN	TEV

ST 5MG TABLET

00882836	APO-PRAZO	APX
00560979	MINIPRESS	ERF
01934228	TEVA-PRAZOSIN	TEV

TERAZOSIN HYDROCHLORIDE

ST 1MG TABLET

02234502	APO-TERAZOSIN	APX
02243746	DOM-TERAZOSIN	DPC
02243518	PMS-TERAZOSIN	PMS
02237476	TERAZOSIN	PDL
02350475	TERAZOSIN	SAN
02230805	TEVA-TERAZOSIN	TEV

ST 2MG TABLET

02234503	APO-TERAZOSIN	APX
02243747	DOM-TERAZOSIN	DPC
02243519	PMS-TERAZOSIN	PMS
02237477	TERAZOSIN	PDL

24:20.00 ALPHA ADRENERGIC BLOCKING AGENTS

TERAZOSIN HYDROCHLORIDE

ST **2MG TABLET**

02350483	TERAZOSIN	SAN
02230806	TEVA-TERAZOSIN	TEV

ST **5MG TABLET**

02234504	APO-TERAZOSIN	APX
02243748	DOM-TERAZOSIN	DPC
02243520	PMS-TERAZOSIN	PMS
02237478	TERAZOSIN	PDL
02350491	TERAZOSIN	SAN
02230807	TEVA-TERAZOSIN	TEV

ST **10MG TABLET**

02234505	APO-TERAZOSIN	APX
02243749	DOM-TERAZOSIN	DPC
02243521	PMS-TERAZOSIN	PMS
02237479	TERAZOSIN	PDL
02350505	TERAZOSIN	SAN
02230808	TEVA-TERAZOSIN	TEV

24:24.00 BETA ADRENERGIC BLOCKING AGENTS

ACEBUTOLOL HYDROCHLORIDE

ST **100MG TABLET**

02164396	ACEBUTOLOL	PDL
02147602	APO-ACEBUTOLOL	APX
02204517	TEVA-ACEBUTOLOL	TEV

ST **200MG TABLET**

02164418	ACEBUTOLOL	PDL
02147610	APO-ACEBUTOLOL	APX
02204525	TEVA-ACEBUTOLOL	TEV

ST **400MG TABLET**

02164426	ACEBUTOLOL	PDL
02147629	APO-ACEBUTOLOL	APX
02204533	TEVA-ACEBUTOLOL	TEV

ATENOLOL

ST **25MG TABLET**

02326701	ATENOLOL	PDL
02392194	BIO-ATENOLOL	BMI
02367556	JAMP-ATENOLOL	JMP
02371979	MAR-ATENOLOL	MAR
02368013	MINT-ATENOL	MIN
02246581	PMS-ATENOLOL	PMS
02373963	RAN-ATENOLOL	RBY
02277379	RIVA-ATENOLOL	RIV
02368633	SEPTA-ATENOLOL	SPT
02266660	TEVA-ATENOLOL	TEV

ST **50MG TABLET**

02255545	ACT ATENOLOL	ACG
00773689	APO-ATENOL	APX
00828807	ATENOLOL	PDL
02238316	ATENOLOL	SIV
02466465	ATENOLOL	SAN
02392178	BIO-ATENOLOL	BMI
02229467	DOM-ATENOLOL	DPC

24:24.00 BETA ADRENERGIC BLOCKING AGENTS

ATENOLOL

ST **50MG TABLET**

02367564	JAMP-ATENOLOL	JMP
02371987	MAR-ATENOLOL	MAR
02368021	MINT-ATENOL	MIN
02237600	PMS-ATENOLOL	PMS
02267985	RAN-ATENOLOL	RBY
02242094	RIVA-ATENOLOL	RIV
02368641	SEPTA-ATENOLOL	SPT
02039532	TENORMIN	AZC
02171791	TEVA-ATENOLOL	TEV

ST **100MG TABLET**

02255553	ACT ATENOLOL	ACG
00773697	APO-ATENOL	APX
00828793	ATENOLOL	PDL
02238318	ATENOLOL	SIV
02466473	ATENOLOL	SAN
02392186	BIO-ATENOLOL	BMI
02229468	DOM-ATENOLOL	DPC
02367572	JAMP-ATENOLOL	JMP
02371995	MAR-ATENOLOL	MAR
02368048	MINT-ATENOL	MIN
02237601	PMS-ATENOLOL	PMS
02267993	RAN-ATENOLOL	RBY
02242093	RIVA-ATENOLOL	RIV
02368668	SEPTA-ATENOLOL	SPT
02039540	TENORMIN	AZC
02171805	TEVA-ATENOLOL	TEV

ATENOLOL, CHLORTHALIDONE

ST **50MG & 25MG TABLET**

02248763	APO-ATENIDONE	APX
02049961	TENORETIC	AZC
02302918	TEVA-ATENOLOL/CHLORTHALIDONE	TEV

ST **100MG & 25MG TABLET**

02248764	APO-ATENIDONE	APX
02049988	TENORETIC	AZC
02302926	TEVA-ATENOLOL/CHLORTHALIDONE	TEV

BISOPROLOL FUMARATE

ST **5MG TABLET**

02256134	APO-BISOPROLOL	APX
02383055	BISOPROLOL	SIV
02391589	BISOPROLOL	SAN
02465612	MINT-BISOPROLOL	MIN
02302632	PMS-BISOPROLOL	PMS
02306999	PRO-BISOPROLOL	PDL
02471264	RIVA-BISOPROLOL	RIV
02247439	SANDOZ BISOPROLOL	SDZ
02267470	TEVA-BISOPROLOL	TEV

ST **10MG TABLET**

02256177	APO-BISOPROLOL	APX
02383063	BISOPROLOL	SIV
02391597	BISOPROLOL	SAN

24:24.00 BETA ADRENERGIC BLOCKING AGENTS

BISOPROLOL FUMARATE

ST **10MG TABLET**

02465620	MINT-BISOPROLOL	MIN
02302640	PMS-BISOPROLOL	PMS
02307006	PRO-BISOPROLOL	PDL
02471272	RIVA-BISOPROLOL	RIV
02247440	SANDOZ BISOPROLOL	SDZ
02267489	TEVA-BISOPROLOL	TEV

CARVEDILOL

ST **3.125MG TABLET**

02247933	APO-CARVEDILOL	APX
02418495	AURO-CARVEDILOL	AUR
02248752	CARVEDILOL	SIV
02324504	CARVEDILOL	PDL
02364913	CARVEDILOL	SAN
02248748	DOM-CARVEDILOL	DPC
02368897	JAMP-CARVEDILOL	JMP
02245914	PMS-CARVEDILOL	PMS
02268027	RAN-CARVEDILOL	RBY
02252309	TEVA-CARVEDILOL	TEV

ST **6.25MG TABLET**

02247934	APO-CARVEDILOL	APX
02418509	AURO-CARVEDILOL	AUR
02248753	CARVEDILOL	SIV
02324512	CARVEDILOL	PDL
02364921	CARVEDILOL	SAN
02248749	DOM-CARVEDILOL	DPC
02368900	JAMP-CARVEDILOL	JMP
02245915	PMS-CARVEDILOL	PMS
02268035	RAN-CARVEDILOL	RBY
02252317	TEVA-CARVEDILOL	TEV

ST **12.5MG TABLET**

02247935	APO-CARVEDILOL	APX
02418517	AURO-CARVEDILOL	AUR
02248754	CARVEDILOL	SIV
02324520	CARVEDILOL	PDL
02364948	CARVEDILOL	SAN
02248750	DOM-CARVEDILOL	DPC
02368919	JAMP-CARVEDILOL	JMP
02245916	PMS-CARVEDILOL	PMS
02268043	RAN-CARVEDILOL	RBY
02252325	TEVA-CARVEDILOL	TEV

ST **25MG TABLET**

02247936	APO-CARVEDILOL	APX
02418525	AURO-CARVEDILOL	AUR
02248755	CARVEDILOL	SIV
02324539	CARVEDILOL	PDL
02364956	CARVEDILOL	SAN
02248751	DOM-CARVEDILOL	DPC
02368927	JAMP-CARVEDILOL	JMP
02245917	PMS-CARVEDILOL	PMS
02268051	RAN-CARVEDILOL	RBY
02252333	TEVA-CARVEDILOL	TEV

24:24.00 BETA ADRENERGIC BLOCKING AGENTS

HYDROCHLOROTHIAZIDE, PINDOLOL

ST **10MG & 25MG TABLET**

00568627	VISKAZIDE	UNK
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ST **10MG & 50MG TABLET**

00568635	VISKAZIDE	UNK
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LABETALOL HYDROCHLORIDE

ST **100MG TABLET**

02106272	TRANDATE	PAL
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ST **200MG TABLET**

02106280	TRANDATE	PAL
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METOPROLOL TARTRATE

ST **25MG TABLET**

02246010	APO-METOPROLOL	APX
02252252	DOM-METOPROLOL-L	DPC
02356813	JAMP-METOPROLOL-L	JMP
02296713	METOPROLOL	PDL
02442116	METOPROLOL-L	SIV
02248855	PMS-METOPROLOL-L	PMS
02315300	RIVA-METOPROLOL L	RIV
02261898	TEVA-METOPROLOL	TEV

ST **50MG TABLET**

00618632	APO METOPROLOL	APX
00749354	APO METOPROLOL (TYPE L)	APX
02172550	DOM-METOPROLOL-B	DPC
02231121	DOM-METOPROLOL-L	DPC
02356821	JAMP-METOPROLOL-L	JMP
00397423	LOPRESOR	NVR
00648019	METOPROLOL	PDL
02350394	METOPROLOL	SAN
02442124	METOPROLOL-L	SIV
02145413	PMS-METOPROLOL-B	PMS
02230803	PMS-METOPROLOL-L	PMS
02315319	RIVA-METOPROLOL L	RIV
00648035	TEVA-METOPROLOL	TEV
00842648	TEVA-METOPROLOL	TEV

ST **100MG TABLET**

00618640	APO METOPROLOL	APX
00751170	APO-METOPROLOL (TYPE L)	APX
02172569	DOM-METOPROLOL-B	DPC
02231122	DOM-METOPROLOL-L	DPC
02356848	JAMP-METOPROLOL-L	JMP
00397431	LOPRESOR	NVR
00648027	METOPROLOL	PDL
02350408	METOPROLOL	SAN
02442132	METOPROLOL-L	SIV
02145421	PMS-METOPROLOL-B	PMS
02230804	PMS-METOPROLOL-L	PMS
02315327	RIVA-METOPROLOL L	RIV
00648043	TEVA-METOPROLOL	TEV
00842656	TEVA-METOPROLOL	TEV

ST **100MG TABLET (EXTENDED RELEASE)**

02285169	APO-METOPROLOL SR	APX
00658855	LOPRESOR SR	NVR
02351404	METOPROLOL SR	PDL

24:24.00 BETA ADRENERGIC BLOCKING AGENTS

METOPROLOL TARTRATE

ST 100MG TABLET (EXTENDED RELEASE)			
02303396	SANDOZ METOPROLOL SR	SDZ	
ST 200MG TABLET (EXTENDED RELEASE)			
02285177	APO-METOPROLOL SR	APX	
00534560	LOPRESOR SR	NVR	
02303418	SANDOZ METOPROLOL SR	SDZ	
ST PDIN FOR EXTEMPORANEOUS MIXTURE			
99503015	METOPROLOL ORAL LIQUID	UNK	

NADOLOL

ST 40MG TABLET			
00782505	NADOLOL	AAP	
ST 80MG TABLET			
00782467	NADOLOL	AAP	
ST 160MG TABLET			
00782475	NADOLOL	AAP	

PINDOLOL

ST 5MG TABLET			
00755877	APO-PINDOL	APX	
00828416	PINDOLOL	PDL	
00869007	TEVA-PINDOLOL	TEV	
00417270	VISKEN	UNK	
ST 10MG TABLET			
00755885	APO-PINDOL	APX	
00828424	PINDOLOL	PDL	
00869015	TEVA-PINDOLOL	TEV	
00443174	VISKEN	UNK	
ST 15MG TABLET			
00755893	APO-PINDOL	APX	
02238047	DOM-PINDOLOL	DPC	
02231539	PMS-PINDOLOL	PMS	
00869023	TEVA-PINDOLOL	TEV	

PROPRANOLOL (HEMANGIOL)

Limited use benefit (prior approval required).

For the treatment of proliferating infantile hemangioma requiring systemic therapy and at least one of the following:

- Life or function-threatening hemangioma, OR
- Ulcerated hemangioma with pain and/or lack of response to simple wound care measures, OR
- Hemangioma with a risk of permanent scarring or disfigurement.

3.75MG SOLUTION

02457857	HEMANGIOL	PFD	
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PROPRANOLOL HYDROCHLORIDE

ST 60MG CAPSULE (SUSTAINED RELEASE)			
02042231	INDERAL LA	PFI	
ST 80MG CAPSULE (SUSTAINED RELEASE)			
02042258	INDERAL LA	PFI	
ST 120MG CAPSULE (SUSTAINED RELEASE)			
02042266	INDERAL LA	PFI	
ST 160MG CAPSULE (SUSTAINED RELEASE)			
02042274	INDERAL LA	PFI	

24:24.00 BETA ADRENERGIC BLOCKING AGENTS

PROPRANOLOL HYDROCHLORIDE

ST 10MG TABLET			
00496480	TEVA-PROPRANOLOL	TEV	
ST 20MG TABLET			
00740675	TEVA-PROPRANOLOL	TEV	
ST 40MG TABLET			
00496499	TEVA-PROPRANOLOL	TEV	
ST 80MG TABLET			
00582271	PMS-PROPRANOLOL	PMS	
00496502	TEVA-PROPRANOLOL	TEV	
ST 120MG TABLET			
00504335	APO PROPRANOLOL	APX	
00582298	PMS-PROPRANOLOL	PMS	
ST PDIN FOR EXTEMPORANEOUS MIXTURE			
99503014	PROPRANOLOL ORAL LIQUID	UNK	

SOTALOL HYDROCHLORIDE

ST 80MG TABLET			
02210428	APO-SOTALOL	APX	
02238634	DOM-SOTALOL	DPC	
02368617	JAMP-SOTALOL	JMP	
02238326	PMS-SOTALOL	PMS	
02316528	PRO-SOTALOL	PDL	
ST 160MG TABLET			
02167794	APO-SOTALOL	APX	
02238635	DOM-SOTALOL	DPC	
02368625	JAMP-SOTALOL	JMP	
02238327	PMS-SOTALOL	PMS	
02316536	PRO-SOTALOL	PDL	
ST PDIN FOR EXTEMPORANEOUS MIXTURE			
99503023	SOTALOL ORAL LIQUID	UNK	

TIMOLOL MALEATE

ST 5MG TABLET			
00755842	TIMOLOL	APX	
ST 10MG TABLET			
00755850	TIMOLOL	APX	
ST 20MG TABLET			
00755869	TIMOLOL	APX	

24:28.08 DIHYDROPYRIDINES

AMLODIPINE BESYLATE

ST 2.5MG TABLET			
02297477	ACT AMLODIPINE	ACG	
02326795	AMLODIPINE	PDL	
02385783	AMLODIPINE	SIV	
02419556	AMLODIPINE BESYLATE	ACC	
02392127	BIO-AMLODIPINE	BMI	
02326825	DOM-AMLODIPINE	DPC	
02280124	GD-AMLODIPINE	PFI	
02357186	JAMP-AMLODIPINE	JMP	
02468018	M-AMLODIPINE	MAN	
02371707	MAR-AMLODIPINE	MAR	
02476452	NRA-AMLODIPINE	UNK	
02469022	PHARMA-AMLODIPINE	PMS	
02295148	PMS-AMLODIPINE	PMS	

24:28.08 DIHYDROPYRIDINES

AMLODIPINE BESYLATE

ST 2.5MG TABLET

02398877	RAN-AMLODIPINE	RBY
02331489	RIVA-AMLODIPINE	RIV
02330474	SANDOZ AMLODIPINE	SDZ
02357704	SEPTA-AMLODIPINE	SPT

ST 5MG TABLET

02297485	ACT AMLODIPINE	ACG
02369230	AG-AMLODIPINE	ANG
02326809	AMLODIPINE	PDL
02331284	AMLODIPINE	SAN
02385791	AMLODIPINE	SIV
02429217	AMLODIPINE	JMP
02419564	AMLODIPINE BESYLATE	ACC
02273373	APO-AMLODIPINE	APX
02397072	AURO-AMLODIPINE	AUR
02392135	BIO-AMLODIPINE	BMI
02326833	DOM-AMLODIPINE	DPC
02280132	GD-AMLODIPINE	PFI
02357194	JAMP-AMLODIPINE	JMP
02468026	M-AMLODIPINE	MAN
02371715	MAR-AMLODIPINE	MAR
02362651	MINT-AMLODIPINE	MIN
02272113	MYLAN-AMLODIPINE	MYL
00878928	NORVASC	PFI
02476460	NRA-AMLODIPINE	UNK
02469030	PHARMA-AMLODIPINE	PMS
02284065	PMS-AMLODIPINE	PMS
02321858	RAN-AMLODIPINE	RBY
02331497	RIVA-AMLODIPINE	RIV
02284383	SANDOZ AMLODIPINE	SDZ
02357712	SEPTA-AMLODIPINE	SPT
02250497	TEVA-AMLODIPINE	TEV
02426986	VAN-AMLODIPINE	VAN

ST 10MG TABLET

02297493	ACT AMLODIPINE	ACG
02369249	AG-AMLODIPINE	ANG
02326817	AMLODIPINE	PDL
02331292	AMLODIPINE	SAN
02385805	AMLODIPINE	SIV
02429225	AMLODIPINE	JMP
02419572	AMLODIPINE BESYLATE	ACC
02273381	APO-AMLODIPINE	APX
02397080	AURO-AMLODIPINE	AUR
02392143	BIO-AMLODIPINE	BMI
02326841	DOM-AMLODIPINE	DPC
02280140	GD-AMLODIPINE	PFI
02357208	JAMP-AMLODIPINE	JMP
02468034	M-AMLODIPINE	MAN
02371723	MAR-AMLODIPINE	MAR
02362678	MINT-AMLODIPINE	MIN
02272121	MYLAN-AMLODIPINE	MYL
00878936	NORVASC	PFI
02476479	NRA-AMLODIPINE	UNK
02469049	PHARMA-AMLODIPINE	PMS
02284073	PMS-AMLODIPINE	PMS

24:28.08 DIHYDROPYRIDINES

AMLODIPINE BESYLATE

ST 10MG TABLET

02321866	RAN-AMLODIPINE	RBY
02331500	RIVA-AMLODIPINE	RIV
02284391	SANDOZ AMLODIPINE	SDZ
02357720	SEPTA-AMLODIPINE	SPT
02250500	TEVA-AMLODIPINE	TEV
02426994	VAN-AMLODIPINE	VAN

ST PDIN FOR EXTEMPORANEOUS MIXTURE

99503003	AMLODIPINE ORAL LIQUID	UNK
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AMLODIPINE BESYLATE, ATORVASTATIN CALCIUM

ST 5MG & 10MG TABLET

02411253	APO-AMLODIPINE-ATORVASTATIN	APX
02273233	CADUET	PFI
02362759	GD-AMLODIPINE-ATORVASTATIN	PFI
02404222	PMS-AMLODIPINE-ATORVASTATIN	PMS

ST 5MG & 20MG TABLET

02411261	APO-AMLODIPINE-ATORVASTATIN	APX
02273241	CADUET	PFI
02362767	GD-AMLODIPINE-ATORVASTATIN	PFI
02404230	PMS-AMLODIPINE-ATORVASTATIN	PMS

ST 5MG & 40MG TABLET

02411288	APO-AMLODIPINE-ATORVASTATIN	APX
02273268	CADUET	PFI
02362775	GD-AMLODIPINE-ATORVASTATIN	PFI

ST 5MG & 80MG TABLET

02411296	APO-AMLODIPINE-ATORVASTATIN	APX
02273276	CADUET	PFI
02362783	GD-AMLODIPINE-ATORVASTATIN	PFI

ST 10MG & 10MG TABLET

02411318	APO-AMLODIPINE-ATORVASTATIN	APX
02273284	CADUET	PFI
02362791	GD-AMLODIPINE-ATORVASTATIN	PFI
02404249	PMS-AMLODIPINE-ATORVASTATIN	PMS

ST 10MG & 20MG TABLET

02411326	APO-AMLODIPINE-ATORVASTATIN	APX
02273292	CADUET	PFI
02362805	GD-AMLODIPINE-ATORVASTATIN	PFI
02404257	PMS-AMLODIPINE-ATORVASTATIN	PMS

ST 10MG & 40MG TABLET

02411334	APO-AMLODIPINE-ATORVASTATIN	APX
02273306	CADUET	PFI
02362813	GD-AMLODIPINE-ATORVASTATIN	PFI

ST 10MG & 80MG TABLET

02411342	APO-AMLODIPINE-ATORVASTATIN	APX
02273314	CADUET	PFI
02362821	GD-AMLODIPINE-ATORVASTATIN	PFI

AMLODIPINE BESYLATE, TELMISARTAN

ST 5MG & 40MG TABLET

02371022	TWYNSTA	BOE
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ST 5MG & 80MG TABLET

02371049	TWYNSTA	BOE
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ST 10MG & 40MG TABLET

02371030	TWYNSTA	BOE
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24:28.08 DIHYDROPYRIDINES

AMLODIPINE BESYLATE, TELMISARTAN

ST **10MG & 80MG TABLET**
02371057 TWYNSTA BOE

FELODIPINE

ST **2.5MG TABLET (EXTENDED RELEASE)**
02452367 APO-FELODIPINE APX
02057778 PLENDIL AZC

ST **5MG TABLET (EXTENDED RELEASE)**
02452375 APO-FELODIPINE APX
00851779 PLENDIL AZC
02280264 SANDOZ FELODIPINE SDZ
09857203 SANDOZ-FELODIPINE SDZ

ST **10MG TABLET (EXTENDED RELEASE)**
02452383 APO-FELODIPINE APX
00851787 PLENDIL AZC
02280272 SANDOZ FELODIPINE SDZ
09857204 SANDOZ-FELODIPINE SDZ

NIFEDIPINE

ST **5MG CAPSULE**
00725110 NIFEDIPINE AAP
02235897 PMS-NIFEDIPINE PMS

ST **10MG CAPSULE**
00755907 NIFEDIPINE AAP
02235898 PMS-NIFEDIPINE PMS

ST **20MG TABLET (EXTENDED RELEASE)**
02237618 ADALAT XL BAY

ST **30MG TABLET (EXTENDED RELEASE)**
02155907 ADALAT XL BAY
02349167 MYLAN-NIFEDIPINE MYL
02421631 NIFEDIPINE PDL
02418630 PMS-NIFEDIPINE PMS

ST **60MG TABLET (EXTENDED RELEASE)**
02155990 ADALAT XL BAY
02321149 MYLAN-NIFEDIPINE MYL
02421658 NIFEDIPINE PDL
02416301 PMS-NIFEDIPINE PMS

NIMODIPINE

ST **30MG TABLET**
02325926 NIMOTOP BAY

24:28.92 MISCELLANEOUS CALCIUM-CHANNEL BLOCKING AGENTS

DILTIAZEM HYDROCHLORIDE

ST **120MG CAPSULE (CONTROLLED DELIVERY)**
02230997 APO-DILTIAZ CD APX
02231472 DILTIAZEM CD PDL
02400421 DILTIAZEM CD SAN
02355752 PMS-DILTIAZEM CD PMS

ST **180MG CAPSULE (CONTROLLED DELIVERY)**
02230998 APO-DILTIAZ CD APX
02231474 DILTIAZEM CD PDL
02400448 DILTIAZEM CD SAN
02355760 PMS-DILTIAZEM CD PMS

24:28.92 MISCELLANEOUS CALCIUM-CHANNEL BLOCKING AGENTS

DILTIAZEM HYDROCHLORIDE

ST **240MG CAPSULE (CONTROLLED DELIVERY)**
02230999 APO-DILTIAZ CD APX
02231475 DILTIAZEM CD PDL
02400456 DILTIAZEM CD SAN
02355779 PMS-DILTIAZEM CD PMS

ST **300MG CAPSULE (CONTROLLED DELIVERY)**
02229526 APO-DILTIAZ CD APX
02231057 DILTIAZEM CD PDL
02400464 DILTIAZEM CD SAN
02355787 PMS-DILTIAZEM CD PMS

ST **120MG CAPSULE (EXTENDED RELEASE)**
02370611 ACT DILTIAZEM CD TEV
02370441 ACT DILTIAZEM T ACG
02097249 CARDIZEM CD VAE
02445999 DILTIAZEM CD SIV
02325306 DILTIAZEM TZ PDL
02465353 MAR-DILTIAZEM T MAR
02243338 SANDOZ DILTIAZEM CD SDZ
02245918 SANDOZ DILTIAZEM T SDZ
02271605 TEVA-DILTIAZEM VAE
02242538 TEVA-DILTIAZEM CD TEV
02231150 TIAZAC VAE

ST **180MG CAPSULE (EXTENDED RELEASE)**
02370638 ACT DILTIAZEM CD TEV
02370492 ACT DILTIAZEM T ACG
02097257 CARDIZEM CD VAE
02446006 DILTIAZEM CD SIV
02325314 DILTIAZEM TZ PDL
02465361 MAR-DILTIAZEM T MAR
02243339 SANDOZ DILTIAZEM CD SDZ
02245919 SANDOZ DILTIAZEM T SDZ
02271613 TEVA-DILTIAZEM VAE
02242539 TEVA-DILTIAZEM CD TEV
02231151 TIAZAC VAE

ST **240MG CAPSULE (EXTENDED RELEASE)**
02370646 ACT DILTIAZEM CD TEV
02370506 ACT DILTIAZEM T ACG
02097265 CARDIZEM CD VAE
02446014 DILTIAZEM CD SIV
02325322 DILTIAZEM TZ PDL
02465388 MAR-DILTIAZEM T MAR
02243340 SANDOZ DILTIAZEM CD SDZ
02245920 SANDOZ DILTIAZEM T SDZ
02271621 TEVA-DILTIAZEM VAE
02242540 TEVA-DILTIAZEM CD TEV
02231152 TIAZAC VAE

ST **300MG CAPSULE (EXTENDED RELEASE)**
02370654 ACT DILTIAZEM CD TEV
02370514 ACT DILTIAZEM T ACG
02097273 CARDIZEM CD VAE
02446022 DILTIAZEM CD SIV
02325330 DILTIAZEM TZ PDL
02465396 MAR-DILTIAZEM T MAR
02243341 SANDOZ DILTIAZEM CD SDZ

24:28.92 MISCELLANEOUS CALCIUM-CHANNEL BLOCKING AGENTS

DILTIAZEM HYDROCHLORIDE

ST 300MG CAPSULE (EXTENDED RELEASE)		
02245921	SANDOZ DILTIAZEM T	SDZ
02271648	TEVA-DILTIAZEM	VAE
02242541	TEVA-DILTIAZEM CD	TEV
02231154	TIAZAC	VAE
ST 360MG CAPSULE (EXTENDED RELEASE)		
02370522	ACT DILTIAZEM T	ACG
02325349	DILTIAZEM TZ	PDL
02465418	MAR-DILTIAZEM T	MAR
02245922	SANDOZ DILTIAZEM T	SDZ
02271656	TEVA-DILTIAZEM	VAE
02231155	TIAZAC	VAE
ST 30MG TABLET		
00771376	APO-DILTIAZ	APX
00862924	TEVA-DILTIAZEM	TEV
ST 60MG TABLET		
00771384	APO-DILTIAZ	APX
00862932	TEVA-DILTIAZEM	TEV
ST 120MG TABLET (EXTENDED RELEASE)		
02256738	TIAZAC XC	VAE
ST 180MG TABLET (EXTENDED RELEASE)		
02256746	TIAZAC XC	VAE
ST 240MG TABLET (EXTENDED RELEASE)		
02256754	TIAZAC XC	VAE
ST 300MG TABLET (EXTENDED RELEASE)		
02256762	TIAZAC XC	VAE
ST 360MG TABLET (EXTENDED RELEASE)		
02256770	TIAZAC XC	VAE

VERAPAMIL HYDROCHLORIDE

120MG CAPSULE (SUSTAINED RELEASE)		
02100479	VERELAN	RGL
ST 180MG CAPSULE (SUSTAINED RELEASE)		
02100487	VERELAN	RGL
ST 240MG CAPSULE (SUSTAINED RELEASE)		
02100495	VERELAN	RGL
ST 80MG TABLET		
00782483	APO-VERAP	APX
02237921	MYLAN-VERAPAMIL	MYL
ST 120MG TABLET		
00782491	APO-VERAP	APX
02237922	MYLAN-VERAPAMIL	MYL
ST 120MG TABLET (EXTENDED RELEASE)		
02246893	APO-VERAP SR	APX
01907123	ISOPTIN SR	BGP
02210347	MYLAN-VERAPAMIL SR	MYL
ST 180MG TABLET (EXTENDED RELEASE)		
02246894	APO-VERAP SR	APX
01934317	ISOPTIN SR	BGP
02450488	MYLAN-VERAPAMIL	MYL
ST 240MG TABLET (EXTENDED RELEASE)		
02246895	APO-VERAP SR	APX
02240321	DOM-VERAPAMIL SR	DPC
00742554	ISOPTIN SR	BGP

24:28.92 MISCELLANEOUS CALCIUM-CHANNEL BLOCKING AGENTS

VERAPAMIL HYDROCHLORIDE

ST 240MG TABLET (EXTENDED RELEASE)		
02450496	MYLAN-VERAPAMIL	MYL
02237791	PMS-VERAPAMIL SR	PMS
02248082	RIVA-VERAPAMIL SR	RIV

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

BENAZEPRIL HYDROCHLORIDE

ST 5MG TABLET		
02290332	BENAZEPRIL	AAP
ST 10MG TABLET		
02290340	BENAZEPRIL	AAP
ST 20MG TABLET		
02273918	BENAZEPRIL	AAP

CAPTOPRIL

ST 6.25MG TABLET		
01999559	APO-CAPTO	APX
ST 12.5MG TABLET		
00893595	APO-CAPTO	APX
01942964	TEVA-CAPTOPRIL	TEV
ST 25MG TABLET		
00893609	APO-CAPTO	APX
01942972	TEVA-CAPTOPRIL	TEV
ST 50MG TABLET		
00893617	APO-CAPTO	APX
01942980	TEVA-CAPTOPRIL	TEV
ST 100MG TABLET		
00893625	APO-CAPTO	APX
02230206	PMS-CAPTOPRIL	PMS
01942999	TEVA-CAPTOPRIL	TEV

CILAZAPRIL

ST 1MG TABLET		
02291134	APO-CILAZAPRIL	APX
02283778	MYLAN-CILAZAPRIL	MYL
02280442	PMS-CILAZAPRIL	PMS
ST 2.5MG TABLET		
02291142	APO-CILAZAPRIL	APX
01911473	INHIBACE	HLR
02283786	MYLAN-CILAZAPRIL	MYL
02280450	PMS-CILAZAPRIL	PMS
ST 5MG TABLET		
02291150	APO-CILAZAPRIL	APX
01911481	INHIBACE	HLR
02283794	MYLAN-CILAZAPRIL	MYL
02280469	PMS-CILAZAPRIL	PMS

CILAZAPRIL, HYDROCHLOROTHIAZIDE

ST 5MG & 12.5MG TABLET		
02284987	APO-CILAZAPRIL/HCTZ	APX
02181479	INHIBACE PLUS	CHE
02313731	TEVA-CILAZAPRIL/HCTZ	TEV

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

ENALAPRIL MALEATE

ST **2.5MG TABLET**

02291878	ACT ENALAPRIL	TEV
02020025	APO-ENALAPRIL	APX
02400650	ENALAPRIL	SAN
02442957	ENALAPRIL	SIV
02459450	MAR-ENALAPRIL	MAR
02300036	MYLAN-ENALAPRIL	MYL
02311402	PRO-ENALAPRIL	PDL
02352230	RAN-ENALAPRIL	RBY
02300796	RIVA-ENALAPRIL	RIV
02299933	SANDOZ ENALAPRIL	SDZ
02300117	TARO-ENALAPRIL	TAR

ST **5MG TABLET**

02291886	ACT ENALAPRIL	TEV
02019884	APO-ENALAPRIL	APX
02400669	ENALAPRIL	SAN
02442965	ENALAPRIL	SIV
02459469	MAR-ENALAPRIL	MAR
02300044	MYLAN-ENALAPRIL	MYL
02311410	PRO-ENALAPRIL	PDL
02352249	RAN-ENALAPRIL	RBY
02300818	RIVA-ENALAPRIL	RIV
02299941	SANDOZ ENALAPRIL	SDZ
02300125	TARO-ENALAPRIL	TAR
02233005	TEVA-ENALAPRIL	TEV
00708879	VASOTEC	FRS

ST **10MG TABLET**

02291894	ACT ENALAPRIL	TEV
02019892	APO-ENALAPRIL	APX
02400677	ENALAPRIL	SAN
02442973	ENALAPRIL	SIV
02444771	MAR-ENALAPRIL	IDE
02300052	MYLAN-ENALAPRIL	MYL
02311429	PRO-ENALAPRIL	PDL
02352257	RAN-ENALAPRIL	RBY
02300826	RIVA-ENALAPRIL	RIV
02299968	SANDOZ ENALAPRIL	SDZ
02300133	TARO-ENALAPRIL	TAR
02233006	TEVA-ENALAPRIL	TEV
00670901	VASOTEC	FRS

ST **20MG TABLET**

02291908	ACT ENALAPRIL	TEV
02019906	APO-ENALAPRIL	APX
02400685	ENALAPRIL	SAN
02442981	ENALAPRIL	SIV
02444798	MAR-ENALAPRIL	IDE
02300060	MYLAN-ENALAPRIL	MYL
02311437	PRO-ENALAPRIL	PDL
02352265	RAN-ENALAPRIL	RBY
02300834	RIVA-ENALAPRIL	RIV
02299976	SANDOZ ENALAPRIL	SDZ
02300141	TARO-ENALAPRIL	TAR
02233007	TEVA-ENALAPRIL	TEV
00670928	VASOTEC	FRS

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

ENALAPRIL MALEATE

ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503013	ENALAPRIL ORAL LIQUID	UNK
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ENALAPRIL MALEATE, HYDROCHLOROTHIAZIDE

ST **5MG & 12.5MG TABLET**

02352923	ENALAPRIL MALEATE/HCTZ	AAP
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ST **10MG & 25MG TABLET**

02352931	ENALAPRIL MALEATE/HCTZ	AAP
00657298	VASERETIC	FRS

FOSINOPRIL SODIUM

ST **10MG TABLET**

02266008	APO-FOSINOPRIL	APX
02303000	FOSINOPRIL	PDL
02332566	FOSINOPRIL	RBY
02459388	FOSINOPRIL	SAN
02331004	JAMP-FOSINOPRIL	JMP
02255944	PMS-FOSINOPRIL	PMS
02294524	RAN-FOSINOPRIL	RBY
02247802	TEVA-FOSINOPRIL	TEV

ST **20MG TABLET**

02266016	APO-FOSINOPRIL	APX
02303019	FOSINOPRIL	PDL
02332574	FOSINOPRIL	RBY
02459396	FOSINOPRIL	SAN
02331012	JAMP-FOSINOPRIL	JMP
02255952	PMS-FOSINOPRIL	PMS
02294532	RAN-FOSINOPRIL	RBY
02247803	TEVA-FOSINOPRIL	TEV

LISINOPRIL

ST **5MG TABLET**

02217481	APO-LISINOPRIL	APX
09853685	APO-LISINOPRIL	APX
02394472	AURO-LISINOPRIL	AUR
02361531	JAMP-LISINOPRIL	JMP
02386232	LISINOPRIL	SIV
02292203	PMS-LISINOPRIL	PMS
02310961	PRO-LISINOPRIL	PDL
02294230	RAN-LISINOPRIL	RBY
02289199	SANDOZ LISINOPRIL	SDZ
02285061	TEVA-LISINOPRIL (TYPE P)	TEV
02285118	TEVA-LISINOPRIL (TYPE Z)	TEV
02049333	ZESTRIL	AZC

ST **10MG TABLET**

02217503	APO-LISINOPRIL	APX
09853960	APO-LISINOPRIL	APX
02394480	AURO-LISINOPRIL	AUR
02361558	JAMP-LISINOPRIL	JMP
02386240	LISINOPRIL	SIV
02292211	PMS-LISINOPRIL	PMS
00839396	PRINIVIL	FRS
02310988	PRO-LISINOPRIL	PDL
02294249	RAN-LISINOPRIL	RBY

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

LISINOPRIL

ST **10MG TABLET**

02289202	SANDOZ LISINOPRIL	SDZ
02285088	TEVA-LISINOPRIL (TYPE P)	TEV
02285126	TEVA-LISINOPRIL (TYPE Z)	TEV
02049376	ZESTRIL	AZC

ST **20MG TABLET**

02217511	APO-LISINOPRIL	APX
09854010	APO-LISINOPRIL	APX
02394499	AURO-LISINOPRIL	AUR
02361566	JAMP-LISINOPRIL	JMP
02386259	LISINOPRIL	SIV
02292238	PMS-LISINOPRIL	PMS
00839418	PRINIVIL	FRS
02310996	PRO-LISINOPRIL	PDL
02294257	RAN-LISINOPRIL	RBV
02289229	SANDOZ LISINOPRIL	SDZ
02285096	TEVA-LISINOPRIL (TYPE P)	TEV
02285134	TEVA-LISINOPRIL (TYPE Z)	TEV
02049384	ZESTRIL	AZC

LISINOPRIL, HYDROCHLOROTHIAZIDE

ST **10MG & 12.5MG TABLET**

02362945	LISINOPRIL/HCTZ (TYPE Z)	SAN
02302365	SANDOZ LISINOPRIL HCT	SDZ
02302136	TEVA-LISINOPRIL/HCTZ (TYPE P)	TEV
02301768	TEVA-LISINOPRIL/HCTZ (TYPE Z)	TEV
02103729	ZESTORETIC	AZC

ST **20MG & 12.5MG TABLET**

02362953	LISINOPRIL/HCTZ (TYPE Z)	SAN
02302373	SANDOZ LISINOPRIL HCT	SDZ
02302144	TEVA-LISINOPRIL/HCTZ (TYPE P)	TEV
02301776	TEVA-LISINOPRIL/HCTZ (TYPE Z)	TEV
02045737	ZESTORETIC	AZC

ST **20MG & 25MG TABLET**

02362961	LISINOPRIL/HCTZ (TYPE Z)	SAN
02302381	SANDOZ LISINOPRIL HCT	SDZ
02302152	TEVA-LISINOPRIL/HCTZ (TYPE P)	TEV
02301784	TEVA-LISINOPRIL/HCTZ (TYPE Z)	TEV
02045729	ZESTORETIC	AZC

PERINDOPRIL ERBUMINE

ST **2MG TABLET**

02289261	APO-PERINDOPRIL	APX
02459817	AURO-PERINDOPRIL	AUR
02123274	COVERSYL	SEV
02476762	MINT-PERINDOPRIL	MIN
02479877	PERINDOPRIL ERBUMINE	SIV
02470675	PMS-PERINDOPRIL	PMS
02472015	RIVA-PERINDOPRIL	RIV
02470225	SANDOZ PERINDOPRIL ERBUMINE	SDZ
02464985	TEVA-PERINDOPRIL	TEV

ST **4MG TABLET**

02289288	APO-PERINDOPRIL	APX
02459825	AURO-PERINDOPRIL	AUR
02123282	COVERSYL	SEV

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

PERINDOPRIL ERBUMINE

ST **4MG TABLET**

02477017	JAMP PERINDOPRIL	JMP
02476770	MINT-PERINDOPRIL	MIN
02479885	PERINDOPRIL ERBUMINE	SIV
02470683	PMS-PERINDOPRIL	PMS
02472023	RIVA-PERINDOPRIL	RIV
02470233	SANDOZ PERINDOPRIL ERBUMINE	SDZ
02464993	TEVA-PERINDOPRIL	TEV

ST **8MG TABLET**

02289296	APO-PERINDOPRIL	APX
02459833	AURO-PERINDOPRIL	AUR
02246624	COVERSYL	SEV
02477025	JAMP PERINDOPRIL	JMP
02476789	MINT-PERINDOPRIL	MIN
02479893	PERINDOPRIL ERBUMINE	SIV
02470691	PMS-PERINDOPRIL	PMS
02472031	RIVA-PERINDOPRIL	RIV
02470241	SANDOZ PERINDOPRIL ERBUMINE	SDZ
02465000	TEVA-PERINDOPRIL	TEV

PERINDOPRIL ERBUMINE, INDAPAMIDE

ST **4MG & 1.25MG TABLET**

02246569	COVERSYL PLUS	SEV
02470438	SANDOZ PERINDOPRIL ERBUMINE/ INDAPAMIDE	SDZ
02464020	TEVA-PERINDOPRIL/INDAPAMIDE	TEV

ST **8MG & 2.5MG TABLET**

02453061	APO-PERINDOPRIL-INDAPAMIDE	APX
02321653	COVERSYL PLUS HD	SEV
02408201	MYLAN-PERINDOPRIL/INDAPAMIDE	MYL
02470446	SANDOZ PERINDOPRIL ERBUMINE/ INDAPAMIDE HD	SDZ
02464039	TEVA-PERINDOPRIL/INDAPAMIDE	TEV

QUINAPRIL

ST **5MG TABLET**

01947664	ACCUPRIL	PFI
02248499	APO-QUINAPRIL	APX
02340550	PMS-QUINAPRIL	PMS

ST **10MG TABLET**

01947672	ACCUPRIL	PFI
02248500	APO-QUINAPRIL	APX
02290995	GD-QUINAPRIL	PFI
02340569	PMS-QUINAPRIL	PMS

ST **20MG TABLET**

01947680	ACCUPRIL	PFI
02248501	APO-QUINAPRIL	APX
02291002	GD-QUINAPRIL	PFI
02340577	PMS-QUINAPRIL	PMS

ST **40MG TABLET**

01947699	ACCUPRIL	PFI
02248502	APO-QUINAPRIL	APX
02340585	PMS-QUINAPRIL	PMS

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

QUINAPRIL, HYDROCHLOROTHIAZIDE

ST 10MG & 12.5MG TABLET

02237367	ACCURETIC	PFI
02408767	APO-QUINAPRIL/HCTZ	APX
02473291	AURO-QUINAPRIL HCTZ	AUR

ST 20MG & 12.5MG TABLET

02237368	ACCURETIC	PFI
02408775	APO-QUINAPRIL/HCTZ	APX
02473305	AURO-QUINAPRIL HCTZ	AUR

ST 20MG & 25MG TABLET

02237369	ACCURETIC	PFI
02408783	APO-QUINAPRIL/HCTZ	APX
02473321	AURO-QUINAPRIL HCTZ	AUR

RAMIPRIL

ST 1.25MG CAPSULE

02295482	ACT RAMIPRIL	ACG
02221829	ALTACE	VAE
02251515	APO-RAMIPRIL	APX
02387387	AURO-RAMIPRIL	AUR
02331101	JAMP-RAMIPRIL	JMP
02420457	MAR-RAMIPRIL	MAR
02469057	PHARMA-RAMIPRIL	PMS
02295369	PMS-RAMIPRIL	PMS
02310023	PRO-RAMIPRIL	PDL
02299372	RAMIPRIL	RIV
02308363	RAMIPRIL	SIV
02310503	RAN-RAMIPRIL	RBV
02438860	VAN-RAMIPRIL	VAN

ST 2.5MG CAPSULE

02295490	ACT RAMIPRIL	ACG
02221837	ALTACE	VAE
02251531	APO-RAMIPRIL	APX
02387395	AURO-RAMIPRIL	AUR
02287951	DOM-RAMIPRIL	DPC
02331128	JAMP-RAMIPRIL	JMP
02420465	MAR-RAMIPRIL	MAR
02421305	MINT-RAMIPRIL	MIN
02469065	PHARMA-RAMIPRIL	PMS
02247917	PMS-RAMIPRIL	PMS
02310066	PRO-RAMIPRIL	PDL
02255316	RAMIPRIL	RIV
02287927	RAMIPRIL	SIV
02374846	RAMIPRIL	SAN
02310511	RAN-RAMIPRIL	RBV
02247945	TEVA-RAMIPRIL	TEV
02438879	VAN-RAMIPRIL	VAN

ST 5MG CAPSULE

02295504	ACT RAMIPRIL	ACG
02221845	ALTACE	VAE
02251574	APO-RAMIPRIL	APX
02387409	AURO-RAMIPRIL	AUR
02287978	DOM-RAMIPRIL	DPC
02331136	JAMP-RAMIPRIL	JMP
02420473	MAR-RAMIPRIL	MAR

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

RAMIPRIL

ST 5MG CAPSULE

02421313	MINT-RAMIPRIL	MIN
02469073	PHARMA-RAMIPRIL	PMS
02247918	PMS-RAMIPRIL	PMS
02310074	PRO-RAMIPRIL	PDL
02255324	RAMIPRIL	RIV
02287935	RAMIPRIL	SIV
02374854	RAMIPRIL	SAN
02310538	RAN-RAMIPRIL	RBV
02247946	TEVA-RAMIPRIL	TEV
02438887	VAN-RAMIPRIL	VAN

ST 10MG CAPSULE

02295512	ACT RAMIPRIL	ACG
02221853	ALTACE	VAE
02251582	APO-RAMIPRIL	APX
02387417	AURO-RAMIPRIL	AUR
02287986	DOM-RAMIPRIL	DPC
02331144	JAMP-RAMIPRIL	JMP
02420481	MAR-RAMIPRIL	MAR
02421321	MINT-RAMIPRIL	MIN
02469081	PHARMA-RAMIPRIL	PMS
02247919	PMS-RAMIPRIL	PMS
02310104	PRO-RAMIPRIL	PDL
02255332	RAMIPRIL	RIV
02287943	RAMIPRIL	SIV
02374862	RAMIPRIL	SAN
02310546	RAN-RAMIPRIL	RBV
02247947	TEVA-RAMIPRIL	TEV
02438895	VAN-RAMIPRIL	VAN

ST 15MG CAPSULE

02325381	APO-RAMIPRIL	APX
02440334	JAMP-RAMIPRIL	JMP
02420503	MAR-RAMIPRIL	MAR
02421348	MINT-RAMIPRIL	MIN
02343932	PMS-RAMIPRIL	PMS
02425548	RAN-RAMIPRIL	RBV
02438909	VAN-RAMIPRIL	VAN

ST 1.25MG TABLET

02291398	SANDOZ RAMIPRIL	SDZ
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ST 2.5MG TABLET

02291401	SANDOZ RAMIPRIL	SDZ
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ST 5MG TABLET

02291428	SANDOZ RAMIPRIL	SDZ
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ST 10MG TABLET

02291436	SANDOZ RAMIPRIL	SDZ
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RAMIPRIL, HYDROCHLOROTHIAZIDE

ST 2.5MG & 12.5MG TABLET

02283131	ALTACE HCT	VAE
02354004	APO-RAMIPRIL/HCTZ	APX
02449439	RAN-RAMIPRIL HCTZ	RBV

ST 5MG & 12.5MG TABLET

02283158	ALTACE HCT	VAE
02354012	APO-RAMIPRIL/HCTZ	APX

24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS

RAMIPRIL, HYDROCHLOROTHIAZIDE

ST **5MG & 12.5MG TABLET**

02449447 RAN-RAMIPRIL HCTZ RBY

ST **5MG & 25MG TABLET**

02283174 ALTACE HCT VAE

02354020 APO-RAMIPRIL/HCTZ APX

02449463 RAN-RAMIPRIL HCTZ RBY

ST **10MG & 12.5MG TABLET**

02283166 ALTACE HCT VAE

02342154 PMS-RAMIPRIL-HCTZ PMS

02415895 RAMIPRIL-HCTZ PDL

02449455 RAN-RAMIPRIL HCTZ RBY

ST **10MG & 25MG TABLET**

02283182 ALTACE HCT VAE

02354039 APO-RAMIPRIL/HCTZ APX

02342170 PMS-RAMIPRIL-HCTZ PMS

02415909 RAMIPRIL-HCTZ PDL

02449471 RAN-RAMIPRIL HCTZ RBY

TRANDOLAPRIL

ST **0.5MG CAPSULE**

02471868 AURO-TRANDOLAPRIL AUR

02231457 MAVIK BGP

02357755 PMS-TRANDOLAPRIL PMS

02325721 SANDOZ TRANDOLAPRIL SDZ

02415429 TEVA-TRANDOLAPRIL TEV

ST **1MG CAPSULE**

02471876 AURO-TRANDOLAPRIL AUR

02231459 MAVIK BGP

02357763 PMS-TRANDOLAPRIL PMS

02325748 SANDOZ TRANDOLAPRIL SDZ

02415437 TEVA-TRANDOLAPRIL TEV

ST **2MG CAPSULE**

02471884 AURO-TRANDOLAPRIL AUR

02231460 MAVIK BGP

02357771 PMS-TRANDOLAPRIL PMS

02325756 SANDOZ TRANDOLAPRIL SDZ

02415445 TEVA-TRANDOLAPRIL TEV

ST **4MG CAPSULE**

02471892 AURO-TRANDOLAPRIL AUR

02239267 MAVIK BGP

02357798 PMS-TRANDOLAPRIL PMS

02325764 SANDOZ TRANDOLAPRIL SDZ

02415453 TEVA-TRANDOLAPRIL TEV

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

AZILSARTAN MEDOXOMIL

ST **40MG TABLET**

02381389 EDARBI VAE

ST **80MG TABLET**

02381397 EDARBI VAE

CANDESARTAN CILEXETIL

ST **4MG TABLET**

02379260 ACH-CANDESARTAN ACC

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

CANDESARTAN CILEXETIL

ST **4MG TABLET**

02376520 ACT CANDESARTAN ACG

02365340 APO-CANDESARTAN APX

02239090 ATACAND AZC

02445786 AURO-CANDESARTAN AUR

02388901 CANDESARTAN SAN

02386496 JAMP-CANDESARTAN JMP

02391171 PMS-CANDESARTAN PMS

02380684 RAN-CANDESARTAN RBY

02326957 SANDOZ CANDESARTAN SDZ

ST **8MG TABLET**

02463768 ACCEL-CANDESARTAN ACP

02379279 ACH-CANDESARTAN ACC

02376539 ACT CANDESARTAN ACG

02365359 APO-CANDESARTAN APX

02239091 ATACAND AZC

02445794 AURO-CANDESARTAN AUR

02377934 CANDESARTAN PDL

02388707 CANDESARTAN SIV

02388928 CANDESARTAN SAN

02386518 JAMP-CANDESARTAN JMP

02476916 MINT-CANDESARTAN MIN

02391198 PMS-CANDESARTAN PMS

02380692 RAN-CANDESARTAN RBY

02326965 SANDOZ CANDESARTAN SDZ

02366312 TEVA-CANDESARTAN TEV

ST **16MG TABLET**

02463776 ACCEL-CANDESARTAN ACP

02379287 ACH-CANDESARTAN ACC

02376547 ACT CANDESARTAN ACG

02365367 APO-CANDESARTAN APX

02239092 ATACAND AZC

02445808 AURO-CANDESARTAN AUR

02377942 CANDESARTAN PDL

02388715 CANDESARTAN SIV

02388936 CANDESARTAN SAN

02386526 JAMP-CANDESARTAN JMP

02476924 MINT-CANDESARTAN MIN

02391201 PMS-CANDESARTAN PMS

02380706 RAN-CANDESARTAN RBY

02326973 SANDOZ CANDESARTAN SDZ

02366320 TEVA-CANDESARTAN TEV

ST **32MG TABLET**

02463784 ACCEL-CANDESARTAN ACP

02379295 ACH-CANDESARTAN ACC

02376555 ACT CANDESARTAN ACG

02399105 APO-CANDESARTAN APX

02311658 ATACAND AZC

02445816 AURO-CANDESARTAN AUR

02422069 CANDESARTAN PDL

02435845 CANDESARTAN SAN

02386534 JAMP-CANDESARTAN JMP

02391228 PMS-CANDESARTAN PMS

02380714 RAN-CANDESARTAN RBY

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

CANDESARTAN CILEXETIL

ST **32MG TABLET**

02417340	SANDOZ CANDESARTAN	SDZ
02366339	TEVA-CANDESARTAN	TEV

CANDESARTAN CILEXETIL, HYDROCHLOROTHIAZIDE

ST **16MG & 12.5MG TABLET**

02463865	ACCEL-CANDESARTAN/HCTZ	ACP
02367866	APO-CANDESARTAN/HCTZ	APX
02244021	ATACAND PLUS	AZC
02421038	AURO-CANDESARTAN HCT	AUR
02394812	CANDESARTAN-HCT	SIV
02392275	CANDESARTAN-HCTZ	PDL
02394804	CANDESARTAN-HCTZ	SAN
02391295	PMS-CANDESARTAN HCTZ	PMS
02327902	SANDOZ CANDESARTAN PLUS	SDZ
02395541	TEVA-CANDESARTAN/HCTZ	TEV

ST **32MG & 12.5MG TABLET**

02463849	ACCEL-CANDESARTAN/HCTZ	ACP
02395126	APO-CANDESARTAN/HCTZ	APX
02332922	ATACAND PLUS	AZC
02421046	AURO-CANDESARTAN HCT	AUR
02420732	SANDOZ CANDESARTAN PLUS	SDZ
02395568	TEVA-CANDESARTAN/HCTZ	TEV

ST **32MG & 25MG TABLET**

02463857	ACCEL-CANDESARTAN/HCTZ	ACP
02395134	APO-CANDESARTAN/HCTZ	APX
02332957	ATACAND PLUS	AZC
02421054	AURO-CANDESARTAN HCT	AUR
02420740	SANDOZ CANDESARTAN PLUS	SDZ

EPOSARTAN MESYLATE

ST **400MG TABLET**

02240432	TEVETEN	BGP
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ST **600MG TABLET**

02243942	TEVETEN	BGP
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EPOSARTAN MESYLATE, HYDROCHLOROTHIAZIDE

ST **600MG & 12.5MG TABLET**

02253631	TEVETEN PLUS	BGP
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IRBESARTAN

75MG TABLET

02474395	AG-IRBESARTAN	ANG
02386968	APO-IRBESARTAN	APX
02406098	AURO-IRBESARTAN	AUR
02237923	AVAPRO	SAC
02446146	BIO-IRBESARTAN	BMI
02365197	IRBESARTAN	PDL
02372347	IRBESARTAN	SAN
02385287	IRBESARTAN	SIV
02418193	JAMP-IRBESARTAN	JMP
02422980	MINT-IRBESARTAN	MIN
02317060	PMS-IRBESARTAN	PMS
02406810	RAN-IRBESARTAN	RBV

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

IRBESARTAN

75MG TABLET

02328461	SANDOZ IRBESARTAN	SDZ
02316390	TEVA-IRBESARTAN	TEV
02427087	VAN-IRBESARTAN	VAN

150MG TABLET

02474409	AG-IRBESARTAN	ANG
02386976	APO-IRBESARTAN	APX
02406101	AURO-IRBESARTAN	AUR
02237924	AVAPRO	SAC
02446154	BIO-IRBESARTAN	BMI
02365200	IRBESARTAN	PDL
02372371	IRBESARTAN	SAN
02385295	IRBESARTAN	SIV
02418207	JAMP-IRBESARTAN	JMP
02422999	MINT-IRBESARTAN	MIN
02317079	PMS-IRBESARTAN	PMS
02406829	RAN-IRBESARTAN	RBV
02328488	SANDOZ IRBESARTAN	SDZ
02316404	TEVA-IRBESARTAN	TEV
02427095	VAN-IRBESARTAN	VAN

300MG TABLET

02474417	AG-IRBESARTAN	ANG
02386984	APO-IRBESARTAN	APX
02406128	AURO-IRBESARTAN	AUR
02237925	AVAPRO	SAC
02446162	BIO-IRBESARTAN	BMI
02365219	IRBESARTAN	PDL
02372398	IRBESARTAN	SAN
02385309	IRBESARTAN	SIV
02418215	JAMP-IRBESARTAN	JMP
02423006	MINT-IRBESARTAN	MIN
02317087	PMS-IRBESARTAN	PMS
02406837	RAN-IRBESARTAN	RBV
02328496	SANDOZ IRBESARTAN	SDZ
02316412	TEVA-IRBESARTAN	TEV
02427109	VAN-IRBESARTAN	VAN

IRBESARTAN, HYDROCHLOROTHIAZIDE

ST **150MG & 12.5MG TABLET**

02387646	APO-IRBESARTAN/HCTZ	APX
02447878	AURO-IRBESARTAN HCT	AUR
02241818	AVALIDE	SAC
02385317	IRBESARTAN HCT	SIV
02372886	IRBESARTAN/HCTZ	SAN
02365162	IRBESARTAN-HCTZ	PDL
02418223	JAMP-IRBESARTAN AND HYDROCHLOROTHIAZIDE	JMP
02392992	MINT-IRBESARTAN/HCTZ	MIN
02328518	PMS-IRBESARTAN-HCTZ	PMS
02363208	RAN-IRBESARTAN HCTZ	RBV
02337428	SANDOZ IRBESARTAN HCT	SDZ
02330512	TEVA-IRBESARTAN HCTZ	TEV

ST **300MG & 12.5MG TABLET**

02387654	APO-IRBESARTAN/HCTZ	APX
02447886	AURO-IRBESARTAN HCT	AUR

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

IRBESARTAN, HYDROCHLOROTHIAZIDE

ST 300MG & 12.5MG TABLET

02241819	AVALIDE	SAC
02385325	IRBESARTAN HCT	SIV
02372894	IRBESARTAN/HCTZ	SAN
02365170	IRBESARTAN-HCTZ	PDL
02418231	JAMP-IRBESARTAN AND HYDROCHLOROTHIAZIDE	JMP
02393018	MINT-IRBESARTAN/HCTZ	MIN
02328526	PMS-IRBESARTAN-HCTZ	PMS
02363216	RAN-IRBESARTAN HCTZ	RBY
02337436	SANDOZ IRBESARTAN HCT	SDZ
02330520	TEVA-IRBESARTAN HCTZ	TEV

ST 300MG & 25MG TABLET

02387662	APO-IRBESARTAN/HCTZ	APX
02447894	AURO-IRBESARTAN HCT	AUR
02385333	IRBESARTAN HCT	SIV
02372908	IRBESARTAN/HCTZ	SAN
02365189	IRBESARTAN-HCTZ	PDL
02418258	JAMP-IRBESARTAN AND HYDROCHLOROTHIAZIDE	JMP
02393026	MINT-IRBESARTAN/HCTZ	MIN
02328534	PMS-IRBESARTAN-HCTZ	PMS
02363224	RAN-IRBESARTAN HCTZ	RBY
02337444	SANDOZ IRBESARTAN HCT	SDZ
02330539	TEVA-IRBESARTAN HCTZ	TEV

LOSARTAN POTASSIUM

ST 25MG TABLET

02354829	ACT LOSARTAN	ACG
02441195	AG-LOSARTAN	ANG
02379058	APO-LOSARTAN	APX
02403323	AURO-LOSARTAN	AUR
02445964	BIO-LOSARTAN	BMI
02182815	COZAAR	FRS
02398834	JAMP-LOSARTAN	JMP
02388790	LOSARTAN	SIV
02388863	LOSARTAN	SAN
02394367	LOSARTAN	PDL
02405733	MINT-LOSARTAN	MIN
02309750	PMS-LOSARTAN	PMS
02313332	SANDOZ LOSARTAN	SDZ
02424967	SEPTA-LOSARTAN	SPT
02380838	TEVA-LOSARTAN	TEV
02426595	VAN-LOSARTAN	VAN

ST 50MG TABLET

02354837	ACT LOSARTAN	ACG
02441209	AG-LOSARTAN	ANG
02353504	APO-LOSARTAN	APX
02403331	AURO-LOSARTAN	AUR
02445972	BIO-LOSARTAN	BMI
02182874	COZAAR	FRS
02398842	JAMP-LOSARTAN	JMP
02388804	LOSARTAN	SIV
02388871	LOSARTAN	SAN
02394375	LOSARTAN	PDL

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

LOSARTAN POTASSIUM

ST 50MG TABLET

02405741	MINT-LOSARTAN	MIN
02309769	PMS-LOSARTAN	PMS
02404478	RAN-LOSARTAN	RBY
02313340	SANDOZ LOSARTAN	SDZ
02424975	SEPTA-LOSARTAN	SPT
02357968	TEVA-LOSARTAN	TEV
02426609	VAN-LOSARTAN	VAN

ST 100MG TABLET

02354845	ACT LOSARTAN	ACG
02441217	AG-LOSARTAN	ANG
02353512	APO-LOSARTAN	APX
02403358	AURO-LOSARTAN	AUR
02445980	BIO-LOSARTAN	BMI
02182882	COZAAR	FRS
02398850	JAMP-LOSARTAN	JMP
02388812	LOSARTAN	SIV
02388898	LOSARTAN	SAN
02394383	LOSARTAN	PDL
02405768	MINT-LOSARTAN	MIN
02309777	PMS-LOSARTAN	PMS
02404486	RAN-LOSARTAN	RBY
02313359	SANDOZ LOSARTAN	SDZ
02424983	SEPTA-LOSARTAN	SPT
02357976	TEVA-LOSARTAN	TEV
02426617	VAN-LOSARTAN	VAN

LOSARTAN POTASSIUM, HYDROCHLOROTHIAZIDE

ST 50MG & 12.5MG TABLET

02371235	APO-LOSARTAN/HCTZ	APX
02423642	AURO-LOSARTAN HCT	AUR
02230047	HYZAAR	FRS
02408244	JAMP-LOSARTAN HCTZ	JMP
02388960	LOSARTAN HCT	SIV
02427648	LOSARTAN/HCTZ	SAN
02394391	LOSARTAN-HCTZ	PDL
02389657	MINT-LOSARTAN/HCTZ	MIN
02392224	PMS-LOSARTAN-HCTZ	PMS
02313375	SANDOZ LOSARTAN HCT	SDZ
02428539	SEPTA-LOSARTAN HCTZ	SPT
02358263	TEVA-LOSARTAN/HCTZ	TEV

ST 100MG & 12.5MG TABLET

02371243	APO-LOSARTAN/HCTZ	APX
02423650	AURO-LOSARTAN HCT	AUR
02297841	HYZAAR	FRS
02388979	LOSARTAN HCT	SIV
02427656	LOSARTAN/HCTZ	SAN
02394405	LOSARTAN-HCTZ	PDL
02389665	MINT-LOSARTAN/HCTZ	MIN
02392232	PMS-LOSARTAN-HCTZ	PMS
02362449	SANDOZ LOSARTAN HCT	SDZ
02377144	TEVA-LOSARTAN/HCTZ	TEV

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

LOSARTAN POTASSIUM, HYDROCHLOROTHIAZIDE

ST **100MG & 25MG TABLET**

02371251	APO-LOSARTAN/HCTZ	APX
02423669	AURO-LOSARTAN HCT	AUR
02241007	HYZAAR DS	FRS
02408252	JAMP-LOSARTAN HCTZ	JMP
02388987	LOSARTAN HCT	SIV
02427664	LOSARTAN/HCTZ	SAN
02394413	LOSARTAN-HCTZ	PDL
02389673	MINT-LOSARTAN/HCTZ	MIN
02392240	PMS-LOSARTAN-HCTZ	PMS
02313383	SANDOZ LOSARTAN HCT	SDZ
02428547	SEPTA-LOSARTAN HCTZ	SPT
02377152	TEVA-LOSARTAN/HCTZ	TEV

OLMESARTAN MEDOXOMIL

ST **20MG TABLET**

02442191	ACT OLMESARTAN	TEV
02453452	APO-OLMESARTAN	APX
02443864	AURO-OLMESARTAN	AUR
02461641	JAMP-OLMESARTAN	JMP
02318660	OLMETEC	FRS
02461307	PMS-OLMESARTAN	PMS
02471353	RIVA-OLMESARTAN	RIV
02443414	SANDOZ OLMESARTAN	SDZ

ST **40MG TABLET**

02442205	ACT OLMESARTAN	TEV
02453460	APO-OLMESARTAN	APX
02443872	AURO-OLMESARTAN	AUR
02461668	JAMP-OLMESARTAN	JMP
02318679	OLMETEC	FRS
02461315	PMS-OLMESARTAN	PMS
02471361	RIVA-OLMESARTAN	RIV
02443422	SANDOZ OLMESARTAN	SDZ

OLMESARTAN MEDOXOMIL, HYDROCHLOROTHIAZIDE

ST **20MG & 12.5MG TABLET**

02443112	ACT OLMESARTAN HCT	TEV
02453606	APO-OLMESARTAN/HCTZ	APX

ST **20MG/12.5MG TABLET**

02319616	OLMETEC PLUS	FRS
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ST **40MG & 12.5MG TABLET**

02443120	ACT OLMESARTAN HCT	TEV
02453614	APO-OLMESARTAN/HCTZ	APX

ST **40MG & 25MG TABLET**

02443139	ACT OLMESARTAN HCT	TEV
02453622	APO-OLMESARTAN/HCTZ	APX

ST **40MG/12.5MG TABLET**

02319624	OLMETEC PLUS	FRS
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ST **40MG/25MG TABLET**

02319632	OLMETEC PLUS	FRS
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24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

TELMISARTAN

ST **40MG TABLET**

02420082	APO-TELMISARTAN	APX
02453568	AURO-TELMISARTAN	AUR
02240769	MICARDIS	BOE
02391236	PMS-TELMISARTAN	PMS
02375958	SANDOZ TELMISARTAN	SDZ
02388944	TELMISARTAN	SAN
02390345	TELMISARTAN	SIV
02395223	TELMISARTAN	PDL
02407485	TELMISARTAN	ACC
02432897	TELMISARTAN	PMS
02320177	TEVA-TELMISARTAN	TEV

ST **80MG TABLET**

02420090	APO-TELMISARTAN	APX
02453576	AURO-TELMISARTAN	AUR
02240770	MICARDIS	BOE
02391244	PMS-TELMISARTAN	PMS
02375966	SANDOZ TELMISARTAN	SDZ
02388952	TELMISARTAN	SAN
02390353	TELMISARTAN	SIV
02395231	TELMISARTAN	PDL
02407493	TELMISARTAN	ACC
02432900	TELMISARTAN	PMS
02320185	TEVA-TELMISARTAN	TEV

TELMISARTAN, HYDROCHLOROTHIAZIDE

ST **80MG & 12.5MG TABLET**

02419114	ACH-TELMISARTAN HCTZ	ACC
02420023	APO-TELMISARTAN/HCTZ	APX
02456389	AURO-TELMISARTAN HCTZ	AUR
02244344	MICARDIS PLUS	BOE
02401665	PMS-TELMISARTAN-HCTZ	PMS
02393557	SANDOZ TELMISARTAN HCT	SDZ
02390302	TELMISARTAN HCTZ	SIV
02395355	TELMISARTAN/HCTZ	SAN
02395525	TELMISARTAN-HCTZ	PDL
02433214	TELMISARTAN-HCTZ	PMS
02330288	TEVA-TELMISARTAN HCTZ	TEV

ST **80MG & 25MG TABLET**

02419122	ACH-TELMISARTAN HCTZ	ACC
02420031	APO-TELMISARTAN/HCTZ	APX
02456397	AURO-TELMISARTAN HCTZ	AUR
02318709	MICARDIS PLUS	BOE
02393565	SANDOZ TELMISARTAN HCT	SDZ
02390310	TELMISARTAN HCTZ	SIV
02395363	TELMISARTAN/HCTZ	SAN
02395533	TELMISARTAN-HCTZ	PDL
02433222	TELMISARTAN-HCTZ	PMS
02379252	TEVA-TELMISARTAN HCTZ	TEV

VALSARTAN

ST **40MG TABLET**

02337487	ACT VALSARTAN	ACG
02371510	APO-VALSARTAN	APX
02414201	AURO-VALSARTAN	AUR

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

VALSARTAN

ST **40MG TABLET**

02270528	DIOVAN	NVR
02363062	RAN-VALSARTAN	RBY
02356740	SANDOZ VALSARTAN	SDZ
02356643	TEVA-VALSARTAN	TEV
02366940	VALSARTAN	SAN
02367726	VALSARTAN	PDL
02384523	VALSARTAN	SIV

ST **80MG TABLET**

02337495	ACT VALSARTAN	ACG
02371529	APO-VALSARTAN	APX
02414228	AURO-VALSARTAN	AUR
02244781	DIOVAN	NVR
02363100	RAN-VALSARTAN	RBY
02356759	SANDOZ VALSARTAN	SDZ
02356651	TEVA-VALSARTAN	TEV
02366959	VALSARTAN	SAN
02367734	VALSARTAN	PDL
02384531	VALSARTAN	SIV

ST **160MG TABLET**

02337509	ACT VALSARTAN	ACG
02371537	APO-VALSARTAN	APX
02414236	AURO-VALSARTAN	AUR
02244782	DIOVAN	NVR
02363119	RAN-VALSARTAN	RBY
02356767	SANDOZ VALSARTAN	SDZ
02356678	TEVA-VALSARTAN	TEV
02366967	VALSARTAN	SAN
02367742	VALSARTAN	PDL
02384558	VALSARTAN	SIV

ST **320MG TABLET**

02337517	ACT VALSARTAN	ACG
02371545	APO-VALSARTAN	APX
02414244	AURO-VALSARTAN	AUR
02289504	DIOVAN	NVR
02356775	SANDOZ VALSARTAN	SDZ
02356686	TEVA-VALSARTAN	TEV
02366975	VALSARTAN	SAN
02367750	VALSARTAN	PDL
02384566	VALSARTAN	SIV

VALSARTAN, HYDROCHLOROTHIAZIDE

ST **80MG & 12.5MG TABLET**

02382547	APO-VALSARTAN/HCTZ	APX
02408112	AURO-VALSARTAN HCT	AUR
02241900	DIOVAN-HCT	NVR
02356694	SANDOZ VALSARTAN HCT	SDZ
02356996	TEVA-VALSARTAN/HCTZ	TEV
02367009	VALSARTAN HCT	SAN
02384736	VALSARTAN HCT	SIV
02367769	VALSARTAN-HCTZ	PDL

ST **160MG & 12.5MG TABLET**

02382555	APO-VALSARTAN/HCTZ	APX
02408120	AURO-VALSARTAN HCT	AUR

24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS

VALSARTAN, HYDROCHLOROTHIAZIDE

ST **160MG & 12.5MG TABLET**

02241901	DIOVAN-HCT	NVR
02356708	SANDOZ VALSARTAN HCT	SDZ
02357003	TEVA-VALSARTAN/HCTZ	TEV
02367017	VALSARTAN HCT	SAN
02384744	VALSARTAN HCT	SIV
02367777	VALSARTAN-HCTZ	PDL

ST **160MG & 25MG TABLET**

02382563	APO-VALSARTAN/HCTZ	APX
02408139	AURO-VALSARTAN HCT	AUR
02246955	DIOVAN-HCT	NVR
02356716	SANDOZ VALSARTAN HCT	SDZ
02357011	TEVA-VALSARTAN/HCTZ	TEV
02367025	VALSARTAN HCT	SAN
02384752	VALSARTAN HCT	SIV
02367785	VALSARTAN-HCTZ	PDL

ST **320MG & 12.5MG TABLET**

02382571	APO-VALSARTAN/HCTZ	APX
02408147	AURO-VALSARTAN HCT	AUR
02308908	DIOVAN-HCT	NVR
02356724	SANDOZ VALSARTAN HCT	SDZ
02357038	TEVA-VALSARTAN/HCTZ	TEV
02367033	VALSARTAN HCT	SAN
02384760	VALSARTAN HCT	SIV

ST **320MG & 25MG TABLET**

02382598	APO-VALSARTAN/HCTZ	APX
02408155	AURO-VALSARTAN HCT	AUR
02308916	DIOVAN-HCT	NVR
02356732	SANDOZ VALSARTAN HCT	SDZ
02357046	TEVA-VALSARTAN/HCTZ	TEV
02367041	VALSARTAN HCT	SAN
02384779	VALSARTAN HCT	SIV

24:32.20 MINERALOCORTICOIDE (ALDOSTERONE) RECEPTOR ANTAGONISTS

ENALAPRIL MALEATE

ST **2.5MG TABLET**

02474786	JAMP ENALAPRIL	JMP
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ST **5MG TABLET**

02474794	JAMP ENALAPRIL	JMP
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ST **10MG TABLET**

02474808	JAMP ENALAPRIL	JMP
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ST **20MG TABLET**

02474816	JAMP ENALAPRIL	JMP
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**24:32.20 MINERALOCORTICOIDE
(ALDOSTERONE) RECEPTOR
ANTAGONISTS**

EPLERENONE

Limited use benefit (prior approval required).

For the treatment of patients with New York Heart Association (NYHA) class II chronic heart failure with left ventricular systolic dysfunction (with ejection fraction \leq 35%), as an adjunct to standard therapy.

Note: Patients must be on optimal therapy with an angiotensin-converting-enzyme (ACE) inhibitor or an angiotensin-receptor blocker (ARB), and a beta-blocker (unless contraindicated) at the recommended dose or

25MG TABLET

02323052	INSPRA	PFI
02471442	MINT-EPLERENONE	MIN

50MG TABLET

02323060	INSPRA	PFI
02471450	MINT-EPLERENONE	MIN

HYDROCHLOROTHIAZIDE, SPIRONOLACTONE

ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503009	ALDACTAZIDE ORAL LIQUID	UNK
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SPIRONOLACTONE

ST **25MG TABLET**

00028606	ALDACTONE	PFI
00613215	TEVA-SPIRONOLACTONE	TEV

ST **100MG TABLET**

00285455	ALDACTONE	PFI
00613223	TEVA-SPIRONOLACTONE	TEV

ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503001	SPIRONOLACTONE ORAL LIQUID	UNK
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24:32.92

VALSARTAN, SACUBITRIL

Limited use benefit (prior approval required).

For the treatment of New York Heart Association (NYHA) class II or III heart failure if the following criteria are met:

- Must be initiated by a physician experienced in the treatment of heart failure; AND

- Left ventricular ejection fraction < 40%; AND
- NYHA class II to III symptoms despite at least four weeks of treatment with an angiotensin-converting enzyme inhibitor (ACEI) or an angiotensin II receptor antagonist (ARB); OR If your patient has a contraindication or intolerance to ACEI or ARBs;

AND

- Must be used in combination with a beta blocker and an aldosterone antagonist (if tolerated); OR If your patient has a contraindication or intolerance to beta blockers or aldosterone antagonists.

26MG & 24MG TABLET

02446928	ENTRESTO	NVR
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51MG & 49MG TABLET

02446936	ENTRESTO	NVR
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103MG & 97MG TABLET

02446944	ENTRESTO	NVR
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28:00 CENTRAL NERVOUS SYSTEM AGENTS

28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

ACETYLSALICYLIC ACID

Limited use benefit (prior approval is not required).

ASA 80 mg tablets are a benefit to clients age 21 years and under to allow access for use in pediatric conditions (e.g. Kawasaki Syndrome).

150MG SUPPOSITORY

00785547 ASA PMS

650MG SUPPOSITORY

00582867 ASA PMS

ST 80MG TABLET

02269139 ACETYLSALICYLIC ACID JMP

02295563 LOWPRIN EUR

02202360 RIVASA RIV

ST 325MG TABLET

00472468 APO ASA APX

00530336 ASA VTH

02150328 ASPIRIN BAY

ST 80MG TABLET (CHEWABLE)

02009013 ASAPHEN PMS

02280167 ASATAB ODN

02250675 EURO-ASA EUR

02296004 LOWPRIN SDZ

02429950 M-ASA MAN

02311518 PRO-AAS PDL

02202352 RIVASA RIV

ST 81MG TABLET (CHEWABLE)

02394790 ASA DAILY LOW DOSE PMS

02243974 ENTROPHEN PED

ST 80MG TABLET (DELAYED RELEASE)

02427176 ASA EC SAN

02238545 ASAPHEN PMS

02283905 JAMP-ASA JMP

02311496 PRO-AAS PDL

ST 81MG TABLET (DELAYED RELEASE)

02461471 APO-ASA LD APX

02244993 ASA PMS

02372177 ASA VTH

02433044 ASA PMS

02449277 ASA TLI

02243101 ASA DAILY LOW DOSE PMS

02377683 ASA DAILY LOW DOSE APX

02426811 ASA EC SAN

02242281 ENTROPHEN PED

02283700 PRAXIS ASA DAILY LOW DOSE PMS

02420279 RIVASA EC RIV

ST 162MG TABLET (DELAYED RELEASE)

02247550 ASAPHEN EC PMS

ST 325MG TABLET (DELAYED RELEASE)

02010526 ASA VTH

02352427 ASATAB EC ODN

02150417 ASPIRIN BAY

00010332 ENTROPHEN PED

28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

ACETYLSALICYLIC ACID

Limited use benefit (prior approval is not required).

ASA 80 mg tablets are a benefit to clients age 21 years and under to allow access for use in pediatric conditions (e.g. Kawasaki Syndrome).

ST 325MG TABLET (DELAYED RELEASE)

02050161 ENTROPHEN PED

00216666 NOVASEN TEV

ST 650MG TABLET (DELAYED RELEASE)

00794244 ASA VTH

02352435 ASATAB EC ODN

00229296 NOVASEN TEV

02284537 PMS-ASA EC PMS

ST 81MG TABLET (ENTERIC COATED)

02243896 ASA DAILY LOW DOSE PMS

02237726 ASPIRIN BAY

02243801 EQUATE DAILY LOW-DOSE PMS

02427206 JAMP-ASA EC VTH

ST 325MG TABLET (ENTERIC COATED)

00510696 ASA APX

02285371 PMS-ASA EC PMS

ST 650MG TABLET (ENTERIC COATED)

00472476 ASA APX

00010340 ENTROPHEN PED

01905392 ENTROPHEN PED

CELECOXIB

ST 100MG CAPSULE

02420155 ACT CELECOXIB ACG

02437570 AG-CELECOXIB ANG

02418932 APO-CELECOXIB APX

02445670 AURO-CELECOXIB AUR

02426382 BIO-CELECOXIB BMI

02239941 CELEBEX PFI

02424371 CELECOXIB PDL

02429675 CELECOXIB SIV

02436299 CELECOXIB SAN

02291975 GD-CELECOXIB PFI

02424533 JAMP-CELECOXIB JMP

02420058 MAR-CELECOXIB MAR

02412497 MINT-CELECOXIB MIN

02355442 PMS-CELECOXIB PMS

02426366 PRIVA-CELECOXIB PHA

02412373 RAN-CELECOXIB RBY

02425386 RIVA-CELECOX RIV

02321246 SANDOZ CELECOXIB SDZ

02442639 SDZ CELECOXIB SDZ

ST 200MG CAPSULE

02420163 ACT CELECOXIB ACG

02437589 AG-CELECOXIB ANG

02418940 APO-CELECOXIB APX

02445689 AURO-CELECOXIB AUR

02426390 BIO-CELECOXIB BMI

02239942 CELEBEX PFI

02424398 CELECOXIB PDL

28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

CELECOXIB

ST 200MG CAPSULE

02429683	CELECOXIB	SIV
02436302	CELECOXIB	SAN
02291983	GD-CELECOXIB	PFI
02424541	JAMP-CELECOXIB	JMP
02420066	MAR-CELECOXIB	MAR
02412500	MINT-CELECOXIB	MIN
02355450	PMS-CELECOXIB	PMS
02426374	PRIVA-CELECOXIB	PHA
02412381	RAN-CELECOXIB	RBY
02425394	RIVA-CELECOX	RIV
02321254	SANDOZ CELECOXIB	SDZ
02442647	SDZ CELECOXIB	SDZ

DICLOFENAC SODIUM

50MG SUPPOSITORY

02231506	PMS-DICLOFENAC	PMS
02261928	SANDOZ-DICLOFENAC	SDZ
00632724	VOLTAREN	NVR

100MG SUPPOSITORY

02231508	PMS-DICLOFENAC	PMS
02261936	SANDOZ-DICLOFENAC	SDZ
00632732	VOLTAREN	NVR

ST 25MG TABLET (DELAYED RELEASE)

02231662	DOM-DICLOFENAC	DPC
02302616	PMS-DICLOFENAC	PMS

ST 50MG TABLET (DELAYED RELEASE)

02231663	DOM-DICLOFENAC	DPC
02302624	PMS-DICLOFENAC	PMS
02261960	SANDOZ-DICLOFENAC	SDZ
00514012	VOLTAREN	NVR

ST 25MG TABLET (ENTERIC COATED)

00839175	APO-DICLO	APX
00808539	TEVA-DICLOFENAC	TEV

ST 50MG TABLET (ENTERIC COATED)

00839183	APO-DICLO	APX
00870978	DICLOFENAC	PDL
02352397	DICLOFENAC EC	SAN
02231503	PMS-DICLOFENAC	PMS
00808547	TEVA-DICLOFENAC	TEV

ST 75MG TABLET (EXTENDED RELEASE)

02162814	APO-DICLO SR	APX
02224119	DICLOFENAC-SR	PDL
02231664	DOM-DICLOFENAC SR	DPC
02231504	PMS-DICLOFENAC	PMS
02261901	SANDOZ-DICLOFENAC SR	SDZ
02158582	TEVA-DICLOFENAC SR	TEV
00782459	VOLTAREN	NVR

ST 100MG TABLET (EXTENDED RELEASE)

02091194	APO-DICLO SR	APX
02224127	DICLOFENAC-SR	PDL
02231505	PMS-DICLOFENAC	PMS
02261944	SANDOZ-DICLOFENAC SR	SDZ
02048698	TEVA-DICLOFENAC SR	TEV

28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

DICLOFENAC SODIUM

ST 100MG TABLET (EXTENDED RELEASE)

00590827	VOLTAREN SR	NVR
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DICLOFENAC SODIUM (TOPICAL)

Limited use benefit (prior approval required).

For the treatment of osteoarthritis when:

- pain is inadequately controlled with acetaminophen AND a non-steroidal anti-inflammatory (NSAID); OR
- there is contraindication to acetaminophen and NSAID; OR
- there is intolerance to acetaminophen and NSAID.

ST 1.5% SOLUTION

02354403	APO-DICLOFENAC	APX
02476134	DICLOFENAC SODIUM	TEL
02434571	DICLOFENAC TOPICAL	RAX
02472309	JAMP DICLOFENAC TOPICAL	JMP
02356783	PMS-DICLOFENAC	PMS
02420988	TARO-DICLOFENAC	TAR

DIFLUNISAL

ST 250MG TABLET

02039486	DIFLUNISAL	AAP
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ST 500MG TABLET

02039494	DIFLUNISAL	AAP
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FLURBIPROFEN

ST 50MG TABLET

01912046	APO-FLURBIPROFEN	AAP
02100509	TEVA-FLURBIPROFEN	TEV

ST 100MG TABLET

01912038	APO-FLURBIPROFEN	AAP
02100517	TEVA-FLURBIPROFEN	TEV

IBUPROFEN

ST 40MG/ML DROP

02242522	ADVIL PEDIATRIC DROPS	PFI
02238626	CHILDREN'S MOTRIN	MCL

ST 20MG/ML SUSPENSION

02232297	CHILDREN'S ADVIL	PFI
02354799	CHILDREN'S EUROPROFEN	PED
02242365	CHILDREN'S MOTRIN	MCL

ST 100MG TABLET

02246403	ADVIL	PFI
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ST 200MG TABLET

01933558	ADVIL	PFI
00441643	APO-IBUPROFEN	APX
02257912	IBUPROFEN	PMT
02314754	IBUPROFEN	PMS
02314762	IBUPROFEN	PMS
02368072	IBUPROFEN	VTH
02186934	MOTRIN	MCL
00629324	NOVO-PROFEN	TEV

ST 300MG TABLET

00441651	APO IBUPROFEN	APX
00629332	NOVO-PROFEN	TEV

ST 400MG TABLET

00506052	APO IBUPROFEN	APX
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28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

IBUPROFEN

ST 400MG TABLET

00636533	IBUPROFEN	PDL
02314770	IBUPROFEN	PMS
02317338	IBUPROFEN	PMT
02401290	JAMP-IBUPROFEN	JMP
00629340	NOVO-PROFEN	TEV
00836133	PMS-IBUPROFEN	PMS

ST 600MG TABLET

00585114	APO IBUPROFEN	APX
00629359	TEVA-PROFEN	TEV

INDOMETHACIN

ST 25MG CAPSULE

00611158	APO INDOMETHACIN	APX
02461811	MINT-INDOMETHACIN	MIN
00337420	TEVA-INDOMETHACIN	TEV

ST 50MG CAPSULE

00611166	APO INDOMETHACIN	APX
02461536	MINT-INDOMETHACIN	MIN
00337439	TEVA-INDOMETHACIN	TEV

50MG SUPPOSITORY

02231799	SANDOZ INDOMETHACIN	SDZ
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100MG SUPPOSITORY

02231800	SANDOZ INDOMETHACIN	SDZ
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KETOPROFEN

ST 50MG CAPSULE

00790427	KETOPROFEN	AAP
02150808	PMS-KETOPROFEN	PMS

100MG SUPPOSITORY

02015951	PMS-KETOPROFEN	PMS
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ST 50MG TABLET (ENTERIC COATED)

00790435	KETOPROFEN-E	AAP
02150816	PMS-KETOPROFEN	PMS

ST 100MG TABLET (ENTERIC COATED)

00842664	KETOPROFEN-E	AAP
02150824	PMS-KETOPROFEN	PMS

ST 200MG TABLET (EXTENDED RELEASE)

02172577	KETOPROFEN SR	AAP
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MEFENAMIC ACID

ST 250MG CAPSULE

02237826	DOM-MEFENAMIC ACID	DPC
02229452	MEFENAMIC	AAP
00155225	PONSTAN	AAP

MELOXICAM

ST 7.5MG TABLET

02250012	ACT MELOXICAM	TEV
02248973	APO-MELOXICAM	APX
02390884	AURO-MELOXICAM	AUR
02248605	DOM-MELOXICAM	DPC
02353148	MELOXICAM	SAN
02248267	PMS-MELOXICAM	PMS
02258315	TEVA-MELOXICAM	TEV

28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

MELOXICAM

ST 15MG TABLET

02250020	ACT MELOXICAM	TEV
02248974	APO-MELOXICAM	APX
02390892	AURO-MELOXICAM	AUR
02248606	DOM-MELOXICAM	DPC
02324334	MELOXICAM	PDL
02353156	MELOXICAM	SAN
02248268	PMS-MELOXICAM	PMS
02258323	TEVA-MELOXICAM	TEV

MISOPROSTOL, DICLOFENAC SODIUM

ST 200MCG & 50MG TABLET

02400596	SANDOZ DICLOFENAC MISOPROSTOL	SDZ
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ST 200MCG & 75MG TABLET

02400618	SANDOZ DICLOFENAC MISOPROSTOL	SDZ
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ST 200MCG & 50MG TABLET (DELAYED RELEASE)

01917056	ARTHROTEC	PFI
02341689	GD-DICLOFENAC/MISOPROSTOL	PFI
02413469	PMS-DICLOFENAC-MISOPROSTOL	PMS

ST 200MCG & 75MG TABLET (DELAYED RELEASE)

02229837	ARTHROTEC	PFI
02341697	GD-DICLOFENAC/MISOPROSTOL	PFI
02413477	PMS-DICLOFENAC-MISOPROSTOL	PMS

ST 200MCG & 50MG TABLET (ENTERIC COATED)

02397145	ACT DICLO-MISO	ACG
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ST 200MCG & 75MG TABLET (ENTERIC COATED)

02397153	ACT DICLO-MISO	ACG
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NAPROXEN

500MG SUPPOSITORY

02017237	PMS-NAPROXEN	PMS
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ST 25MG/ML SUSPENSION

02162431	NAPROXEN	PEI
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ST 125MG TABLET

00522678	APO NAPROXEN	APX
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ST 220MG TABLET

02362430	NAPROXEN	PMS
02385007	NAPROXEN SODIUM	APX

ST 250MG TABLET

00522651	APO-NAPROXEN	APX
00590762	NAPROXEN	PDL
02350750	NAPROXEN	SAN
00565350	TEVA-NAPROXEN	TEV

ST 275MG TABLET

02162725	ANAPROX	APU
00784354	APO-NAPRO-NA	APX
02351013	NAPROXEN SODIUM	SAN
00887056	NAPROXEN-NA	PDL
00778389	TEVA-NAPROXEN	TEV

ST 375MG TABLET

00600806	APO-NAPROXEN	APX
00655686	NAPROXEN	PDL
02350769	NAPROXEN	SAN

28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

NAPROXEN

ST 375MG TABLET		
00627097	TEVA-NAPROXEN	TEV
ST 500MG TABLET		
00592277	APO-NAPROXEN	APX
00618721	NAPROXEN	PDL
02350777	NAPROXEN	SAN
00589861	TEVA-NAPROXEN	TEV
ST 500MG TABLET		
02162717	ANAPROX DS	APU
01940309	APO-NAPRO-NA DS	APX
02351021	NAPROXEN SODIUM DS	SAN
02153386	NAPROXEN-NA DF	PDL
02026600	TEVA-NAPROXEN DS	TEV
ST 250MG TABLET (ENTERIC COATED)		
02246699	APO-NAPROXEN EC	APX
02350785	NAPROXEN EC	SAN
02243312	TEVA-NAPROXEN	TEV
ST 375MG TABLET (ENTERIC COATED)		
02246700	APO-NAPROXEN EC	APX
02162415	NAPROSYN	APU
02350793	NAPROXEN EC	SAN
02294702	PMS-NAPROXEN EC	PMS
02310945	PRO-NAPROXEN	PDL
02243313	TEVA-NAPROXEN	TEV
ST 500MG TABLET (ENTERIC COATED)		
02246701	APO-NAPROXEN EC	APX
02162423	NAPROSYN	APU
02350807	NAPROXEN EC	SAN
02294710	PMS-NAPROXEN EC	PMS
02310953	PRO-NAPROXEN	PDL
02243314	TEVA-NAPROXEN	TEV
ST 750MG TABLET (EXTENDED RELEASE)		
02162466	NAPROSYN	APU

PIROXICAM

ST 10MG CAPSULE		
00642886	APO PIROXICAM	APX
00695718	TEVA-PIROXICAM	TEV
ST 20MG CAPSULE		
00642894	APO PIROXICAM	APX
00695696	TEVA-PIROXICAM	TEV

SULINDAC

ST 150MG TABLET		
00745588	TEVA-SULINDAC	TEV
ST 200MG TABLET		
00745596	TEVA-SULINDAC	TEV

TIAPROFENIC ACID

ST 200MG TABLET		
02230827	PMS-TIAPROFENIC	PMS
02179679	TEVA-TIAPROFENIC	TEV
ST 300MG TABLET		
02231060	DOM-TIAPROFENIC	DPC
02179687	TEVA-TIAPROFENIC	TEV

28:08.08 OPIATE AGONISTS

ACETAMINOPHEN, CAFFEINE CITRATE, CODEINE PHOSPHATE

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

300MG & 15MG & 15MG TABLET		
00653241	RATIO-LENOLTEC NO 2	TEV
02163934	TYLENOL WITH CODEINE NO.2	JSO
300MG & 15MG & 30MG TABLET		
00653276	RATIO-LENOLTEC NO 3	TEV
02163926	TYLENOL WITH CODEINE NO.3	JSO
325MG & 30MG & 15MG TABLET		
00293504	ATASOL 15	CHU
325MG & 30MG & 30MG TABLET		
00293512	ATASOL 30	CHU

ACETAMINOPHEN, CODEINE PHOSPHATE

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

32MG & 1.6MG/ML ELIXIR		
00816027	PMS-ACETAMINOPHEN	PMS
300MG & 30MG TABLET		
02232658	PROCET-30	PDL
00608882	TEVA-EMTEC-30	TEV
00789828	TRIA TEC-30	RIV

ACETAMINOPHEN, OXYCODONE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

325MG & 5MG TABLET		
02324628	APO-OXYCODONE/ACET	APX
02361361	OXYCODONE/ACET	SAN
02327171	OXYCODONE-ACET	PDL
02242468	RIVACOCET	RIV
02307898	SANDOZ OXYCODONE/ACETAMINOPHEN	SDZ
00608165	TEVA-OXYCOCET	TEV

28:08.08 OPIATE AGONISTS

ACETYSALICYLIC ACID, OXYCODONE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

325MG & 5MG TABLET

00608157 TEVA-OXYCODAN TEV

CODEINE MONOHYDRATE, CODEINE SULFATE TRIHYDRATE

Limited use benefit (prior approval required).

For treatment of:

- chronic pain and palliative care patients as an alternative to products containing codeine in combination with acetaminophen or ASA with or without caffeine; OR
- chronic pain and palliative care patients as an alternative to regular release codeine tablets when large doses are required.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

50MG TABLET (EXTENDED RELEASE)

02230302 CODEINE CONTIN CR PFR

100MG TABLET (EXTENDED RELEASE)

02163748 CODEINE CONTIN CR PFR

150MG TABLET (EXTENDED RELEASE)

02163780 CODEINE CONTIN CR PFR

200MG TABLET (EXTENDED RELEASE)

02163799 CODEINE CONTIN CR PFR

CODEINE PHOSPHATE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

5MG/ML LIQUID

00050024 CODEINE PHOSPHATE ATL

2MG/ML SOLUTION

00380571 LINCTUS CODEINE ATL

15MG TABLET

02009889 CODEINE RIV

00593435 TEVA-CODEINE TEV

30MG TABLET

02009757 CODEINE RIV

00593451 TEVA-CODEINE TEV

28:08.08 OPIATE AGONISTS

FENTANYL

Limited use benefit (prior approval required).

For the management of chronic pain in patients who are unresponsive or intolerant to at least one long-acting oral sustained released product, such as morphine, hydromorphone and oxycodone, despite appropriate dose titration and adjunctive therapy including laxatives and antiemetics.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

12MCG/HR PATCH

02395657 FENTANYL PDL

02341379 PMS-FENTANYL MTX PMS

02330105 RAN-FENTANYL MATRIX RBY

02327112 SANDOZ FENTANYL SDZ

02311925 TEVA-FENTANYL TEV

25MCG/HR PATCH

02395665 FENTANYL PDL

02341387 PMS-FENTANYL MTX PMS

02330113 RAN-FENTANYL MATRIX RBY

02327120 SANDOZ FENTANYL SDZ

02282941 TEVA-FENTANYL TEV

50MCG/HR PATCH

02395673 FENTANYL PDL

02341395 PMS-FENTANYL MTX PMS

02330121 RAN-FENTANYL MATRIX RBY

02327147 SANDOZ FENTANYL SDZ

02282968 TEVA-FENTANYL TEV

75MCG/HR PATCH

02395681 FENTANYL PDL

02341409 PMS-FENTANYL MTX PMS

02330148 RAN-FENTANYL MATRIX RBY

02327155 SANDOZ FENTANYL SDZ

02282976 TEVA-FENTANYL TEV

100MCG/HR PATCH

02395703 FENTANYL PDL

02341417 PMS-FENTANYL MTX PMS

02330156 RAN-FENTANYL MATRIX RBY

02327163 SANDOZ FENTANYL SDZ

02282984 TEVA-FENTANYL TEV

28:08.08 OPIATE AGONISTS

HYDROMORPHONE HYDROCHLORIDE

Limited use benefit.
Prior approval required for controlled release capsules only.
Regular release dosage forms are full benefits and do not require prior approval.

For treatment of moderate to severe chronic pain when other opioids such as morphine have been ineffective in controlling pain or in patients experiencing intolerable side effects.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

3MG CAPSULE (EXTENDED RELEASE)		
02476614	APO-HYDROMORPHONE	APX
4.5MG CAPSULE (EXTENDED RELEASE)		
02476622	APO-HYDROMORPHONE	APX
6MG CAPSULE (EXTENDED RELEASE)		
02476630	APO-HYDROMORPHONE	APX
9MG CAPSULE (EXTENDED RELEASE)		
02476649	APO-HYDROMORPHONE	APX
12MG CAPSULE (EXTENDED RELEASE)		
02476657	APO-HYDROMORPHONE	APX
18MG CAPSULE (EXTENDED RELEASE)		
02476665	APO-HYDROMORPHONE	APX
24MG CAPSULE (EXTENDED RELEASE)		
02476673	APO-HYDROMORPHONE	APX
30MG CAPSULE (EXTENDED RELEASE)		
02476681	APO-HYDROMORPHONE	APX
3MG CAPSULE (SUSTAINED RELEASE)		
02125323	HYDROMORPH CONTIN	PFR
4.5MG CAPSULE (SUSTAINED RELEASE)		
02359502	HYDROMORPH CONTIN	PFR
6MG CAPSULE (SUSTAINED RELEASE)		
02125331	HYDROMORPH CONTIN	PFR
9MG CAPSULE (SUSTAINED RELEASE)		
02359510	HYDROMORPH CONTIN	PFR
12MG CAPSULE (SUSTAINED RELEASE)		
02125366	HYDROMORPH CONTIN	PFR
18MG CAPSULE (SUSTAINED RELEASE)		
02243562	HYDROMORPH CONTIN	PFR
24MG CAPSULE (SUSTAINED RELEASE)		
02125382	HYDROMORPH CONTIN	PFR
30MG CAPSULE (SUSTAINED RELEASE)		
02125390	HYDROMORPH CONTIN	PFR
1MG/ML LIQUID		
01916386	PMS HYDROMORPHONE	PMS
3MG SUPPOSITORY		
01916394	PMS HYDROMORPHONE	PMS
1MG TABLET		
02364115	APO-HYDROMORPHONE	APX
00705438	DILAUDID	PFR
00885444	PMS-HYDROMORPHONE	PMS
02319403	TEVA-HYDROMORPHONE	TEV

28:08.08 OPIATE AGONISTS

HYDROMORPHONE HYDROCHLORIDE

Limited use benefit.
Prior approval required for controlled release capsules only.
Regular release dosage forms are full benefits and do not require prior approval.

For treatment of moderate to severe chronic pain when other opioids such as morphine have been ineffective in controlling pain or in patients experiencing intolerable side effects.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

2MG TABLET		
02364123	APO-HYDROMORPHONE	APX
00125083	DILAUDID	PFR
00885436	PMS-HYDROMORPHONE	PMS
02319411	TEVA-HYDROMORPHONE	TEV
4MG TABLET		
02364131	APO-HYDROMORPHONE	APX
00125121	DILAUDID	PFR
00885401	PMS-HYDROMORPHONE	PMS
02319438	TEVA-HYDROMORPHONE	TEV
8MG TABLET		
02364158	APO-HYDROMORPHONE	APX
00786543	DILAUDID	PFR
00885428	PMS-HYDROMORPHONE	PMS
02319446	TEVA-HYDROMORPHONE	TEV
METHADONE HYDROCHLORIDE		
POWDER		
00908835	METHADONE POWDER (OAT)	MDS
10MG/ML SOLUTION		
02244290	METADOL-D	PAL
02394596	METHADOSE	MAT
02394618	METHADOSE	MAT
METHADONE HYDROCHLORIDE (BC ONLY)		
10MG/ML ORAL LIQUID		
66999999	METHADOSE DEL. W DIRECT INTER (OAT)	UNK
67000000	METHADOSE DEL. W/OUT DIR INTER (OAT)	UNK
66999997	METHADOSE W DIRECT INTERACTION (OAT)	UNK
66999998	METHADOSE W/OUT DIRECT INTER (OAT)	UNK

28:08.08 OPIATE AGONISTS

METHADONE HYDROCHLORIDE (METADOL)

Limited use benefit (prior approval required) with the following criteria:

Prescriber is registered with Health Canada and is eligible to prescribe methadone for the management of pain; AND For the management of moderate to severe cancer pain or chronic non-cancer pain, as an alternative to other opioids; OR For the management of pain for palliative care patients. Pharmacists may only dispense a maximum supply of 30 days at one time.

1MG/ML SOLUTION		
02247694 METADOL		PAL
10MG/ML SOLUTION		
02241377 METADOL		PAL
1MG TABLET		
02247698 METADOL		PAL
5MG TABLET		
02247699 METADOL		PAL
10MG TABLET		
02247700 METADOL		PAL
25MG TABLET		
02247701 METADOL		PAL

MORPHINE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

1MG/ML SYRUP		
00614491 DOLORAL 1		ATL
5MG/ML SYRUP		
00614505 DOLORAL 5		ATL

MORPHINE SULFATE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

10MG CAPSULE (EXTENDED RELEASE)		
02019930 M-ESLON		ETH
15MG CAPSULE (EXTENDED RELEASE)		
02177749 M-ESLON		ETH
30MG CAPSULE (EXTENDED RELEASE)		
02019949 M-ESLON		ETH
60MG CAPSULE (EXTENDED RELEASE)		
02019957 M-ESLON		ETH
100MG CAPSULE (EXTENDED RELEASE)		
02019965 M-ESLON		ETH
200MG CAPSULE (EXTENDED RELEASE)		
02177757 M-ESLON		ETH

28:08.08 OPIATE AGONISTS

MORPHINE SULFATE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

5MG SUPPOSITORY		
00632228 STATEX		PAL
10MG SUPPOSITORY		
00632201 STATEX		PAL
20MG SUPPOSITORY		
00596965 STATEX		PAL
1MG/ML SYRUP		
00591467 STATEX		PAL
5MG TABLET		
00594652 STATEX		PAL
10MG TABLET		
00594644 STATEX		PAL
25MG TABLET		
00594636 STATEX		PAL
50MG TABLET		
00675962 STATEX		PAL
15MG TABLET (EXTENDED RELEASE)		
02350815 MORPHINE SR		SAN
02015439 MS CONTIN SR		PFR
02244790 SANDOZ MORPHINE SR		SDZ
02302764 TEVA-MORPHINE SR		TEV
30MG TABLET (EXTENDED RELEASE)		
02350890 MORPHINE SR		SAN
02014297 MS CONTIN SR		PFR
02244791 SANDOZ MORPHINE SR		SDZ
02302772 TEVA-MORPHINE SR		TEV
60MG TABLET (EXTENDED RELEASE)		
02350912 MORPHINE SR		SAN
02014300 MS CONTIN SR		PFR
02244792 SANDOZ MORPHINE SR		SDZ
02302780 TEVA-MORPHINE SR		TEV
100MG TABLET (EXTENDED RELEASE)		
02014319 MS CONTIN SR		PFR
02302799 TEVA-MORPHINE SR		TEV
200MG TABLET (EXTENDED RELEASE)		
02014327 MS CONTIN SR		PFR
02302802 TEVA-MORPHINE SR		TEV
5MG TABLET (IMMEDIATE RELEASE)		
02014203 MS IR		PFR
10MG TABLET (IMMEDIATE RELEASE)		
02014211 MS IR		PFR
20MG TABLET (IMMEDIATE RELEASE)		
02014238 MS IR		PFR
30MG TABLET (IMMEDIATE RELEASE)		
02014254 MS IR		PFR

28:08.08 OPIATE AGONISTS

MORPHINE SULFATE (KADIAN)

Limited use benefit (prior approval required).

- For the treatment of opioid dependence where methadone and Suboxone are not available or not appropriate; OR
- For the treatment of chronic pain.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

10MG CAPSULE (SUSTAINED RELEASE)

02242163	KADIAN	BGP
09991310	KADIAN	MAY

20MG CAPSULE (SUSTAINED RELEASE)

02184435	KADIAN	BGP
09991311	KADIAN	MAY

50MG CAPSULE (SUSTAINED RELEASE)

02184443	KADIAN	BGP
09991312	KADIAN	MAY

100MG CAPSULE (SUSTAINED RELEASE)

02184451	KADIAN	BGP
09991313	KADIAN	MAY

OXYCODONE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

10MG SUPPOSITORY

00392480	SUPEUDOL	SDZ
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20MG SUPPOSITORY

00392472	SUPEUDOL	SDZ
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5MG TABLET

02325950	OXYCODONE	PDL
02231934	OXY-IR	PFR
02319977	PMS-OXYCODONE	PMS
00789739	SUPEUDOL	SDZ

10MG TABLET

02325969	OXYCODONE	PDL
02240131	OXY-IR	PFR
02319985	PMS-OXYCODONE	PMS
00443948	SUPEUDOL	SDZ

20MG TABLET

02325977	OXYCODONE	PDL
02319993	PMS-OXYCODONE	PMS
02262983	SUPEUDOL	SDZ

20MG TABLET (IMMEDIATE RELEASE)

02240132	OXY-IR	PFR
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28:08.12 OPIATE PARTIAL AGONISTS

BUPRENORPHINE HYDROCHLORIDE, NALOXONE HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of opioid dependence when:

- The client must be 16 years or older.
- In cases where the client lives in a remote or isolated location, confirmation is required that the community has the ability to support buprenorphine/naloxone administration. These supports include the safe daily witnessing, storage and handling of the buprenorphine/naloxone doses. After this confirmation, NIHB will approve the buprenorphine/naloxone for the client.

2MG & 0.5MG TABLET

02453908	ACT BUPRENORPHINE/NALOXONE	ACG
02408090	MYLAN-BUPRENORPHINE/NALOXONE	MYL
02424851	PMS-BUPRENORPHINE-NALOXONE	PMS
02295695	SUBOXONE	IND

8MG & 2MG TABLET

02453916	ACT BUPRENORPHINE/NALOXONE	ACG
02408104	MYLAN-BUPRENORPHINE/NALOXONE	MYL
02424878	PMS-BUPRENORPHINE-NALOXONE	PMS
02295709	SUBOXONE	IND

28:08.92 MISCELLANEOUS ANALGESICS AND ANTIPYRETICS

ACETAMINOPHEN

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

ST **80MG/ML DROP**

01904140	ACETAMINOPHEN	TAN
01905864	ACETAMINOPHEN	TLI
02263793	PEDIAPHEN	EUR
02027801	PEDIATRIX	TEV
00875988	TEMPRA INFANT	PAL
02046059	TYLENOL	MCL

ST **16MG/ML LIQUID**

01905848	ACETAMINOPHEN	TLI
00792713	PDP-ACETAMINOPHEN	PED
02263807	PEDIAPHEN	EUR
00884553	TEMPRA CHILDREN'S	PAL

ST **32MG/ML LIQUID**

01901389	ACETAMINOPHEN	JMP
01958836	ACETAMINOPHEN	TLI
00792691	PDP-ACETAMINOPHEN	PED
02263831	PEDIAPHEN	EUR
02027798	PEDIATRIX	TEV
00875996	TEMPRA CHILDREN'S DOUBLE STRENGTH	PAL

28:08.92 MISCELLANEOUS ANALGESICS AND ANTIPYRETICS

ACETAMINOPHEN

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

ST 32MG/ML LIQUID		
02046040	TYLENOL	MCL
120MG SUPPOSITORY		
00553328	ABENOL	GSK
02230434	ACET 120	PED
02046660	PMS-ACETAMINOPHEN	PMS
160MG SUPPOSITORY		
02230435	ACET	PED
325MG SUPPOSITORY		
01919393	ABENOL	PED
02230436	ACET 325	PED
02046687	PMS-ACETAMINOPHEN	PMS
650MG SUPPOSITORY		
02230437	ACET 650	PED
02046695	PMS-ACETAMINOPHEN	PMS
ST 80MG TABLET		
02015676	ACETAMINOPHEN	TAN
02263815	PEDIAPHEN	EUR
ST 160MG TABLET		
02230934	ACETAMINOPHEN	TAN
ST 325MG TABLET		
00605751	ACETAMINOPHEN	VTH
00743542	ACETAMINOPHEN	PMT
00789801	ACETAMINOPHEN	TLI
01938088	ACETAMINOPHEN	JMP
02022214	ACÉTAMINOPHÈNE	RIV
02362198	ACÉTAMINOPHÈNE	RIV
00544981	APO ACETAMINOPHEN	APX
02229873	APO-ACETAMINOPHEN	APX
00389218	NOVO-GESIC	TEV
00559393	TYLENOL	MCL
00723894	TYLENOL	MCL
ST 500MG TABLET		
00549703	ACETAMINOPHEN	PMT
00605778	ACETAMINOPHEN	VTH
00789798	ACETAMINOPHEN	TLI
01939122	ACETAMINOPHEN	JMP
01962353	ACETAMINOPHEN	TAN
02252813	ACETAMINOPHEN	PMT
02255251	ACETAMINOPHEN	PMT
02022222	ACÉTAMINOPHÈNE	RIV
02362228	ACÉTAMINOPHÈNE	RIV
02362201	ACÉTAMINOPHÈNE BLASON SHIELD	RIV
00545007	APO ACETAMINOPHEN	APX
02229977	APO-ACETAMINOPHEN	APX

28:08.92 MISCELLANEOUS ANALGESICS AND ANTIPYRETICS

ACETAMINOPHEN

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

ST 500MG TABLET		
00013668	ATASOL FORTE	CHU
02355299	JAMP ACETAMINOPHEN BLAZON	JMP
00482323	NOVO-GESIC FORTE	TEV
00892505	PMS-ACETAMINOPHEN	PMS
00723908	TYLENOL	MCL
00559407	TYLENOL EXTRA STRENGTH	MCL
ST 80MG TABLET (CHEWABLE)		
01905856	ACETAMINOPHEN	TLI
02017458	ACETAMINOPHEN	RIV
02129957	ACETAMINOPHEN	VTH
ST 160MG TABLET (CHEWABLE)		
02017431	ACETAMINOPHEN	RIV
02142805	ACETAMINOPHEN	VTH
02263823	PEDIAPHEN	EUR
02347792	TYLENOL JR STRENGTH FASTMELTS	MCL
02241361	TYLENOL JUNIOR STRENGTH	MCL

FLOCTAFENINE

ST 200MG TABLET		
02244680	FLOCTAFENINE	AAP
ST 400MG TABLET		
02244681	FLOCTAFENINE	AAP

28:10.00 OPIATE ANTAGONISTS

NALOXONE HYDROCHLORIDE

INJECTION		
09991488	NALOXONE KIT	UNK
0.4MG/ML INJECTION		
09991460	NALOXONE KIT	UNK
0.4MG SOLUTION		
02453258	S.O.S NALOXONE HYDROCHLORIDE	SDZ
0.4MG/ML SOLUTION		
02148706	NALOXONE	SDZ
02382482	NALOXONE	TEL
02382601	NALOXONE	SDZ
02393034	NALOXONE	OMG
1MG/ML SOLUTION		
02148714	NALOXONE	SDZ
02393042	NALOXONE	OMG
4MG SPRAY		
02458187	NARCAN	UNK

28:10.00 OPIATE ANTAGONISTS

NALTREXONE HYDROCHLORIDE

50MG TABLET

02444275	APO-NALTREXONE	APX
02451883	NALTREXONE HYDROCHLORIDE	UNK
02213826	REVIA	TEV

28:12.04 ANTICONVULSANTS - BARBITURATES

PHENOBARBITAL

5MG/ML ELIXIR

00645575	PHENOBARB	PED
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100MG TABLET

00178829	PHENOBARB	PED
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PRIMIDONE

ST **125MG TABLET**

00399310	PRIMIDONE	AAP
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ST **250MG TABLET**

00396761	PRIMIDONE	AAP
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28:12.08 ANTICONVULSANTS - BENZODIAZEPINES

CLOBAZAM

ST **10MG TABLET**

02244638	APO-CLOBAZAM	APX
02244474	PMS-CLOBAZAM	PMS
02238334	TEVA-CLOBAZAM	TEV

CLONAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **0.25MG TABLET**

02442027	CLONAZEPAM	SIV
02179660	PMS-CLONAZEPAM	PMS

ST **0.5MG TABLET**

02177889	APO-CLONAZEPAM	APX
02230366	CLONAPAM	VAE
02442035	CLONAZEPAM	SIV
02048701	PMS-CLONAZEPAM	PMS
02207818	PMS-CLONAZEPAM-R	PMS
02311593	PRO-CLONAZEPAM	PDL
02242077	RIVA-CLONAZEPAM	RIV
00382825	RIVOTRIL	HLR
02239024	TEVA-CLONAZEPAM	TEV

ST **1MG TABLET**

02230368	CLONAPAM	VAE
02442043	CLONAZEPAM	SIV

28:12.08 ANTICONVULSANTS - BENZODIAZEPINES

CLONAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **1MG TABLET**

02048728	PMS-CLONAZEPAM	PMS
02311607	PRO-CLONAZEPAM	PDL

ST **2MG TABLET**

02177897	APO-CLONAZEPAM	APX
02230369	CLONAPAM	VAE
02442051	CLONAZEPAM	SIV
02048736	PMS-CLONAZEPAM	PMS
02311615	PRO-CLONAZEPAM	PDL
02242078	RIVA-CLONAZEPAM	RIV
00382841	RIVOTRIL	HLR
02239025	TEVA-CLONAZEPAM	TEV

ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503020	BENZODIAZEPINE ORAL LIQUID	UNK
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28:12.12 ANTICONVULSANTS - HYDANTOINS

PHENYTOIN

ST **30MG CAPSULE**

00022772	DILANTIN	PFI
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ST **100MG CAPSULE**

02460912	APO-PHENYTOIN SODIUM	APX
00022780	DILANTIN	PFI

ST **6MG/ML SUSPENSION**

00023442	DILANTIN	PFI
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ST **25MG/ML SUSPENSION**

00023450	DILANTIN	PFI
02250896	TARO-PHENYTOIN	TAR

ST **50MG TABLET**

00023698	DILANTIN INFATABS	PFI
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28:12.20 ANTICONVULSANTS - SUCCINIMIDES

ETHOSUXIMIDE

ST **250MG CAPSULE**

00022799	ZARONTIN	ERF
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ST **50MG/ML SYRUP**

00023485	ZARONTIN	ERF
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**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

BRIVARACETAM

Limited use benefit (prior approval required).

For adjunctive therapy in adult patients with refractory partial-onset seizures who meet all of the following criteria:

- Are under the care of a physician experienced in the treatment of epilepsy; AND
- Are currently receiving two or more antiepileptic medications; AND
- Have failed or demonstrated intolerance to at least two other antiepileptic medications; AND
- Are not receiving concurrent therapy with levetiracetam.

10MG TABLET

02452936 BRIVLERA UCB

25MG TABLET

02452944 BRIVLERA UCB

50MG TABLET

02452952 BRIVLERA UCB

75MG TABLET

02452960 BRIVLERA UCB

100MG TABLET

02452979 BRIVLERA UCB

CARBAMAZEPINE

ST **20MG/ML SUSPENSION**

02367394 TARO-CARBAMAZEPINE TAR

02194333 TEGRETOL NVR

ST **200MG TABLET**

00402699 APO CARBAMAZEPINE APX

00504742 MAZEPINE BMI

02407515 TARO-CARBAMAZEPINE TAR

00010405 TEGRETOL NVR

00782718 TEVA-CARBAMAZEPINE TEV

ST **100MG TABLET (CHEWABLE)**

02244403 TARO-CARBAMAZEPINE TAR

ST **200MG TABLET (CHEWABLE)**

02244404 TARO-CARBAMAZEPINE TAR

ST **200MG TABLET (EXTENDED RELEASE)**

02413590 CARBAMAZEPINE PDL

02238222 DOM-CARBAMAZEPINE DPC

02231543 PMS-CARBAMAZEPINE PMS

02261839 SANDOZ-CARBAMAZEPINE SDZ

02237907 TARO-CARBAMAZEPINE TAR

00773611 TEGRETOL NVR

ST **400MG TABLET (EXTENDED RELEASE)**

02413604 CARBAMAZEPINE PDL

02238223 DOM-CARBAMAZEPINE DPC

02231544 PMS-CARBAMAZEPINE PMS

02261847 SANDOZ-CARBAMAZEPINE SDZ

02237908 TARO-CARBAMAZEPINE TAR

00755583 TEGRETOL NVR

**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

ESLICARBAZEPINE ACETATE

Limited use benefit (prior approval required).

For adjunctive therapy in adult patients with refractory partial-onset seizures who meet all of the following criteria:

- Are under the care of a physician experienced in the treatment of epilepsy; AND
- Are currently receiving two or more antiepileptic medications; AND
- Have failed or demonstrated intolerance to at least two other antiepileptic medications.

ST **200MG TABLET**

02426862 APTIOM SPC

ST **400MG TABLET**

02426870 APTIOM SPC

ST **600MG TABLET**

02426889 APTIOM SPC

ST **800MG TABLET**

02426897 APTIOM SPC

GABAPENTIN

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on gabapentin. The limit accumulates against the amount of gabapentin claimed to the program. A total of 400 grams of gabapentin is permitted in a 100-day period, for a total daily dose of 4000 mg/day.

ST **100MG CAPSULE**

02244304 APO-GABAPENTIN APX

02321203 AURO-GABAPENTIN AUR

02243743 DOM-GABAPENTIN DPC

02246314 GABAPENTIN SIV

02353245 GABAPENTIN SAN

02416840 GABAPENTIN ACC

02285819 GD-GABAPENTIN PFI

02361469 JAMP-GABAPENTIN JMP

02391473 MAR-GABAPENTIN MAR

02084260 NEURONTIN PFI

02243446 PMS-GABAPENTIN PMS

02310449 PRO-GABAPENTIN PDL

02319055 RAN-GABAPENTIN RBY

02251167 RIVA-GABAPENTIN RIV

02244513 TEVA-GABAPENTIN TEV

ST **300MG CAPSULE**

02244305 APO-GABAPENTIN APX

02321211 AURO-GABAPENTIN AUR

02243744 DOM-GABAPENTIN DPC

02246315 GABAPENTIN SIV

02353253 GABAPENTIN SAN

02416859 GABAPENTIN ACC

02285827 GD-GABAPENTIN PFI

02361485 JAMP-GABAPENTIN JMP

02391481 MAR-GABAPENTIN MAR

02084279 NEURONTIN PFI

02243447 PMS-GABAPENTIN PMS

02310457 PRO-GABAPENTIN PDL

02319063 RAN-GABAPENTIN RBY

**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

GABAPENTIN

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on gabapentin. The limit accumulates against the amount of gabapentin claimed to the program. A total of 400 grams of gabapentin is permitted in a 100-day period, for a total daily dose of 4000 mg/day.

ST **300MG CAPSULE**

02251175	RIVA-GABAPENTIN	RIV
02244514	TEVA-GABAPENTIN	TEV

ST **400MG CAPSULE**

02244306	APO-GABAPENTIN	APX
02321238	AURO-GABAPENTIN	AUR
02243745	DOM-GABAPENTIN	DPC
02246316	GABAPENTIN	SIV
02353261	GABAPENTIN	SAN
02416867	GABAPENTIN	ACC
02361493	JAMP-GABAPENTIN	JMP
02391503	MAR-GABAPENTIN	MAR
02084287	NEURONTIN	PFI
02243448	PMS-GABAPENTIN	PMS
02310465	PRO-GABAPENTIN	PDL
02319071	RAN-GABAPENTIN	RBV
02251183	RIVA-GABAPENTIN	RIV
02244515	TEVA-GABAPENTIN	TEV

ST **600MG TABLET**

02293358	APO-GABAPENTIN	APX
02388200	GABAPENTIN	SIV
02392526	GABAPENTIN	ACC
02431289	GABAPENTIN	SAN
02285843	GD-GABAPENTIN	PFI
02402289	JAMP-GABAPENTIN	JMP
02239717	NEURONTIN	PFI
02255898	PMS-GABAPENTIN	PMS
02310473	PRO-GABAPENTIN	PDL
02259796	RIVA-GABAPENTIN	RIV
02248457	TEVA-GABAPENTIN	TEV

ST **800MG TABLET**

02293366	APO-GABAPENTIN	APX
02388219	GABAPENTIN	SIV
02392534	GABAPENTIN	ACC
02431297	GABAPENTIN	SAN
02402297	JAMP-GABAPENTIN	JMP
02239718	NEURONTIN	PFI
02255901	PMS-GABAPENTIN	PMS
02310481	PRO-GABAPENTIN	PDL
02259818	RIVA-GABAPENTIN	RIV
02247346	TEVA-GABAPENTIN	TEV

ST **600MG TABLET (IMMEDIATE RELEASE)**

02410990	GLN-GABAPENTIN	GLK
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ST **800MG TABLET (IMMEDIATE RELEASE)**

02411008	GLN-GABAPENTIN	GLK
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**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

LACOSAMIDE

Limited use benefit (prior approval required).

For adjunctive therapy in adult patients with refractory partial-onset seizures who meet all of the following criteria:

- Are under the care of a physician experienced in the treatment of epilepsy; AND
- Are currently receiving two or more antiepileptic medications; AND
- Have failed or demonstrated intolerance to at least two other antiepileptic medications.

ST **50MG TABLET**

02475332	AURO-LACOSAMIDE	AUR
02478196	PHARMA-LACOSAMIDE	PMS
02474670	SANDOZ LACOSAMIDE	SDZ
02472902	TEVA-LACOSAMIDE	TEV
02357615	VIMPAT	UCB

ST **100MG TABLET**

02475340	AURO-LACOSAMIDE	AUR
02478218	PHARMA-LACOSAMIDE	PMS
02474689	SANDOZ LACOSAMIDE	SDZ
02472910	TEVA-LACOSAMIDE	TEV
02357623	VIMPAT	UCB

ST **150MG TABLET**

02475359	AURO-LACOSAMIDE	AUR
02478226	PHARMA-LACOSAMIDE	PMS
02474697	SANDOZ LACOSAMIDE	SDZ
02472929	TEVA-LACOSAMIDE	TEV
02357631	VIMPAT	UCB

ST **200MG TABLET**

02475367	AURO-LACOSAMIDE	AUR
02478234	PHARMA-LACOSAMIDE	PMS
02474700	SANDOZ LACOSAMIDE	SDZ
02472937	TEVA-LACOSAMIDE	TEV
02357658	VIMPAT	UCB

LAMOTRIGINE

ST **2MG TABLET**

02243803	LAMICTAL	GSK
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ST **5MG TABLET**

02240115	LAMICTAL	GSK
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ST **25MG TABLET**

02245208	APO-LAMOTRIGINE	APX
02381354	AURO-LAMOTRIGINE	AUR
02142082	LAMICTAL	GSK
02302969	LAMOTRIGINE	PDL
02343010	LAMOTRIGINE	SAN
02428202	LAMOTRIGINE	SIV
02265494	MYLAN-LAMOTRIGINE	MYL
02246897	PMS-LAMOTRIGINE	PMS
02248232	TEVA-LAMOTRIGINE	TEV

ST **100MG TABLET**

02245209	APO-LAMOTRIGINE	APX
02381362	AURO-LAMOTRIGINE	AUR
02142104	LAMICTAL	GSK
02302985	LAMOTRIGINE	PDL
02343029	LAMOTRIGINE	SAN

**28:12.92 MISCELLANEOUS
ANTICONSULSANTS**

LAMOTRIGINE

ST 100MG TABLET

02428210	LAMOTRIGINE	SIV
02265508	MYLAN-LAMOTRIGINE	MYL
02246898	PMS-LAMOTRIGINE	PMS
02248233	TEVA-LAMOTRIGINE	TEV

ST 150MG TABLET

02245210	APO-LAMOTRIGINE	APX
02381370	AURO-LAMOTRIGINE	AUR
02142112	LAMICTAL	GSK
02302993	LAMOTRIGINE	PDL
02343037	LAMOTRIGINE	SAN
02428229	LAMOTRIGINE	SIV
02265516	MYLAN-LAMOTRIGINE	MYL
02246899	PMS-LAMOTRIGINE	PMS
02248234	TEVA-LAMOTRIGINE	TEV

LEVETIRACETAM

ST 250MG TABLET

02274183	ACT LEVETIRACETAM	ACG
02285924	APO-LEVETIRACETAM	APX
02375249	AURO-LEVETIRACETAM	AUR
02450348	BIO-LEVETIRACETAM	BMI
02403005	JAMP-LEVETIRACETAM	JMP
02247027	KEPPRA	UCB
02353342	LEVETIRACETAM	SAN
02399776	LEVETIRACETAM	ACC
02442531	LEVETIRACETAM	SIV
02454653	LEVETIRACETAM	PMS
02474468	LEVETIRACETAM	RIV
02440202	NAT-LEVETIRACETAM	NPH
02296101	PMS-LEVETIRACETAM	PMS
02396106	RAN-LEVETIRACETAM	RBV
02461986	SANDOZ LEVETIRACETAM	SDZ

ST 500MG TABLET

02274191	ACT LEVETIRACETAM	ACG
02285932	APO-LEVETIRACETAM	APX
02375257	AURO-LEVETIRACETAM	AUR
02450356	BIO-LEVETIRACETAM	BMI
02297418	DOM-LEVETIRACETAM	DPC
02403021	JAMP-LEVETIRACETAM	JMP
02247028	KEPPRA	UCB
02353350	LEVETIRACETAM	SAN
02399784	LEVETIRACETAM	ACC
02442558	LEVETIRACETAM	SIV
02454661	LEVETIRACETAM	PMS
02474476	LEVETIRACETAM	RIV
02440210	NAT-LEVETIRACETAM	NPH
02296128	PMS-LEVETIRACETAM	PMS
02311380	PRO-LEVETIRACETAM	PDL
02396114	RAN-LEVETIRACETAM	RBV
02461994	SANDOZ LEVETIRACETAM	SDZ

ST 750MG TABLET

02274205	ACT LEVETIRACETAM	ACG
02285940	APO-LEVETIRACETAM	APX

**28:12.92 MISCELLANEOUS
ANTICONSULSANTS**

LEVETIRACETAM

ST 750MG TABLET

02375265	AURO-LEVETIRACETAM	AUR
02450364	BIO-LEVETIRACETAM	BMI
02403048	JAMP-LEVETIRACETAM	JMP
02247029	KEPPRA	UCB
02353369	LEVETIRACETAM	SAN
02399792	LEVETIRACETAM	ACC
02442566	LEVETIRACETAM	SIV
02454688	LEVETIRACETAM	PMS
02474484	LEVETIRACETAM	RIV
02440229	NAT-LEVETIRACETAM	NPH
02296136	PMS-LEVETIRACETAM	PMS
02311399	PRO-LEVETIRACETAM	PDL
02396122	RAN-LEVETIRACETAM	RBV
02462001	SANDOZ LEVETIRACETAM	SDZ

PDIN FOR EXTEMPORANEOUS MIXTURE

99503026	LEVETIRACETAM ORAL LIQUID	UNK
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OXCARBAZEPINE

150MG TABLET

02284294	APO-OXCARBAZEPINE	APX
02348381	APX-OXCARBAZEPINE	APX
02440717	JAMP-OXCARBAZEPINE	JMP

300MG TABLET

02284308	APO-OXCARBAZEPINE	APX
02348403	APX-OXCARBAZEPINE	APX
02440725	JAMP-OXCARBAZEPINE	JMP
02242068	TRILEPTAL	NVR

600MG TABLET

02284316	APO-OXCARBAZEPINE	APX
02348411	APX-OXCARBAZEPINE	APX
02440733	JAMP-OXCARBAZEPINE	JMP
02242069	TRILEPTAL	NVR

OXCARBAZEPINE (SUSPENSION)

Limited use benefit (prior approval is not required).

For patients 19 years of age or over who are unable to swallow the tablet formulation due to:

- Tube feeding; OR
- Severe dysphagia

Note:

Trileptal (oxcarbazepine) suspension is an open benefit for patients 18 years of age and under and does not require prior approval for these patients.

Oxcarbazepine tablets are an open benefit for patients of all ages and do not require prior approval.

60MG SUSPENSION

02244673	TRILEPTAL	NVR
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**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

PREGABALIN

Limited use benefit (prior approval required).

For the treatment of neuropathic pain in patients who have failed to effectively treat their pain with a tricyclic antidepressant (TCA);

OR

For the treatment of neuropathic pain in patients who have a contraindication or intolerance to a tricyclic antidepressant (TCA).

Coverage is limited to a maximum of 600mg per day.

ST **25MG CAPSULE**

02402912	ACT PREGABALIN	ACG
02394235	APO-PREGABALIN	APX
02433869	AURO-PREGABALIN	AUR
02402556	DOM-PREGABALIN	DPC
02435977	JAMP-PREGABALIN	JMP
02268418	LYRICA	PFI
02417529	MAR-PREGABALIN	MAR
02423804	MINT-PREGABALIN	MIN
02467291	M-PREGABALIN	MAN
02479117	NRA-PREGABALIN	UNK
02359596	PMS-PREGABALIN	PMS
02396483	PREGABALIN	PDL
02403692	PREGABALIN	SIV
02405539	PREGABALIN	SAN
02476304	PREGABALIN	RIV
02392801	RAN-PREGABALIN	RBV
02377039	RIVA-PREGABALIN	RIV
02390817	SANDOZ PREGABALIN	SDZ
02361159	TEVA-PREGABALIN	TEV

ST **50MG CAPSULE**

02402920	ACT PREGABALIN	ACG
02394243	APO-PREGABALIN	APX
02433877	AURO-PREGABALIN	AUR
02402564	DOM-PREGABALIN	DPC
02435985	JAMP-PREGABALIN	JMP
02268426	LYRICA	PFI
02417537	MAR-PREGABALIN	MAR
02423812	MINT-PREGABALIN	MIN
02467305	M-PREGABALIN	MAN
02479125	NRA-PREGABALIN	UNK
02359618	PMS-PREGABALIN	PMS
02396505	PREGABALIN	PDL
02403706	PREGABALIN	SIV
02405547	PREGABALIN	SAN
02476312	PREGABALIN	RIV
02392828	RAN-PREGABALIN	RBV
02377047	RIVA-PREGABALIN	RIV
02390825	SANDOZ PREGABALIN	SDZ
02361175	TEVA-PREGABALIN	TEV

ST **75MG CAPSULE**

02402939	ACT PREGABALIN	ACG
02394251	APO-PREGABALIN	APX
02433885	AURO-PREGABALIN	AUR
02402572	DOM-PREGABALIN	DPC

**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

PREGABALIN

Limited use benefit (prior approval required).

For the treatment of neuropathic pain in patients who have failed to effectively treat their pain with a tricyclic antidepressant (TCA);

OR

For the treatment of neuropathic pain in patients who have a contraindication or intolerance to a tricyclic antidepressant (TCA).

Coverage is limited to a maximum of 600mg per day.

ST **75MG CAPSULE**

02435993	JAMP-PREGABALIN	JMP
02268434	LYRICA	PFI
02417545	MAR-PREGABALIN	MAR
02424185	MINT-PREGABALIN	MIN
02467313	M-PREGABALIN	MAN
02479133	NRA-PREGABALIN	UNK
02359626	PMS-PREGABALIN	PMS
02396513	PREGABALIN	PDL
02403714	PREGABALIN	SIV
02405555	PREGABALIN	SAN
02476320	PREGABALIN	RIV
02392836	RAN-PREGABALIN	RBV
02377055	RIVA-PREGABALIN	RIV
02390833	SANDOZ PREGABALIN	SDZ
02361183	TEVA-PREGABALIN	TEV

ST **150MG CAPSULE**

02402955	ACT PREGABALIN	ACG
02394278	APO-PREGABALIN	APX
02433907	AURO-PREGABALIN	AUR
02402580	DOM-PREGABALIN	DPC
02436000	JAMP-PREGABALIN	JMP
02268450	LYRICA	PFI
02417561	MAR-PREGABALIN	MAR
02424207	MINT-PREGABALIN	MIN
02467321	M-PREGABALIN	MAN
02479168	NRA-PREGABALIN	UNK
02359634	PMS-PREGABALIN	PMS
02396521	PREGABALIN	PDL
02403722	PREGABALIN	SIV
02405563	PREGABALIN	SAN
02476347	PREGABALIN	RIV
02392844	RAN-PREGABALIN	RBV
02377063	RIVA-PREGABALIN	RIV
02390841	SANDOZ PREGABALIN	SDZ
02361205	TEVA-PREGABALIN	TEV

ST **300MG CAPSULE**

02402998	ACT PREGABALIN	ACG
02394294	APO-PREGABALIN	APX
02436019	JAMP-PREGABALIN	JMP
02268485	LYRICA	PFI
02359642	PMS-PREGABALIN	PMS
02396548	PREGABALIN	PDL
02403730	PREGABALIN	SIV
02405598	PREGABALIN	SAN

**28:12.92 MISCELLANEOUS
ANTICONSULSANTS**

PREGABALIN

Limited use benefit (prior approval required).

For the treatment of neuropathic pain in patients who have failed to effectively treat their pain with a tricyclic antidepressant (TCA);

OR

For the treatment of neuropathic pain in patients who have a contraindication or intolerance to a tricyclic antidepressant (TCA).

Coverage is limited to a maximum of 600mg per day.

ST 300MG CAPSULE

02476371	PREGABALIN	RIV
02392860	RAN-PREGABALIN	RBY
02377071	RIVA-PREGABALIN	RIV
02390868	SANDOZ PREGABALIN	SDZ
02361248	TEVA-PREGABALIN	TEV

RUFINAMIDE

Limited use benefit (prior approval required).

- For the adjunctive treatment of seizures associated with Lennox-Gastaut syndrome in adults and children 4 years and older when prescribed by a neurologist or experienced specialist.
- Patient has failed or is intolerant to or has contraindications to at least two adjunctive antiepileptic drugs.

ST 100MG TABLET

02369613	BANZEL	EIS
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ST 200MG TABLET

02369621	BANZEL	EIS
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ST 400MG TABLET

02369648	BANZEL	EIS
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TOPIRAMATE

ST 15MG CAPSULE

02239907	TOPAMAX	JSO
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ST 25MG CAPSULE

02239908	TOPAMAX	JSO
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ST 25MG TABLET

02351307	ACCEL-TOPIRAMATE	ACP
02279614	APO-TOPIRAMATE	APX
02345803	AURO-TOPIRAMATE	APL
02271141	DOM-TOPIRAMATE	DPC
02287765	GLN-TOPIRAMATE	GLK
02435608	JAMP-TOPIRAMATE	JMP
02432099	MAR-TOPIRAMATE	MAR
02315645	MINT-TOPIRAMATE	MIN
02263351	MYLAN-TOPIRAMATE	MYL
02262991	PMS-TOPIRAMATE	PMS
02313650	PRO-TOPIRAMATE	PDL
02396076	RAN-TOPIRAMATE	RBY
02260050	SANDOZ TOPIRAMATE	SDZ
02431807	SANDOZ TOPIRAMATE	SDZ
02248860	TEVA-TOPIRAMATE	TEV
02230893	TOPAMAX	JSO
02356856	TOPIRAMATE	SAN
02389460	TOPIRAMATE	SIV

**28:12.92 MISCELLANEOUS
ANTICONSULSANTS**

TOPIRAMATE

ST 25MG TABLET

02395738	TOPIRAMATE	ACC
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ST 50MG TABLET

02312085	PMS-TOPIRAMATE	PMS
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ST 100MG TABLET

02351315	ACCEL-TOPIRAMATE	ACP
02279630	APO-TOPIRAMATE	APX
02345838	AURO-TOPIRAMATE	APL
02271168	DOM-TOPIRAMATE	DPC
02287773	GLN-TOPIRAMATE	GLK
02435616	JAMP-TOPIRAMATE	JMP
02432102	MAR-TOPIRAMATE	MAR
02315653	MINT-TOPIRAMATE	MIN
02263378	MYLAN-TOPIRAMATE	MYL
02263009	PMS-TOPIRAMATE	PMS
02313669	PRO-TOPIRAMATE	PDL
02396084	RAN-TOPIRAMATE	RBY
02260069	SANDOZ TOPIRAMATE	SDZ
02431815	SANDOZ TOPIRAMATE	SDZ
02248861	TEVA-TOPIRAMATE	TEV
02230894	TOPAMAX	JSO
02356864	TOPIRAMATE	SAN
02389487	TOPIRAMATE	SIV
02395746	TOPIRAMATE	ACC

ST 200MG TABLET

02351323	ACCEL-TOPIRAMATE	ACP
02279649	APO-TOPIRAMATE	APX
02345846	AURO-TOPIRAMATE	APL
02271176	DOM-TOPIRAMATE	DPC
02287781	GLN-TOPIRAMATE	GLK
02435624	JAMP-TOPIRAMATE	JMP
02432110	MAR-TOPIRAMATE	MAR
02315661	MINT-TOPIRAMATE	MIN
02263386	MYLAN-TOPIRAMATE	MYL
02263017	PMS-TOPIRAMATE	PMS
02313677	PRO-TOPIRAMATE	PDL
02396092	RAN-TOPIRAMATE	RBY
02267837	SANDOZ TOPIRAMATE	SDZ
02431823	SANDOZ TOPIRAMATE	SDZ
02248862	TEVA-TOPIRAMATE	TEV
02230896	TOPAMAX	JSO
02356872	TOPIRAMATE	SAN
02395754	TOPIRAMATE	ACC

PDIN FOR EXTEMPORANEOUS MIXTURE

99503027	TOPIRAMATE ORAL LIQUID	UNK
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VALPROIC ACID (DIVALPROEX SODIUM)

ST 125MG TABLET (ENTERIC COATED)

02239698	APO-DIVALPROEX	APX
02400499	DIVALPROEX	SAN
00596418	EPIVAL	BGP
02458926	MYLAN-DIVALPROEX	MYL
02244138	PMS-DIVALPROEX	PMS
02239701	TEVA-DIVALPROEX	TEV

**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

VALPROIC ACID (DIVALPROEX SODIUM)

ST 250MG TABLET (ENTERIC COATED)

02239699	APO-DIVALPROEX	APX
02400502	DIVALPROEX	SAN
00596426	EPIVAL	BGP
02458934	MYLAN-DIVALPROEX	MYL
02244139	PMS-DIVALPROEX	PMS
02239702	TEVA-DIVALPROEX	TEV

ST 500MG TABLET (ENTERIC COATED)

02239700	APO-DIVALPROEX	APX
02400510	DIVALPROEX	SAN
00596434	EPIVAL	BGP
02459019	MYLAN-DIVALPROEX	MYL
02244140	PMS-DIVALPROEX	PMS
02239703	TEVA-DIVALPROEX	TEV

VALPROIC ACID (SODIUM VALPROATE)

ST 250MG CAPSULE

02238048	APO-VALPROIC	APX
02231030	DOM-VALPROIC ACID	DPC
02230768	PMS-VALPROIC ACID	PMS
02239714	SANDOZ VALPROIC	SDZ

ST 500MG CAPSULE (ENTERIC COATED)

02231031	DOM-VALPROIC ACID	DPC
02229628	PMS-VALPROIC ACID	PMS

ST 50MG/ML SOLUTION

02238817	DOM-VALPROIC ACID	DPC
02236807	PMS-VALPROIC ACID	PMS

ST 50MG/ML SYRUP

02238370	APO-VALPROIC	APX
00443832	DEPAKENE	BGP

VIGABATRIN

ST 500MG POWDER FOR SOLUTION

02068036	SABRIL	LUK
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ST 500MG TABLET

02065819	SABRIL	LUK
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28:16.04 ANTIDEPRESSANTS

AMITRIPTYLINE HYDROCHLORIDE

ST 10MG TABLET

00370991	AMITRIPTYLINE	PDL
02403137	APO-AMITRIPTYLINE	APX
00335053	ELAVIL	AAP
02435527	JAMP-AMITRIPTYLINE	JMP
00293911	LEVATE	BMI
02429861	MAR-AMITRIPTYLINE	MAR
00654523	PMS-AMITRIPTYLINE	PMS
02326043	TEVA-AMITRIPTYLINE	TEV

ST 25MG TABLET

00371009	AMITRIPTYLINE	PDL
02403145	APO-AMITRIPTYLINE	APX
00335061	ELAVIL	AAP
02435535	JAMP-AMITRIPTYLINE	JMP
02429888	MAR-AMITRIPTYLINE	MAR
00654515	PMS-AMITRIPTYLINE	PMS

28:16.04 ANTIDEPRESSANTS

AMITRIPTYLINE HYDROCHLORIDE

ST 25MG TABLET

02326051	TEVA-AMITRIPTYLINE	TEV
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ST 50MG TABLET

00456349	AMITRIPTYLINE	PDL
02403153	APO-AMITRIPTYLINE	APX
00335088	ELAVIL	AAP
02435543	JAMP-AMITRIPTYLINE	JMP
00271152	LEVATE	BMI
02429896	MAR-AMITRIPTYLINE	MAR
00654507	PMS-AMITRIPTYLINE	PMS
02326078	TEVA-AMITRIPTYLINE	TEV

ST 75MG TABLET

02403161	APO-AMITRIPTYLINE	APX
00754129	ELAVIL	AAP
02435551	JAMP-AMITRIPTYLINE	JMP
00405612	LEVATE	BMI
02429918	MAR-AMITRIPTYLINE	MAR

BUPROPION HYDROCHLORIDE (WELLBUTRIN)

ST 100MG TABLET (EXTENDED RELEASE)

02331616	BUPROPION SR	PDL
02391562	BUPROPION SR	SAN
02325373	PMS-BUPROPION SR	PMS
02275074	SANDOZ BUPROPION SR	SDZ

ST 150MG TABLET (EXTENDED RELEASE)

02439654	ACT BUPROPION XL	ACG
02325357	BUPROPION SR	PDL
02391570	BUPROPION SR	SAN
02382075	MYLAN-BUPROPION XL	MYL
02313421	PMS-BUPROPION SR	PMS
02275082	SANDOZ BUPROPION SR	SDZ
02237825	WELLBUTRIN SR	VAE
02275090	WELLBUTRIN XL	VAE

ST 300MG TABLET (EXTENDED RELEASE)

02439662	ACT BUPROPION XL	ACG
02382083	MYLAN-BUPROPION XL	MYL
02275104	WELLBUTRIN XL	VAE

BUPROPION HYDROCHLORIDE (ZYBAN)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 180 tablets during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached the client is eligible again for coverage for bupropion hydrochloride when one year has elapsed from the day the initial prescription was filled.

ST 150MG TABLET (EXTENDED RELEASE)

02238441	ZYBAN	VAE
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CITALOPRAM HYDROBROMIDE

ST 10MG TABLET

02355248	ACCEL-CITALOPRAM	ACP
02374617	AG-CITALOPRAM	ANG
02448475	BIO-CITALOPRAM	BMI
02325047	CITALOPRAM	PDL

**28:16.04 ANTIDEPRESSANTS
CITALOPRAM HYDROBROMIDE**

ST 10MG TABLET

02387948	CITALOPRAM	SIV
02430517	CITALOPRAM	JMP
02445719	CITALOPRAM	SAN
02273055	DOM-CITALOPRAM	DPC
02370085	JAMP-CITALOPRAM	JMP
02371871	MAR-CITALOPRAM	MAR
02429691	MINT-CITALOPRAM	MIN
02409003	NAT-CITALOPRAM	NPH
02477637	NRA-CITALOPRAM	UNK
02270609	PMS-CITALOPRAM	PMS
02303256	RIVA-CITALOPRAM	RIV
02431629	SEPTA-CITALOPRAM	SPT
02312336	TEVA-CITALOPRAM	TEV

ST 20MG TABLET

02355256	ACCEL-CITALOPRAM	ACP
02248050	ACT CITALOPRAM	TEV
02339390	AG-CITALOPRAM	ANG
02246056	APO-CITALOPRAM	APX
02275562	AURO-CITALOPRAM	AUR
02448491	BIO-CITALOPRAM	BMI
02239607	CELEXA	LUD
02257513	CITALOPRAM	PDL
02353660	CITALOPRAM	SAN
02387956	CITALOPRAM	SIV
02430541	CITALOPRAM	JMP
02248942	DOM-CITALOPRAM	DPC
02313405	JAMP-CITALOPRAM	JMP
02371898	MAR-CITALOPRAM	MAR
02429705	MINT-CITALOPRAM	MIN
02409011	NAT-CITALOPRAM	NPH
02477645	NRA-CITALOPRAM	UNK
02248010	PMS-CITALOPRAM	PMS
02285622	RAN-CITALO	RBV
02303264	RIVA-CITALOPRAM	RIV
02248170	SANDOZ CITALOPRAM	SDZ
02355272	SEPTA-CITALOPRAM	SPT
02293218	TEVA-CITALOPRAM	TEV

ST 30MG TABLET

02296152	CTP 30	SPC
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ST 40MG TABLET

02355264	ACCEL-CITALOPRAM	ACP
02248051	ACT CITALOPRAM	TEV
02339404	AG-CITALOPRAM	ANG
02246057	APO-CITALOPRAM	APX
02275570	AURO-CITALOPRAM	AUR
02448513	BIO-CITALOPRAM	BMI
02239608	CELEXA	LUD
02257521	CITALOPRAM	PDL
02353679	CITALOPRAM	SAN
02387964	CITALOPRAM	SIV
02430568	CITALOPRAM	JMP
02248943	DOM-CITALOPRAM	DPC
02313413	JAMP-CITALOPRAM	JMP
02371901	MAR-CITALOPRAM	MAR

**28:16.04 ANTIDEPRESSANTS
CITALOPRAM HYDROBROMIDE**

ST 40MG TABLET

02429713	MINT-CITALOPRAM	MIN
02409038	NAT-CITALOPRAM	NPH
02477653	NRA-CITALOPRAM	UNK
02248011	PMS-CITALOPRAM	PMS
02285630	RAN-CITALO	RBV
02303272	RIVA-CITALOPRAM	RIV
02248171	SANDOZ CITALOPRAM	SDZ
02355280	SEPTA-CITALOPRAM	SPT
02293226	TEVA-CITALOPRAM	TEV

CLOMIPRAMINE HYDROCHLORIDE

ST 10MG TABLET

00330566	ANAFRANIL	AAP
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ST 25MG TABLET

00324019	ANAFRANIL	AAP
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ST 50MG TABLET

00402591	ANAFRANIL	AAP
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DESIPRAMINE HYDROCHLORIDE

ST 10MG TABLET

02216248	DESIPRAMINE	AAP
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ST 25MG TABLET

02216256	DESIPRAMINE	AAP
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ST 50MG TABLET

02216264	DESIPRAMINE	AAP
01946277	PMS DESIPRAMINE	PMS

ST 75MG TABLET

02216272	DESIPRAMINE	AAP
01946242	PMS DESIPRAMINE	PMS

ST 100MG TABLET

02216280	DESIPRAMINE	AAP
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DOXEPIN HYDROCHLORIDE

ST 10MG CAPSULE

02049996	DOXEPIN	APX
00024325	SINEQUAN	AAP

ST 25MG CAPSULE

02050005	DOXEPIN	APX
00024333	SINEQUAN	AAP

ST 50MG CAPSULE

02050013	DOXEPIN	APX
00024341	SINEQUAN	AAP

ST 75MG CAPSULE

02050021	DOXEPIN	APX
00400750	SINEQUAN	AAP

ST 100MG CAPSULE

02050048	DOXEPIN	APX
00326925	SINEQUAN	AAP

ST 150MG CAPSULE

02050056	DOXEPIN	APX
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DULOXETINE HYDROCHLORIDE

30MG CAPSULE (DELAYED RELEASE)

02475308	AG-DULOXETINE	ANG
02440423	APO-DULOXETINE	APX
02436647	AURO-DULOXETINE	AUR

28:16.04 ANTIDEPRESSANTS

DULOXETINE HYDROCHLORIDE

30MG CAPSULE (DELAYED RELEASE)

02301482	CYMBALTA	LIL
02452650	DULOXETINE	PDL
02453630	DULOXETINE	SIV
02437082	DULOXETINE DR	TEV
02451913	JAMP-DULOXETINE	JMP
02446081	MAR-DULOXETINE	MAR
02473208	M-DULOXETINE	MAN
02438984	MINT-DULOXETINE	MIN
02429446	PMS-DULOXETINE	PMS
02438259	RAN-DULOXETINE	RBV
02451077	RIVA-DULOXETINE	RIV
02439948	SANDOZ DULOXETINE	SDZ

60MG CAPSULE (DELAYED RELEASE)

02475316	AG-DULOXETINE	ANG
02440431	APO-DULOXETINE	APX
02436655	AURO-DULOXETINE	AUR
02301490	CYMBALTA	LIL
02452669	DULOXETINE	PDL
02453649	DULOXETINE	SIV
02437090	DULOXETINE DR	TEV
02451921	JAMP-DULOXETINE	JMP
02446103	MAR-DULOXETINE	MAR
02473216	M-DULOXETINE	MAN
02438992	MINT-DULOXETINE	MIN
02429454	PMS-DULOXETINE	PMS
02438267	RAN-DULOXETINE	RBV
02451085	RIVA-DULOXETINE	RIV
02439956	SANDOZ DULOXETINE	SDZ

ESCITALOPRAM OXALATE

ST **10MG TABLET**

02434652	ACH-ESCITALOPRAM	ACC
02295016	APO-ESCITALOPRAM	APX
02397358	AURO-ESCITALOPRAM	AUR
02263238	CIPRALEX	LUD
02424401	ESCITALOPRAM	PDL
02429039	ESCITALOPRAM	SIV
02430118	ESCITALOPRAM	SAN
02429780	JAMP-ESCITALOPRAM	JMP
02423480	MAR-ESCITALOPRAM	MAR
02407418	MINT-ESCITALOPRAM	MIN
02309467	MYLAN-ESCITALOPRAM	MYL
02440296	NAT-ESCITALOPRAM	NPH
02476851	NRA-ESCITALOPRAM	UNK
02469243	PHARMA-ESCITALOPRAM	PMS
02303949	PMS-ESCITALOPRAM	PMS
02426331	PRIVA-ESCITALOPRAM	PHA
02385481	RAN-ESCITALOPRAM	RBV
02428830	RIVA-ESCITALOPRAM	RIV
02364077	SANDOZ ESCITALOPRAM	SDZ
02318180	TEVA-ESCITALOPRAM	TEV

ST **20MG TABLET**

02434660	ACH-ESCITALOPRAM	ACC
02295024	APO-ESCITALOPRAM	APX
02397374	AURO-ESCITALOPRAM	AUR

28:16.04 ANTIDEPRESSANTS

ESCITALOPRAM OXALATE

ST **20MG TABLET**

02263254	CIPRALEX	LUD
02424428	ESCITALOPRAM	PDL
02429047	ESCITALOPRAM	SIV
02430126	ESCITALOPRAM	SAN
02429799	JAMP-ESCITALOPRAM	JMP
02423502	MAR-ESCITALOPRAM	MAR
02407434	MINT-ESCITALOPRAM	MIN
02309475	MYLAN-ESCITALOPRAM	MYL
02440318	NAT-ESCITALOPRAM	NPH
02476878	NRA-ESCITALOPRAM	UNK
02469251	PHARMA-ESCITALOPRAM	PMS
02303965	PMS-ESCITALOPRAM	PMS
02426358	PRIVA-ESCITALOPRAM	PHA
02385503	RAN-ESCITALOPRAM	RBV
02428857	RIVA-ESCITALOPRAM	RIV
02364085	SANDOZ ESCITALOPRAM	SDZ
02318202	TEVA-ESCITALOPRAM	TEV

ST **10MG TABLET (ORALLY DISINTEGRATING)**

02454297	ACT ESCITALOPRAM ODT	ACC
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ST **20MG TABLET (ORALLY DISINTEGRATING)**

02454300	ACT ESCITALOPRAM ODT	ACC
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FLUOXETINE HYDROCHLORIDE

ST **10MG CAPSULE**

02393441	ACH-FLUOXETINE	ACC
02242177	ACT FLUOXETINE	REC
02216353	APO-FLUOXETINE	APX
02385627	AURO-FLUOXETINE	AUR
02448424	BIO-FLUOXETINE	BMI
02177617	DOM-FLUOXETINE	DPC
02286068	FLUOXETINE	SAN
02374447	FLUOXETINE	SIV
02401894	JAMP-FLUOXETINE	JMP
02380560	MINT-FLUOXETINE	MIN
02177579	PMS-FLUOXETINE	PMS
02314991	PRO-FLUOXETINE	PDL
02018985	PROZAC	LIL
02405695	RAN-FLUOXETINE	RBV
02479486	SANDOZ FLUOXETINE	SDZ
02216582	TEVA-FLUOXETINE	TEV
02432412	VAN-FLUOXETINE	VAN

ST **20MG CAPSULE**

02383241	ACH-FLUOXETINE	ACC
02242178	ACT FLUOXETINE	REC
02216361	APO-FLUOXETINE	APX
02385635	AURO-FLUOXETINE	AUR
02448432	BIO-FLUOXETINE	BMI
02177625	DOM-FLUOXETINE	DPC
02286076	FLUOXETINE	SAN
02374455	FLUOXETINE	SIV
02386402	JAMP-FLUOXETINE	JMP
02380579	MINT-FLUOXETINE	MIN
02177587	PMS-FLUOXETINE	PMS
02315009	PRO-FLUOXETINE	PDL
00636622	PROZAC	LIL

28:16.04 ANTIDEPRESSANTS

FLUOXETINE HYDROCHLORIDE

ST 20MG CAPSULE

02405709	RAN-FLUOXETINE	RBY
02305488	RIVA-FLUOXETINE	RIV
02479494	SANDOZ FLUOXETINE	SDZ
02216590	TEVA-FLUOXETINE	TEV
02432420	VAN-FLUOXETINE	VAN

ST 4MG/ML SOLUTION

02231328	APO-FLUOXETINE	APX
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20MG SOLUTION

02459361	ODAN-FLUOXETINE	ODN
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FLUVOXAMINE MALEATE

ST 50MG TABLET

02255529	ACT FLUVOXAMINE	ACG
02231329	APO-FLUVOXAMINE	APX
02236753	FLUVOXAMINE	PDL
01919342	LUVOX	BGP
02303345	RIVA-FLUVOX	RIV
02247054	SANDOZ FLUVOXAMINE	SDZ

ST 100MG TABLET

02255537	ACT FLUVOXAMINE	ACG
02231330	APO-FLUVOXAMINE	APX
02236754	FLUVOXAMINE	PDL
01919369	LUVOX	BGP
02303361	RIVA-FLUVOX	RIV
02247055	SANDOZ FLUVOXAMINE	SDZ

IMIPRAMINE HYDROCHLORIDE

ST 10MG TABLET

00360201	IMIPRAMINE	AAP
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ST 25MG TABLET

00312797	IMIPRAMINE	AAP
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ST 50MG TABLET

00326852	IMIPRAMINE	AAP
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ST 75MG TABLET

00644579	IMIPRAMINE	AAP
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MIRTAZAPINE

ST 15MG TABLET

02286610	APO-MIRTAZAPINE	APX
02411695	AURO-MIRTAZAPINE	AUR
02256096	MYLAN-MIRTAZAPINE	MYL
02273942	PMS-MIRTAZAPINE	PMS
02312778	PRO-MIRTAZAPINE	PDL
02250594	SANDOZ MIRTAZAPINE	SDZ

ST 30MG TABLET

02286629	APO-MIRTAZAPINE	APX
02411709	AURO-MIRTAZAPINE	AUR
02252287	DOM-MIRTAZAPINE	DPC
02370689	MIRTAZAPINE	SAN
02256118	MYLAN-MIRTAZAPINE	MYL
02248762	PMS-MIRTAZAPINE	PMS
02312786	PRO-MIRTAZAPINE	PDL
02243910	REMERON	FRS
02265265	RIVA-MIRTAZAPINE	RIV
02250608	SANDOZ MIRTAZAPINE	SDZ

28:16.04 ANTIDEPRESSANTS

MIRTAZAPINE

ST 30MG TABLET

02259354	TEVA-MIRTAZAPINE	TEV
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ST 45MG TABLET

02286637	APO-MIRTAZAPINE	APX
02411717	AURO-MIRTAZAPINE	AUR
02256126	MYLAN-MIRTAZAPINE	MYL

ST 15MG TABLET (ORALLY DISINTEGRATING)

02299801	AURO-MIRTAZAPINE OD	AUR
02248542	REMERON RD	FRS
02279894	TEVA-MIRTAZAPINE OD	TEV

ST 30MG TABLET (ORALLY DISINTEGRATING)

02299828	AURO-MIRTAZAPINE OD	AUR
02248543	REMERON RD	FRS
02279908	TEVA-MIRTAZAPINE OD	TEV

ST 45MG TABLET (ORALLY DISINTEGRATING)

02299836	AURO-MIRTAZAPINE OD	AUR
02248544	REMERON RD	FRS
02279916	TEVA-MIRTAZAPINE OD	TEV

MOCLOBEMIDE

ST 100MG TABLET

02232148	MOCLOBEMIDE	AAP
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ST 150MG TABLET

00899356	MANERIX	VAE
02232150	MOCLOBEMIDE	AAP
02243218	PMS-MOCLOBEMIDE	PMS

ST 300MG TABLET

02166747	MANERIX	VAE
02240456	MOCLOBEMIDE	AAP
02243219	PMS-MOCLOBEMIDE	PMS

NORTRIPTYLINE HYDROCHLORIDE

ST 10MG CAPSULE

02223511	APO-NORTRIPTYLINE	APX
00015229	AVENTYL	AAP

ST 25MG CAPSULE

02223538	APO-NORTRIPTYLINE	APX
00015237	AVENTYL	AAP

PAROXETINE HYDROCHLORIDE

ST 10MG TABLET

02262746	ACT PAROXETINE	ACG
02475537	AG-PAROXETINE	ANG
02240907	APO-PAROXETINE	APX
02383276	AURO-PAROXETINE	AUR
02444909	BIO-PAROXETINE	BMI
02248447	DOM-PAROXETINE	DPC
02368862	JAMP-PAROXETINE	JMP
02411946	MAR-PAROXETINE	MAR
02421372	MINT-PAROXETINE	MIN
02467402	M-PAROXETINE	MAN
02479753	NRA-PAROXETINE	UNK
02248913	PAROXETINE	PDL
02282844	PAROXETINE	SAN
02388227	PAROXETINE	SIV
02027887	PAXIL	GSK

28:16.04 ANTIDEPRESSANTS

PAROXETINE HYDROCHLORIDE

ST **10MG TABLET**

02247750	PMS-PAROXETINE	PMS
02248559	RIVA-PAROXETINE	RIV
02269422	SANDOZ PAROXETINE	SDZ
02431777	SANDOZ PAROXETINE	SDZ
02248556	TEVA-PAROXETINE	TEV

ST **20MG TABLET**

02262754	ACT PAROXETINE	ACG
02475545	AG-PAROXETINE	ANG
02240908	APO-PAROXETINE	APX
02383284	AURO-PAROXETINE	AUR
02444917	BIO-PAROXETINE	BMI
02248448	DOM-PAROXETINE	DPC
02368870	JAMP-PAROXETINE	JMP
02411954	MAR-PAROXETINE	MAR
02421380	MINT-PAROXETINE	MIN
02467410	M-PAROXETINE	MAN
02479761	NRA-PAROXETINE	UNK
02248914	PAROXETINE	PDL
02282852	PAROXETINE	SAN
02388235	PAROXETINE	SIV
01940481	PAXIL	GSK
02247751	PMS-PAROXETINE	PMS
02248560	RIVA-PAROXETINE	RIV
02269430	SANDOZ PAROXETINE	SDZ
02431785	SANDOZ PAROXETINE	SDZ
02248557	TEVA-PAROXETINE	TEV

ST **30MG TABLET**

02262762	ACT PAROXETINE	ACG
02475553	AG-PAROXETINE	ANG
02240909	APO-PAROXETINE	APX
02383292	AURO-PAROXETINE	AUR
02444925	BIO-PAROXETINE	BMI
02248449	DOM-PAROXETINE	DPC
02368889	JAMP-PAROXETINE	JMP
02411962	MAR-PAROXETINE	MAR
02421399	MINT-PAROXETINE	MIN
02467429	M-PAROXETINE	MAN
02479788	NRA-PAROXETINE	UNK
02248915	PAROXETINE	PDL
02282860	PAROXETINE	SAN
02388243	PAROXETINE	SIV
01940473	PAXIL	GSK
02247752	PMS-PAROXETINE	PMS
02248561	RIVA-PAROXETINE	RIV
02269449	SANDOZ PAROXETINE	SDZ
02431793	SANDOZ PAROXETINE	SDZ
02248558	TEVA-PAROXETINE	TEV

ST **40MG TABLET**

02293749	PMS-PAROXETINE	PMS
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PHENELZINE SULFATE

ST **15MG TABLET**

00476552	NARDIL	ERF
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28:16.04 ANTIDEPRESSANTS

SERTRALINE HYDROCHLORIDE

ST **25MG CAPSULE**

02238280	APO-SERTRALINE	APX
02390906	AURO-SERTRALINE	AUR
02445042	BIO-SERTRALINE	BMI
02245748	DOM-SERTRALINE	DPC
02357143	JAMP-SERTRALINE	JMP
02399415	MAR-SERTRALINE	MAR
02402378	MINT-SERTRALINE	MIN
02244838	PMS-SERTRALINE	PMS
02374552	RAN-SERTRALINE	RBY
02248496	RIVA-SERTRALINE	RIV
02245159	SANDOZ SERTRALINE	SDZ
02353520	SERTRALINE	SAN
02386070	SERTRALINE	SIV
02469626	SERTRALINE	JMP
02241302	SERTRALINE-25	PDL
02240485	TEVA-SERTRALINE	TEV
02132702	ZOLOFT	PFI

ST **50MG CAPSULE**

02238281	APO-SERTRALINE	APX
02390914	AURO-SERTRALINE	AUR
02445050	BIO-SERTRALINE	BMI
02245749	DOM-SERTRALINE	DPC
02357151	JAMP-SERTRALINE	JMP
02399423	MAR-SERTRALINE	MAR
02402394	MINT-SERTRALINE	MIN
02244839	PMS-SERTRALINE	PMS
02374560	RAN-SERTRALINE	RBY
02248497	RIVA-SERTRALINE	RIV
02245160	SANDOZ SERTRALINE	SDZ
02353539	SERTRALINE	SAN
02386089	SERTRALINE	SIV
02469634	SERTRALINE	JMP
02241303	SERTRALINE-50	PDL
02240484	TEVA-SERTRALINE	TEV
01962817	ZOLOFT	PFI

ST **100MG CAPSULE**

02238282	APO-SERTRALINE	APX
02390922	AURO-SERTRALINE	AUR
02445069	BIO-SERTRALINE	BMI
02245750	DOM-SERTRALINE	DPC
02357178	JAMP-SERTRALINE	JMP
02399431	MAR-SERTRALINE	MAR
02402408	MINT-SERTRALINE	MIN
02244840	PMS-SERTRALINE	PMS
02374579	RAN-SERTRALINE	RBY
02248498	RIVA-SERTRALINE	RIV
02245161	SANDOZ SERTRALINE	SDZ
02353547	SERTRALINE	SAN
02386097	SERTRALINE	SIV
02469642	SERTRALINE	JMP
02241304	SERTRALINE-100	PDL
02240481	TEVA-SERTRALINE	TEV
01962779	ZOLOFT	PFI

28:16.04 ANTIDEPRESSANTS

TRANLYCYPROMINE SULFATE

ST **10MG TABLET**

01919598 PARNATE GSK

TRAZODONE HYDROCHLORIDE

ST **50MG TABLET**

02147637 APO-TRAZODONE APX
 02128950 DOM-TRAZODONE DPC
 01937227 PMS TRAZODONE PMS
 02144263 TEVA-TRAZODONE TEV
 02164353 TRAZODONE PDL
 02348772 TRAZODONE SAN

ST **75MG TABLET**

02237339 PMS-TRAZODONE PMS

ST **100MG TABLET**

02147645 APO-TRAZODONE APX
 02128969 DOM-TRAZODONE DPC
 01937235 PMS TRAZODONE PMS
 02144271 TEVA-TRAZODONE TEV
 02164361 TRAZODONE PDL
 02348780 TRAZODONE SAN

ST **150MG TABLET**

02147653 APO-TRAZODONE D APX
 02144298 TEVA-TRAZODONE TEV
 02164388 TRAZODONE PDL
 02348799 TRAZODONE SAN

TRIMIPRAMINE MALEATE

ST **75MG CAPSULE**

02070987 TRIMIPRAMINE AAP

ST **12.5MG TABLET**

00740799 TRIMIPRAMINE AAP

ST **25MG TABLET**

00740802 TRIMIPRAMINE AAP

ST **50MG TABLET**

00740810 TRIMIPRAMINE AAP

ST **100MG TABLET**

00740829 TRIMIPRAMINE AAP

VENLAFAXINE HYDROCHLORIDE

ST **37.5MG CAPSULE (EXTENDED RELEASE)**

02304317 ACT VENLAFAXINE XR TEV
 02331683 APO-VENLAFAXINE XR APX
 02452839 AURO-VENLAFAXINE XR AUR
 02299291 DOM-VENLAFAXINE XR DPC
 02237279 EFFEXOR XR PFI
 02471280 M-VENLAFAXINE XR MAN
 02278545 PMS-VENLAFAXINE XR PMS
 02380072 RAN-VENLAFAXINE XR RBY
 02307774 RIVA-VENLAFAXINE XR RIV
 02310317 SANDOZ VENLAFAXINE XR SDZ
 02275023 TEVA-VENLAFAXINE XR TEV
 02339242 VENLAFAXINE XR PDL
 02354713 VENLAFAXINE XR SAN
 02385929 VENLAFAXINE XR SIV

ST **75MG CAPSULE (EXTENDED RELEASE)**

02304325 ACT VENLAFAXINE XR TEV

28:16.04 ANTIDEPRESSANTS

VENLAFAXINE HYDROCHLORIDE

ST **75MG CAPSULE (EXTENDED RELEASE)**

02331691 APO-VENLAFAXINE XR APX
 02452847 AURO-VENLAFAXINE XR AUR
 02299305 DOM-VENLAFAXINE XR DPC
 02237280 EFFEXOR XR PFI
 02471299 M-VENLAFAXINE XR MAN
 02278553 PMS-VENLAFAXINE XR PMS
 02380080 RAN-VENLAFAXINE XR RBY
 02307782 RIVA-VENLAFAXINE XR RIV
 02310325 SANDOZ VENLAFAXINE XR SDZ
 02275031 TEVA-VENLAFAXINE XR TEV
 02339250 VENLAFAXINE XR PDL
 02354721 VENLAFAXINE XR SAN
 02385937 VENLAFAXINE XR SIV

ST **150MG CAPSULE (EXTENDED RELEASE)**

02304333 ACT VENLAFAXINE XR TEV
 02331705 APO-VENLAFAXINE XR APX
 02452855 AURO-VENLAFAXINE XR AUR
 02299313 DOM-VENLAFAXINE XR DPC
 02237282 EFFEXOR XR PFI
 02471302 M-VENLAFAXINE XR MAN
 02278561 PMS-VENLAFAXINE XR PMS
 02380099 RAN-VENLAFAXINE XR RBY
 02307790 RIVA-VENLAFAXINE XR RIV
 02310333 SANDOZ VENLAFAXINE XR SDZ
 02275058 TEVA-VENLAFAXINE XR TEV
 02339269 VENLAFAXINE XR PDL
 02354748 VENLAFAXINE XR SAN
 02385945 VENLAFAXINE XR SIV

28:16.08 ANTIPSYCHOTIC AGENTS

ARIPIPRAZOLE

Limited use benefit (prior approval required).

For the treatment of schizophrenia and schizoaffective disorders in patients:

- who have intolerance or lack of response to an adequate trial of another antipsychotic agent; OR
- a contraindication to another antipsychotic agent.

ST **2MG TABLET**

02322374 ABILIFY OTS
 02471086 APO-ARIPIPRAZOLE APX
 02460025 AURO-ARIPIPRAZOLE PMS
 02466635 PMS-ARIPIPRAZOLE PMS
 02479346 RIVA-ARIPIPRAZOLE RIV
 02473658 SANDOZ ARIPIPRAZOLE SDZ
 02464144 TEVA-ARIPIPRAZOLE TEV

ST **5MG TABLET**

02322382 ABILIFY OTS
 02471094 APO-ARIPIPRAZOLE APX
 02460033 AURO-ARIPIPRAZOLE PMS
 02466643 PMS-ARIPIPRAZOLE PMS
 02479354 RIVA-ARIPIPRAZOLE RIV
 02473666 SANDOZ ARIPIPRAZOLE SDZ
 02464152 TEVA-ARIPIPRAZOLE TEV

28:16.08 ANTIPSYCHOTIC AGENTS

ARIPIPIRAZOLE

Limited use benefit (prior approval required).

For the treatment of schizophrenia and schizoaffective disorders in patients:
 • who have intolerance or lack of response to an adequate trial of another antipsychotic agent; OR
 • a contraindication to another antipsychotic agent.

ST **10MG TABLET**

02322390	ABILIFY	OTS
02471108	APO-ARIPIPIRAZOLE	APX
02460041	AURO-ARIPIPIRAZOLE	PMS
02466651	PMS-ARIPIPIRAZOLE	PMS
02479362	RIVA-ARIPIPIRAZOLE	RIV
02473674	SANDOZ ARIPIPIRAZOLE	SDZ
02464160	TEVA-ARIPIPIRAZOLE	TEV

ST **15MG TABLET**

02322404	ABILIFY	OTS
02471116	APO-ARIPIPIRAZOLE	APX
02460068	AURO-ARIPIPIRAZOLE	PMS
02466678	PMS-ARIPIPIRAZOLE	PMS
02479370	RIVA-ARIPIPIRAZOLE	RIV
02473682	SANDOZ ARIPIPIRAZOLE	SDZ
02464179	TEVA-ARIPIPIRAZOLE	TEV

ST **20MG TABLET**

02322412	ABILIFY	OTS
02471124	APO-ARIPIPIRAZOLE	APX
02460076	AURO-ARIPIPIRAZOLE	PMS
02466686	PMS-ARIPIPIRAZOLE	PMS
02479389	RIVA-ARIPIPIRAZOLE	RIV
02473690	SANDOZ ARIPIPIRAZOLE	SDZ
02464187	TEVA-ARIPIPIRAZOLE	TEV

ST **30MG TABLET**

02322455	ABILIFY	OTS
02471132	APO-ARIPIPIRAZOLE	APX
02460084	AURO-ARIPIPIRAZOLE	PMS
02466694	PMS-ARIPIPIRAZOLE	PMS
02479397	RIVA-ARIPIPIRAZOLE	RIV
02473704	SANDOZ ARIPIPIRAZOLE	SDZ
02464195	TEVA-ARIPIPIRAZOLE	TEV

ARIPIPIRAZOLE (MAINTENA)

300MG INJECTION

02420864	ABILIFY MAINTENA	OTS
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400MG INJECTION

02420872	ABILIFY MAINTENA	OTS
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28:16.08 ANTIPSYCHOTIC AGENTS

ASENAPINE MALEATE

Limited use benefit (prior approval required).

For the acute treatment of manic or mixed episodes associated with bipolar I disorder as either:
 • Monotherapy, after a trial of lithium or divalproex sodium has failed or is contraindicated, and trials of two atypical antipsychotic agents have failed due to intolerance or lack of response; OR
 • Co-therapy with lithium or divalproex sodium, after trials of two atypical antipsychotic agents have failed due to intolerance or lack of response.

ST **5MG TABLET**

02374803	SAPHRIS	FRS
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ST **10MG TABLET**

02374811	SAPHRIS	FRS
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BREXPIPIRAZOLE

0.25MG TABLET

02461749	REXULTI	OTS
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0.5MG TABLET

02461757	REXULTI	OTS
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1MG TABLET

02461765	REXULTI	OTS
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2MG TABLET

02461773	REXULTI	OTS
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3MG TABLET

02461781	REXULTI	OTS
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4MG TABLET

02461803	REXULTI	OTS
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CHLORPROMAZINE HYDROCHLORIDE

25MG/ML SOLUTION

00743518	CHLORPROMAZINE	SDZ
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ST **25MG TABLET**

00232823	TEVA-CHLORPROMAZINE	TEV
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ST **50MG TABLET**

00232807	TEVA-CHLORPROMAZINE	TEV
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ST **100MG TABLET**

00232831	TEVA-CHLORPROMAZINE	TEV
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CLOZAPINE

ST **25MG TABLET**

02248034	AA-CLOZAPINE	AAP
00894737	CLOZARIL	HLS
02247243	GEN-CLOZAPINE	MYL

ST **50MG TABLET**

02458748	AA-CLOZAPINE	AAP
02305003	GEN-CLOZAPINE	MYL

ST **100MG TABLET**

02248035	AA-CLOZAPINE	AAP
00894745	CLOZARIL	HLS
02247244	GEN-CLOZAPINE	MYL

ST **200MG TABLET**

02458756	AA-CLOZAPINE	AAP
02305011	GEN-CLOZAPINE	MYL

FLUPENTHIXOL DIHYDROCHLORIDE

ST **0.5MG TABLET**

02156008	FLUANXOL	LUD
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28:16.08 ANTIPSYCHOTIC AGENTS

FLUPENTHIXOL DIHYDROCHLORIDE

ST **3MG TABLET**
02156016 FLUANXOL LUD

FLUPENTIXOL DECANOATE

20MG/ML SOLUTION
02156032 FLUANXOL DEPOT LUD

100MG/ML SOLUTION
02156040 FLUANXOL DEPOT LUD

FLUPHENAZINE DECANOATE

25MG/ML LIQUID
02091275 PMS-FLUPHENAZINE PMS

100MG/ML LIQUID
02241928 PMS-FLUPHENAZINE PMS

FLUPHENAZINE HYDROCHLORIDE

ST **1MG TABLET**
00405345 FLUPHENAZINE AAP

ST **2MG TABLET**
00410632 FLUPHENAZINE AAP

ST **5MG TABLET**
00405361 FLUPHENAZINE AAP
00726354 PMS FLUPHENAZINE PMS

HALOPERIDOL

ST **2MG/ML SOLUTION**
00759503 PMS-HALOPERIDOL PMS

5MG/ML SOLUTION
00808652 HALOPERIDOL SDZ
02366010 HALOPERIDOL OMG

ST **0.5MG TABLET**
00396796 APO HALOPERIDOL APX
00363685 TEVA-HALOPERIDOL TEV

ST **1MG TABLET**
00396818 APO HALOPERIDOL APX
00363677 TEVA-HALOPERIDOL TEV

ST **2MG TABLET**
00363669 TEVA-HALOPERIDOL TEV

ST **5MG TABLET**
00363650 TEVA-HALOPERIDOL TEV

ST **10MG TABLET**
00463698 APO-HALOPERIDOL APX
00713449 TEVA-HALOPERIDOL TEV

ST **20MG TABLET**
00768820 TEVA-HALOPERIDOL TEV

HALOPERIDOL DECANOATE

50MG/ML LIQUID
02130297 HALOPERIDOL LA SDZ
02230707 PMS-HALOPERIDOL PMS

100MG/ML LIQUID
02130300 HALOPERIDOL LA SDZ
02239640 HALOPERIDOL LA OMG
02230708 PMS-HALOPERIDOL PMS

LOXAPINE HYDROCHLORIDE

ST **25MG/ML SOLUTION**
02239101 XYLAC PED

28:16.08 ANTIPSYCHOTIC AGENTS

LOXAPINE SUCCINATE

ST **2.5MG TABLET**
02242868 XYLAC PED

ST **5MG TABLET**
02239918 DOM-LOXAPINE DPC
02230837 XYLAC PED

ST **10MG TABLET**
02239919 DOM-LOXAPINE DPC
02230838 XYLAC PED

ST **25MG TABLET**
02239920 DOM-LOXAPINE DPC
02230839 XYLAC PED

ST **50MG TABLET**
02239921 DOM-LOXAPINE DPC
02230840 XYLAC PED

LURASIDONE HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of schizophrenia and schizoaffective disorders in patients:

- who have intolerance or lack of response to an adequate trial of another antipsychotic agent; OR
- a contraindication to another antipsychotic agent.

ST **20MG TABLET**
02422050 LATUDA SPC

ST **40MG TABLET**
02387751 LATUDA SPC

ST **60MG TABLET**
02413361 LATUDA SPC

ST **80MG TABLET**
02387778 LATUDA SPC

ST **120MG TABLET**
02387786 LATUDA SPC

METHOTRIMEPRAZINE MALEATE

ST **2MG TABLET**
02238403 METHOPRAZINE AAP

ST **5MG TABLET**
02238404 METHOPRAZINE AAP

ST **25MG TABLET**
02238405 METHOPRAZINE AAP

ST **50MG TABLET**
02238406 METHOPRAZINE AAP

OLANZAPINE

ST **2.5MG TABLET**
02281791 APO-OLANZAPINE APX
02417243 JAMP-OLANZAPINE JMP

02311968 OLANZAPINE PDL
02372819 OLANZAPINE SAN

02385864 OLANZAPINE SIV
02303116 PMS-OLANZAPINE PMS

02403064 RAN-OLANZAPINE RBY
02337126 RIVA-OLANZAPINE RIV

02310341 SANDOZ OLANZAPINE SDZ
02276712 TEVA-OLANZAPINE TEV

02229250 ZYPREXA LIL

28:16.08 ANTIPSYCHOTIC AGENTS

OLANZAPINE

ST 5MG TABLET

02281805	APO-OLANZAPINE	APX
02417251	JAMP-OLANZAPINE	JMP
02311976	OLANZAPINE	PDL
02372827	OLANZAPINE	SAN
02385872	OLANZAPINE	SIV
02303159	PMS-OLANZAPINE	PMS
02403072	RAN-OLANZAPINE	RBY
02337134	RIVA-OLANZAPINE	RIV
02310368	SANDOZ OLANZAPINE	SDZ
02276720	TEVA-OLANZAPINE	TEV
02229269	ZYPREXA	LIL

ST 7.5MG TABLET

02281813	APO-OLANZAPINE	APX
02417278	JAMP-OLANZAPINE	JMP
02311984	OLANZAPINE	PDL
02372835	OLANZAPINE	SAN
02385880	OLANZAPINE	SIV
02303167	PMS-OLANZAPINE	PMS
02403080	RAN-OLANZAPINE	RBY
02337142	RIVA-OLANZAPINE	RIV
02310376	SANDOZ OLANZAPINE	SDZ
02276739	TEVA-OLANZAPINE	TEV
02229277	ZYPREXA	LIL

ST 10MG TABLET

02281821	APO-OLANZAPINE	APX
02417286	JAMP-OLANZAPINE	JMP
02311992	OLANZAPINE	PDL
02372843	OLANZAPINE	SAN
02385899	OLANZAPINE	SIV
02303175	PMS-OLANZAPINE	PMS
02403099	RAN-OLANZAPINE	RBY
02337150	RIVA-OLANZAPINE	RIV
02310384	SANDOZ OLANZAPINE	SDZ
02276747	TEVA-OLANZAPINE	TEV
02229285	ZYPREXA	LIL

ST 15MG TABLET

02281848	APO-OLANZAPINE	APX
02417294	JAMP-OLANZAPINE	JMP
02312018	OLANZAPINE	PDL
02372851	OLANZAPINE	SAN
02385902	OLANZAPINE	SIV
02303183	PMS-OLANZAPINE	PMS
02403102	RAN-OLANZAPINE	RBY
02337169	RIVA-OLANZAPINE	RIV
02310392	SANDOZ OLANZAPINE	SDZ
02276755	TEVA-OLANZAPINE	TEV
02238850	ZYPREXA	LIL

ST 20MG TABLET

02417308	JAMP-OLANZAPINE	JMP
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ST 5MG TABLET (ORALLY DISINTEGRATING)

02327562	ACT OLANZAPINE ODT	TEV
02360616	APO-OLANZAPINE ODT	APX
02448726	AURO-OLANZAPINE ODT	AUR
02406624	JAMP OLANZAPINE ODT	JMP

28:16.08 ANTIPSYCHOTIC AGENTS

OLANZAPINE

ST 5MG TABLET (ORALLY DISINTEGRATING)

02389088	MAR-OLANZAPINE ODT	MAR
02436965	MINT-OLANZAPINE ODT	MIN
02338645	OLANZAPINE ODT	PDL
02343665	OLANZAPINE ODT	SIV
02352974	OLANZAPINE ODT	SAN
02303191	PMS-OLANZAPINE ODT	PMS
02414090	RAN-OLANZAPINE ODT	RBY
02327775	SANDOZ OLANZAPINE ODT	SDZ
02243086	ZYPREXA ZYDIS	LIL

ST 10MG TABLET (ORALLY DISINTEGRATING)

02327570	ACT OLANZAPINE ODT	TEV
02360624	APO-OLANZAPINE ODT	APX
02448734	AURO-OLANZAPINE ODT	AUR
02406632	JAMP OLANZAPINE ODT	JMP
02389096	MAR-OLANZAPINE ODT	MAR
02436973	MINT-OLANZAPINE ODT	MIN
02338653	OLANZAPINE ODT	PDL
02343673	OLANZAPINE ODT	SIV
02352982	OLANZAPINE ODT	SAN
02303205	PMS-OLANZAPINE ODT	PMS
02414104	RAN-OLANZAPINE ODT	RBY
02327783	SANDOZ OLANZAPINE ODT	SDZ
02243087	ZYPREXA ZYDIS	LIL

ST 15MG TABLET (ORALLY DISINTEGRATING)

02327589	ACT OLANZAPINE ODT	TEV
02360632	APO-OLANZAPINE ODT	APX
02448742	AURO-OLANZAPINE ODT	AUR
02406640	JAMP OLANZAPINE ODT	JMP
02389118	MAR-OLANZAPINE ODT	MAR
02436981	MINT-OLANZAPINE ODT	MIN
02338661	OLANZAPINE ODT	PDL
02343681	OLANZAPINE ODT	SIV
02352990	OLANZAPINE ODT	SAN
02303213	PMS-OLANZAPINE ODT	PMS
02414112	RAN-OLANZAPINE ODT	RBY
02327791	SANDOZ OLANZAPINE ODT	SDZ
02243088	ZYPREXA ZYDIS	LIL

PALIPERIDONE PALMITATE

50MG/0.5ML SUSPENSION (EXTENDED RELEASE)

02354217	INVEGA SUSTENNA	JSO
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75MG/0.75ML SUSPENSION (EXTENDED RELEASE)

02354225	INVEGA SUSTENNA	JSO
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100MG/ML SUSPENSION (EXTENDED RELEASE)

02354233	INVEGA SUSTENNA	JSO
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150MG/1.5ML SUSPENSION (EXTENDED RELEASE)

02354241	INVEGA SUSTENNA	JSO
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175MG SUSPENSION (EXTENDED RELEASE)

02455943	INVEGA TRINZA	JSO
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263MG SUSPENSION (EXTENDED RELEASE)

02455986	INVEGA TRINZA	JSO
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350MG SUSPENSION (EXTENDED RELEASE)

02455994	INVEGA TRINZA	JSO
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28:16.08 ANTIPSYCHOTIC AGENTS

PALIPERIDONE PALMITATE

525MG SUSPENSION (EXTENDED RELEASE)

02456001 INVEGA TRINZA JSO

PERICYAZINE

ST **5MG CAPSULE**

01926780 NEULEPTIL ERF

ST **10MG CAPSULE**

01926772 NEULEPTIL ERF

ST **20MG CAPSULE**

01926764 NEULEPTIL ERF

ST **10MG/ML DROP**

01926756 NEULEPTIL ERF

PERPHENAZINE

ST **3.2MG/ML LIQUID**

00751898 PMS PERPHENAZINE PMS

ST **2MG TABLET**

00335134 PERPHENAZINE AAP

ST **4MG TABLET**

00335126 PERPHENAZINE AAP

ST **8MG TABLET**

00335118 PERPHENAZINE AAP

ST **16MG TABLET**

00335096 PERPHENAZINE AAP

00726206 PMS PERPHENAZINE PMS

PIMOZIDE

ST **2MG TABLET**

00313815 ORAP AAP

02245432 PIMOZIDE AAP

ST **4MG TABLET**

00313823 ORAP AAP

02245433 PIMOZIDE AAP

PIPOTIAZINE PALMITATE

50MG/ML INJECTION

00894672 PIPORTIL L4 SAC

PROCHLORPERAZINE

10MG SUPPOSITORY

00753688 PMS-PROCHLORPERAZINE PMS

00789720 SANDOZ PROCHLORPERAZINE SDZ

PROCHLORPERAZINE MALEATE

ST **5MG TABLET**

00753661 PMS-PROCHLORPERAZINE PMS

00886440 PROCHLORAZINE AAP

ST **10MG TABLET**

00753637 PMS-PROCHLORPERAZINE PMS

00886432 PROCHLORAZINE AAP

PROCHLORPERAZINE MESYLATE

5MG/ML SOLUTION

00753645 PMS PROCHLORPERAZINE PMS

QUETIAPINE FUMARATE

ST **25MG TABLET**

02316080 ACT QUETIAPINE ACG

02313901 APO-QUETIAPINE APX

28:16.08 ANTIPSYCHOTIC AGENTS

QUETIAPINE FUMARATE

ST **25MG TABLET**

02390205 AURO-QUETIAPINE AUR

02447193 BIO-QUETIAPINE BMI

02298996 DOM-QUETIAPINE DPC

02330415 JAMP-QUETIAPINE JMP

02438003 MINT-QUETIAPINE MIN

02439158 NAT-QUETIAPINE NPH

02296551 PMS-QUETIAPINE PMS

02317346 PRO-QUETIAPINE PDL

02317893 QUETIAPINE SIV

02353164 QUETIAPINE SAN

02387794 QUETIAPINE ACC

02397099 RAN-QUETIAPINE RBY

02316692 RIVA-QUETIAPINE RIV

02313995 SANDOZ QUETIAPINE SDZ

02236951 SEROQUEL AZC

02284235 TEVA-QUETIAPINE TEV

02434024 VAN-QUETIAPINE VAN

ST **50MG TABLET**

02361892 PMS-QUETIAPINE PMS

ST **100MG TABLET**

02316099 ACT QUETIAPINE ACG

02313928 APO-QUETIAPINE APX

02390213 AURO-QUETIAPINE AUR

02447207 BIO-QUETIAPINE BMI

02299003 DOM-QUETIAPINE DPC

02330423 JAMP-QUETIAPINE JMP

02438011 MINT-QUETIAPINE MIN

02439166 NAT-QUETIAPINE NPH

02296578 PMS-QUETIAPINE PMS

02317354 PRO-QUETIAPINE PDL

02317907 QUETIAPINE SIV

02353172 QUETIAPINE SAN

02387808 QUETIAPINE ACC

02397102 RAN-QUETIAPINE RBY

02316706 RIVA-QUETIAPINE RIV

02314002 SANDOZ QUETIAPINE SDZ

02236952 SEROQUEL AZC

02284243 TEVA-QUETIAPINE TEV

02434032 VAN-QUETIAPINE VAN

ST **200MG TABLET**

02316110 ACT QUETIAPINE ACG

02313936 APO-QUETIAPINE APX

02390248 AURO-QUETIAPINE AUR

02447223 BIO-QUETIAPINE BMI

02299038 DOM-QUETIAPINE DPC

02330458 JAMP-QUETIAPINE JMP

02438046 MINT-QUETIAPINE MIN

02439182 NAT-QUETIAPINE NPH

02296594 PMS-QUETIAPINE PMS

02317362 PRO-QUETIAPINE PDL

02317923 QUETIAPINE SIV

02353199 QUETIAPINE SAN

02387824 QUETIAPINE ACC

02397110 RAN-QUETIAPINE RBY

28:16.08 ANTIPSYCHOTIC AGENTS

QUETIAPINE FUMARATE

ST 200MG TABLET

02316722	RIVA-QUETIAPINE	RIV
02314010	SANDOZ QUETIAPINE	SDZ
02236953	SEROQUEL	AZC
02284278	TEVA-QUETIAPINE	TEV
02434040	VAN-QUETIAPINE	VAN

ST 300MG TABLET

02316129	ACT QUETIAPINE	ACG
02313944	APO-QUETIAPINE	APX
02390256	AURO-QUETIAPINE	AUR
02447258	BIO-QUETIAPINE	BMI
02299046	DOM-QUETIAPINE	DPC
02330466	JAMP-QUETIAPINE	JMP
02438054	MINT-QUETIAPINE	MIN
02439190	NAT-QUETIAPINE	NPH
02296608	PMS-QUETIAPINE	PMS
02317370	PRO-QUETIAPINE	PDL
02317931	QUETIAPINE	SIV
02353202	QUETIAPINE	SAN
02387832	QUETIAPINE	ACC
02397129	RAN-QUETIAPINE	RBY
02316730	RIVA-QUETIAPINE	RIV
02314029	SANDOZ QUETIAPINE	SDZ
02244107	SEROQUEL	AZC
02284286	TEVA-QUETIAPINE	TEV
02434059	VAN-QUETIAPINE	VAN

50MG TABLET (EXTENDED RELEASE)

02457229	APO-QUETIAPINE XR	APX
02417359	QUETIAPINE XR	SIV
02417782	QUETIAPINE XR	PDL
02407671	SANDOZ QUETIAPINE XRT	SDZ
02300184	SEROQUEL XR	AZC
02395444	TEVA-QUETIAPINE XR	TEV

ST 150MG TABLET (EXTENDED RELEASE)

02457237	APO-QUETIAPINE XR	APX
02417367	QUETIAPINE XR	SIV
02417790	QUETIAPINE XR	PDL
02407698	SANDOZ QUETIAPINE XRT	SDZ
02321513	SEROQUEL XR	AZC
02395452	TEVA-QUETIAPINE XR	TEV

ST 200MG TABLET (EXTENDED RELEASE)

02457245	APO-QUETIAPINE XR	APX
02417375	QUETIAPINE XR	SIV
02417804	QUETIAPINE XR	PDL
02407701	SANDOZ QUETIAPINE XRT	SDZ
02300192	SEROQUEL XR	AZC
02395460	TEVA-QUETIAPINE XR	TEV

300MG TABLET (EXTENDED RELEASE)

02457253	APO-QUETIAPINE XR	APX
02417383	QUETIAPINE XR	SIV
02417812	QUETIAPINE XR	PDL
02407728	SANDOZ QUETIAPINE XRT	SDZ
02300206	SEROQUEL XR	AZC
02395479	TEVA-QUETIAPINE XR	TEV

28:16.08 ANTIPSYCHOTIC AGENTS

QUETIAPINE FUMARATE

400MG TABLET (EXTENDED RELEASE)

02457261	APO-QUETIAPINE XR	APX
02417391	QUETIAPINE XR	SIV
02417820	QUETIAPINE XR	PDL
02407736	SANDOZ QUETIAPINE XRT	SDZ
02300214	SEROQUEL XR	AZC
02395487	TEVA-QUETIAPINE XR	TEV

ST 25MG TABLET (IMMEDIATE RELEASE)

02399822	MAR-QUETIAPINE	MAR
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ST 100MG TABLET (IMMEDIATE RELEASE)

02399830	MAR-QUETIAPINE	MAR
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ST 200MG TABLET (IMMEDIATE RELEASE)

02399849	MAR-QUETIAPINE	MAR
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ST 300MG TABLET (IMMEDIATE RELEASE)

02399857	MAR-QUETIAPINE	MAR
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RISPERIDONE

ST 1MG SOLUTION

02454319	JAMP-RISPERIDONE	JMP
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ST 1MG/ML SOLUTION

02280396	APO-RISPERIDONE	APX
02279266	PMS-RISPERIDONE	PMS
02236950	RISPERDAL	JSO

ST 0.25MG TABLET

02282119	APO-RISPERIDONE	APX
02359529	JAMP-RISPERIDONE	JMP
02371766	MAR-RISPERIDONE	MAR
02359790	MINT-RISPERIDON	MIN
02252007	PMS-RISPERIDONE	PMS
02312700	PRO-RISPERIDONE	PDL
02328305	RAN-RISPERIDONE	RBY
02356880	RISPERIDONE	SAN
02283565	RIVA-RISPERIDONE	RIV
02303655	SANDOZ RISPERIDONE	SDZ
02282690	TEVA-RISPERIDONE	TEV

ST 0.5MG TABLET

02282127	APO-RISPERIDONE	APX
02359537	JAMP-RISPERIDONE	JMP
02371774	MAR-RISPERIDONE	MAR
02359804	MINT-RISPERIDON	MIN
02252015	PMS-RISPERIDONE	PMS
02312719	PRO-RISPERIDONE	PDL
02328313	RAN-RISPERIDONE	RBY
02356899	RISPERIDONE	SAN
02283573	RIVA-RISPERIDONE	RIV
02303663	SANDOZ RISPERIDONE	SDZ
02264188	TEVA-RISPERIDONE	TEV

ST 1MG TABLET

02282135	APO-RISPERIDONE	APX
02359545	JAMP-RISPERIDONE	JMP
02371782	MAR-RISPERIDONE	MAR
02359812	MINT-RISPERIDON	MIN
02252023	PMS-RISPERIDONE	PMS
02312727	PRO-RISPERIDONE	PDL
02328321	RAN-RISPERIDONE	RBY

28:16.08 ANTIPSYCHOTIC AGENTS

RISPERIDONE

ST 1MG TABLET		
02356902	RISPERIDONE	SAN
02283581	RIVA-RISPERIDONE	RIV
02279800	SANDOZ RISPERIDONE	SDZ
02264196	TEVA-RISPERIDONE	TEV
ST 2MG TABLET		
02282143	APO-RISPERIDONE	APX
02359553	JAMP-RISPERIDONE	JMP
02371790	MAR-RISPERIDONE	MAR
02359820	MINT-RISPERIDONE	MIN
02252031	PMS-RISPERIDONE	PMS
02312735	PRO-RISPERIDONE	PDL
02328348	RAN-RISPERIDONE	RBV
02356910	RISPERIDONE	SAN
02283603	RIVA-RISPERIDONE	RIV
02279819	SANDOZ RISPERIDONE	SDZ
02264218	TEVA-RISPERIDONE	TEV
ST 3MG TABLET		
02282151	APO-RISPERIDONE	APX
02359561	JAMP-RISPERIDONE	JMP
02371804	MAR-RISPERIDONE	MAR
02359839	MINT-RISPERIDONE	MIN
02252058	PMS-RISPERIDONE	PMS
02312743	PRO-RISPERIDONE	PDL
02328364	RAN-RISPERIDONE	RBV
02356929	RISPERIDONE	SAN
02283611	RIVA-RISPERIDONE	RIV
02279827	SANDOZ RISPERIDONE	SDZ
02264226	TEVA-RISPERIDONE	TEV
ST 4MG TABLET		
02282178	APO-RISPERIDONE	APX
02359588	JAMP-RISPERIDONE	JMP
02371812	MAR-RISPERIDONE	MAR
02359847	MINT-RISPERIDONE	MIN
02252066	PMS-RISPERIDONE	PMS
02312751	PRO-RISPERIDONE	PDL
02328372	RAN-RISPERIDONE	RBV
02356937	RISPERIDONE	SAN
02283638	RIVA-RISPERIDONE	RIV
02279835	SANDOZ RISPERIDONE	SDZ
02264234	TEVA-RISPERIDONE	TEV
ST 0.5MG TABLET (ORALLY DISINTEGRATING)		
02413485	MYLAN-RISPERIDONE ODT	MYL
ST 1MG TABLET (ORALLY DISINTEGRATING)		
02413493	MYLAN-RISPERIDONE ODT	MYL
ST 2MG TABLET (ORALLY DISINTEGRATING)		
02413507	MYLAN-RISPERIDONE ODT	MYL
ST 3MG TABLET (ORALLY DISINTEGRATING)		
02413515	MYLAN-RISPERIDONE ODT	MYL
ST 4MG TABLET (ORALLY DISINTEGRATING)		
02413523	MYLAN-RISPERIDONE ODT	MYL

RISPERIDONE (CONSTA)

12.5MG INJECTION		
02298465	RISPERDAL CONSTA	JSO

28:16.08 ANTIPSYCHOTIC AGENTS

RISPERIDONE (CONSTA)

25MG INJECTION		
02255707	RISPERDAL CONSTA	JSO
ST 37.5MG INJECTION		
02255723	RISPERDAL CONSTA	JSO
ST 50MG INJECTION		
02255758	RISPERDAL CONSTA	JSO
THIOPROPERAZINE MESYLATE		
ST 10MG TABLET		
01927639	MAJEPTIL	ERF
THIOTHIXENE		
ST 5MG CAPSULE		
00024449	NAVANE	ERF
TRIFLUOPERAZINE HYDROCHLORIDE		
ST 1MG TABLET		
00345539	TRIFLUOPERAZINE	AAP
ST 2MG TABLET		
00312754	TRIFLUOPERAZINE	AAP
ST 5MG TABLET		
00312746	TRIFLUOPERAZINE	AAP
ST 10MG TABLET		
00326836	TRIFLUOPERAZINE	AAP
ST 20MG TABLET		
00595942	TRIFLUOPERAZINE	AAP
ZIPRASIDONE HYDROCHLORIDE MONOHYDRATE		
ST 20MG CAPSULE		
02449544	AURO-ZIPRASIDONE	AUR
02298597	ZELDOX	PFI
ST 40MG CAPSULE		
02449552	AURO-ZIPRASIDONE	AUR
02298600	ZELDOX	PFI
ST 60MG CAPSULE		
02449560	AURO-ZIPRASIDONE	AUR
02298619	ZELDOX	PFI
ST 80MG CAPSULE		
02449579	AURO-ZIPRASIDONE	AUR
02298627	ZELDOX	PFI
ZUCLOPENTHIXOL ACETATE		
50MG/ML SOLUTION		
02230405	CLOPIXOL-ACUPHASE	LUD
ZUCLOPENTHIXOL DIHYDROCHLORIDE		
200MG/ML SOLUTION		
02230406	CLOPIXOL DEPOT	LUD
ST 10MG TABLET		
02230402	CLOPIXOL	LUD
ST 25MG TABLET		
02230403	CLOPIXOL	LUD

28:20.04 AMPHETAMINES

AMPHETAMINE, DEXTROAMPHETAMINE

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST 5MG CAPSULE (EXTENDED RELEASE)

02439239	ACT AMPHETAMINE XR	TEV
02248808	ADDERALL XR	UNK
02440369	PMS-AMPHETAMINES XR	PMS
02457288	SANDOZ AMPHETAMINE XR	SDZ

ST 10MG CAPSULE (EXTENDED RELEASE)

02439247	ACT AMPHETAMINE XR	TEV
02248809	ADDERALL XR	UNK
02440377	PMS-AMPHETAMINES XR	PMS
02457296	SANDOZ AMPHETAMINE XR	SDZ

ST 15MG CAPSULE (EXTENDED RELEASE)

02439255	ACT AMPHETAMINE XR	TEV
02248810	ADDERALL XR	UNK
02440385	PMS-AMPHETAMINES XR	PMS
02457318	SANDOZ AMPHETAMINE XR	SDZ

ST 20MG CAPSULE (EXTENDED RELEASE)

02439263	ACT AMPHETAMINE XR	TEV
02248811	ADDERALL XR	UNK
02440393	PMS-AMPHETAMINES XR	PMS
02457326	SANDOZ AMPHETAMINE XR	SDZ

ST 25MG CAPSULE (EXTENDED RELEASE)

02439271	ACT AMPHETAMINE XR	TEV
02248812	ADDERALL XR	UNK
02440407	PMS-AMPHETAMINES XR	PMS
02457334	SANDOZ AMPHETAMINE XR	SDZ

ST 30MG CAPSULE (EXTENDED RELEASE)

02439298	ACT AMPHETAMINE XR	TEV
02248813	ADDERALL XR	UNK
02440415	PMS-AMPHETAMINES XR	PMS
02457342	SANDOZ AMPHETAMINE XR	SDZ

28:20.04 AMPHETAMINES

DEXTROAMPHETAMINE SULFATE

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST 10MG CAPSULE (SUSTAINED RELEASE)

02448319	ACT DEXTROAMPHETAMINE SR	ACG
01924559	DEXEDRINE SPANSULE	PAL

ST 15MG CAPSULE (SUSTAINED RELEASE)

02448327	ACT DEXTROAMPHETAMINE SR	ACG
01924567	DEXEDRINE SPANSULE	PAL

ST 5MG TABLET

01924516	DEXEDRINE	PAL
02443236	DEXTROAMPHETAMINE	AAP

LISDEXAMFETAMINE DIMESYLATE

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST 10MG CAPSULE

02439603	VYVANSE	SHI
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ST 20MG CAPSULE

02347156	VYVANSE	SHI
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ST 30MG CAPSULE

02322951	VYVANSE	SHI
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ST 40MG CAPSULE

02347164	VYVANSE	SHI
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ST 50MG CAPSULE

02322978	VYVANSE	SHI
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ST 60MG CAPSULE

02347172	VYVANSE	SHI
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28:20.32 CNS STIMULANTS

METHYLPHENIDATE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST 5MG TABLET

02273950	APO-METHYLPHENIDATE	APX
02326221	METHYLPHENIDATE	PDL
02234749	PMS-METHYLPHENIDATE	PMS

ST 10MG TABLET

02249324	APO-METHYLPHENIDATE	APX
02326248	METHYLPHENIDATE	PDL
00584991	PMS-METHYLPHENIDATE	PMS

ST 20MG TABLET

02249332	APO-METHYLPHENIDATE	APX
02326256	METHYLPHENIDATE	PDL
00585009	PMS-METHYLPHENIDATE	PMS

ST 18MG TABLET (EXTENDED RELEASE)

02441934	ACT METHYLPHENIDATE ER	ACG
02452731	APO-METHYLPHENIDATE ER	APX
02247732	CONCERTA	JSO
02413728	PMS-METHYLPHENIDATE ER	PMS
02315068	TEVA-METHYLPHENIDATE	TEV

ST 20MG TABLET (EXTENDED RELEASE)

02266687	APO-METHYLPHENIDATE SR	APX
02320312	SANDOZ METHYLPHENIDATE SR	SDZ

ST 27MG TABLET (EXTENDED RELEASE)

02441942	ACT METHYLPHENIDATE ER	ACG
02452758	APO-METHYLPHENIDATE ER	APX
02250241	CONCERTA	JSO
02413736	PMS-METHYLPHENIDATE ER	PMS
02315076	TEVA-METHYLPHENIDATE	TEV

ST 36MG TABLET (EXTENDED RELEASE)

02441950	ACT METHYLPHENIDATE ER	ACG
02452766	APO-METHYLPHENIDATE ER	APX
02247733	CONCERTA	JSO
02413744	PMS-METHYLPHENIDATE ER	PMS
02315084	TEVA-METHYLPHENIDATE	TEV

ST 54MG TABLET (EXTENDED RELEASE)

02441969	ACT METHYLPHENIDATE ER	ACG
02330377	APO-METHYLPHENIDATE ER	APX
02247734	CONCERTA	JSO
02413752	PMS-METHYLPHENIDATE ER	PMS
02315092	TEVA-METHYLPHENIDATE	TEV

28:20.80 WAKEFULNESS-PROMOTING AGENTS

MODAFINIL

ST 100MG TABLET

02239665	ALERTEC	TEV
02285398	APO-MODAFINIL	APX
02430487	AURO-MODAFINIL	AUR
02442078	BIO-MODAFINIL	BMI
02432560	MAR-MODAFINIL	MAR
02420260	TEVA-MODAFINIL	TEV

28:20.92 MISC ANOREXIGENIC AGENTS & RESPIRATORY & CEREBRAL STIMULANT

CAFFEINE CITRATE

Limited use benefit (prior approval not required).

For children up to 1 year of age

POWDER

00972037	CAFFEINE CITRATE	MDS
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28:24.04 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BARBITURATES

PHENOBARBITAL

15MG TABLET

00178799	PHENOBARB	PED
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30MG TABLET

00178802	PHENOBARB	PED
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60MG TABLET

00178810	PHENOBARB	PED
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28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

ALPRAZOLAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 0.25MG TABLET

01908189	ALPRAZOLAM	PDL
02349191	ALPRAZOLAM	SAN
00865397	APO-ALPRAZ	APX
02400111	JAMP-ALPRAZOLAM	JMP
01913484	TEVA-ALPRAZOLAM	TEV
00548359	XANAX	PFI

ST 0.5MG TABLET

01908170	ALPRAZOLAM	PDL
02349205	ALPRAZOLAM	SAN
00865400	APO-ALPRAZ	APX

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

ALPRAZOLAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **0.5MG TABLET**

02400138	JAMP-ALPRAZOLAM	JMP
01913492	TEVA-ALPRAZOLAM	TEV
00548367	XANAX	PFI

ST **1MG TABLET**

02248706	ALPRAZOLAM	PDL
02243611	APO-ALPRAZ	APX
02400146	JAMP-ALPRAZOLAM	JMP
00723770	XANAX	PFI

ST **2MG TABLET**

02243612	APO-ALPRAZ	APX
02400154	JAMP-ALPRAZOLAM	JMP
00813958	XANAX TS	PFI

BROMAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **1.5MG TABLET**

02177153	APO-BROMAZEPAM	APX
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ST **3MG TABLET**

02177161	APO-BROMAZEPAM	APX
02220520	BROMAZEPAM	PDL
02230584	TEVA-BROMAZEPAM	TEV

ST **6MG TABLET**

02177188	APO-BROMAZEPAM	APX
02220539	BROMAZEPAM	PDL
02230585	TEVA-BROMAZEPAM	TEV

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

DIAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **1MG/ML SOLUTION**

00891797	PMS-DIAZEPAM	PMS
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ST **2MG TABLET**

00405329	DIAZEPAM	AAP
02247490	PMS-DIAZEPAM	PMS

ST **5MG TABLET**

00313580	DIAZEPAM	PDL
00362158	DIAZEPAM	AAP
02247491	PMS-DIAZEPAM	PMS
00013285	VALIUM	HLR

ST **10MG TABLET**

00405337	DIAZEPAM	AAP
02247492	PMS-DIAZEPAM	PMS

DIAZEPAM (DIASTAT)

Limited use benefit (prior approval not required).

For children 12 years of age or under.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4,000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **5MG/ML GEL**

02238162	DIASTAT	VAE
09853340	DIASTAT 2X10MG RECTAL PACK	ELN
09853430	DIASTAT 2X15MG RECTAL PACK	ELN

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

LORAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **0.5MG TABLET**

00655740	APO-LORAZEPAM	APX
02041413	ATIVAN	PFI
02041456	ATIVAN SUBLINGUAL	PFI
02351072	LORAZEPAM	SAN
02410745	LORAZEPAM SUBLINGUAL	AAP
00728187	PMS-LORAZEPAM	PMS
00655643	PRO-LORAZEPAM	PDL
00711101	TEVA-LORAZEPAM	TEV

ST **1MG TABLET**

00655759	APO-LORAZEPAM	APX
02041421	ATIVAN	PFI
02041464	ATIVAN SUBLINGUAL	PFI
02351080	LORAZEPAM	SAN
02410753	LORAZEPAM SUBLINGUAL	AAP
00728195	PMS-LORAZEPAM	PMS
00655651	PRO-LORAZEPAM	PDL
00637742	TEVA-LORAZEPAM	TEV

ST **2MG TABLET**

00655767	APO-LORAZEPAM	APX
02041448	ATIVAN	PFI
02041472	ATIVAN SUBLINGUAL	PFI
02351099	LORAZEPAM	SAN
02410761	LORAZEPAM SUBLINGUAL	AAP
00728209	PMS-LORAZEPAM	PMS
00655678	PRO-LORAZEPAM	PDL
00637750	TEVA-LORAZEPAM	TEV

NITRAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **5MG TABLET**

00511528	MOGADON	AAP
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28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

NITRAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **10MG TABLET**

00511536	MOGADON	AAP
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OXAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **10MG TABLET**

00402680	APO OXAZEPAM	APX
00497754	OXAZEPAM	PDL
00414247	OXPAM	BMI
00568392	RIVA OXAZEPAM	RIV

ST **15MG TABLET**

00402745	APO OXAZEPAM	APX
00497762	OXAZEPAM	PDL
00568406	RIVA OXAZEPAM	RIV

ST **30MG TABLET**

00402737	APO OXAZEPAM	APX
00497770	OXAZEPAM	PDL
00414263	OXPAM	BMI
00568414	RIVA OXAZEPAM	RIV

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

TEMAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 15MG CAPSULE

00604453	RESTORIL	AAP
02225964	TEMAZEPAM	APX
02229760	TEMAZEPAM	PDL
02230095	TEVA-TEMAZEPAM	TEV

ST 30MG CAPSULE

00604461	RESTORIL	AAP
02225972	TEMAZEPAM	APX
02229761	TEMAZEPAM	PDL
02230102	TEVA-TEMAZEPAM	TEV

TRIAZOLAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 0.25MG TABLET

00808571	TRIAZOLAM	AAP
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28:24.92 MISCELLANEOUS ANXIOLYTICS, SEDATIVES, AND HYPNOTICS

BUSPIRONE HYDROCHLORIDE

ST 10MG TABLET

02211076	APO-BUSPIRONE	APX
02223163	BUSPIRONE	PDL
02447851	BUSPIRONE	SAN
02230942	PMS-BUSPIRONE	PMS
02231492	TEVA-BUSPIRONE	TEV

HYDROXYZINE HYDROCHLORIDE

ST 10MG CAPSULE

00646059	HYDROXYZINE	APX
00738824	NOVO-HYDROXYZIN	TEV

ST 25MG CAPSULE

00646024	HYDROXYZINE	APX
00738832	NOVO-HYDROXYZIN	TEV

28:24.92 MISCELLANEOUS ANXIOLYTICS, SEDATIVES, AND HYPNOTICS

HYDROXYZINE HYDROCHLORIDE

ST 50MG CAPSULE

00646016	HYDROXYZINE	APX
00738840	NOVO-HYDROXYZIN	TEV

ST 2MG/ML SYRUP

00024694	ATARAX	ERF
00741817	PMS HYDROXYZINE	PMS

28:28.00 ANTIMANIC AGENTS

LITHIUM CARBONATE

ST 150MG CAPSULE

02242837	APO-LITHIUM CARBONATE	APX
09857532	APO-LITHIUM CARBONATE	APX
00461733	CARBOLITH	VAE
02013231	LITHANE	ERF
02216132	PMS-LITHIUM CARBONATE	PMS

ST 300MG CAPSULE

02242838	APO-LITHIUM CARBONATE	APX
09857540	APO-LITHIUM CARBONATE	APX
00236683	CARBOLITH	VAE
00406775	LITHANE	ERF
02216140	PMS-LITHIUM CARBONATE	PMS

ST 600MG CAPSULE

02011239	CARBOLITH	VAE
02216159	PMS-LITHIUM CARBONATE	PMS

ST 300MG TABLET (EXTENDED RELEASE)

02266695	LITHMAX	AAP
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LITHIUM CITRATE

ST 60MG/ML SYRUP

02074834	PMS-LITHIUM CITRATE	PMS
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28:32.28 SELECTIVE SEROTONIN AGONISTS

ALMOTRIPTAN MALATE

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

6.25MG TABLET

02405792	APO-ALMOTRIPTAN	APX
02248128	AXERT	MCL
02398435	MYLAN-ALMOTRIPTAN	MYL

12.5MG TABLET

02424029	ALMOTRIPTAN	PDL
02466821	ALMOTRIPTAN	SAN
02405806	APO-ALMOTRIPTAN	APX
02248129	AXERT	MCL
02398443	MYLAN-ALMOTRIPTAN	MYL
02405334	SANDOZ ALMOTRIPTAN	SDZ
02434849	TEVA-ALMOTRIPTAN	TEV

NARATRIPTAN HYDROCHLORIDE

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

1MG TABLET

02237820	AMERGE	GSK
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28:32.28 SELECTIVE SEROTONIN AGONISTS

NARATRIPTAN HYDROCHLORIDE

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

1MG TABLET

02314290 TEVA-NARATRIPTAN TEV

2.5MG TABLET

02237821 AMERGE GSK

02322323 SANDOZ NARATRIPTAN SDZ

02314304 TEVA-NARATRIPTAN TEV

RIZATRIPTAN BENZOATE

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

5MG TABLET

02393468 APO-RIZATRIPTAN APX

02380455 JAMP-RIZATRIPTAN JMP

02429233 JAMP-RIZATRIPTAN IR JMP

02379651 MAR-RIZATRIPTAN MAR

02428512 VAN-RIZATRIPTAN VAN

10MG TABLET

02381702 ACT RIZATRIPTAN ACG

02393476 APO-RIZATRIPTAN APX

02441144 AURO-RIZATRIPTAN AUR

02380463 JAMP-RIZATRIPTAN JMP

02429241 JAMP-RIZATRIPTAN IR JMP

02379678 MAR-RIZATRIPTAN MAR

02240521 MAXALT FRS

02428520 VAN-RIZATRIPTAN VAN

5MG TABLET (ORALLY DISINTEGRATING)

02374730 ACT RIZATRIPTAN ODT ACG

02393484 APO-RIZATRIPTAN RPD APX

02465086 JAMP-RIZATRIPTAN ODT JMP

02462788 MAR-RIZATRIPTAN ODT MAR

02240518 MAXALT RPD FRS

02379198 MYLAN-RIZATRIPTAN ODT MYL

02436604 NAT-RIZATRIPTAN ODT NPH

02393360 PMS-RIZATRIPTAN RDT PMS

02442906 RIZATRIPTAN ODT SAN

02446111 RIZATRIPTAN ODT SIV

02415798 RIZATRIPTAN RDT PDL

02351870 SANDOZ RIZATRIPTAN ODT SDZ

02396661 TEVA-RIZATRIPTAN ODT TEV

10MG TABLET (ORALLY DISINTEGRATING)

02374749 ACT RIZATRIPTAN ODT ACG

02393492 APO-RIZATRIPTAN RPD APX

02396203 DOM-RIZATRIPTAN RDT DPC

02465094 JAMP-RIZATRIPTAN RDT JMP

02462796 MAR-RIZATRIPTAN ODT MAR

02240519 MAXALT RPD FRS

02379201 MYLAN-RIZATRIPTAN ODT MYL

02436612 NAT-RIZATRIPTAN ODT NPH

02393379 PMS-RIZATRIPTAN RDT PMS

02442914 RIZATRIPTAN ODT SAN

02446138 RIZATRIPTAN ODT SIV

28:32.28 SELECTIVE SEROTONIN AGONISTS

RIZATRIPTAN BENZOATE

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

10MG TABLET (ORALLY DISINTEGRATING)

02415801 RIZATRIPTAN RDT PDL

02351889 SANDOZ RIZATRIPTAN ODT SDZ

02396688 TEVA-RIZATRIPTAN ODT TEV

02448505 VAN-RIZATRIPTAN ODT VAN

SUMATRIPTAN HEMISULFATE

5MG SPRAY

02230418 IMITREX GSK

20MG SPRAY

02230420 IMITREX GSK

SUMATRIPTAN SUCCINATE

Limited use benefit (prior approval is not required).

A total of 12 tablets (or injections) are permitted in a 30-day period.

6MG/0.5ML INJECTION

99000598 IMITREX STAT DOSE KIT GSK

12MG/ML SOLUTION

02212188 IMITREX GSK

02361698 TARO-SUMATRIPTAN TAR

25MG TABLET

02270749 DOM-SUMATRIPTAN DPC

02268906 MYLAN-SUMATRIPTAN MYL

02256428 PMS-SUMATRIPTAN PMS

02286815 TEVA-SUMATRIPTAN DF TEV

50MG TABLET

02268388 APO-SUMATRIPTAN APX

02270757 DOM-SUMATRIPTAN DPC

02212153 IMITREX DF GSK

02268914 MYLAN-SUMATRIPTAN MYL

02256436 PMS-SUMATRIPTAN PMS

02263025 SANDOZ SUMATRIPTAN SDZ

02286521 SUMATRIPTAN SAN

02324652 SUMATRIPTAN PDL

02385570 SUMATRIPTAN DF SIV

02286823 TEVA-SUMATRIPTAN DF TEV

100MG TABLET

02257904 ACT SUMATRIPTAN ACG

02268396 APO-SUMATRIPTAN APX

02270765 DOM-SUMATRIPTAN DPC

02212161 IMITREX DF GSK

02268922 MYLAN-SUMATRIPTAN MYL

02256444 PMS-SUMATRIPTAN PMS

02263033 SANDOZ SUMATRIPTAN SDZ

02286548 SUMATRIPTAN SAN

02324660 SUMATRIPTAN PDL

02385589 SUMATRIPTAN DF SIV

02239367 TEVA-SUMATRIPTAN TEV

02286831 TEVA-SUMATRIPTAN DF TEV

28:32.28 SELECTIVE SEROTONIN AGONISTS

ZOLMITRIPTAN

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

2.5MG SPRAY

02248992 ZOMIG AZC

5MG SPRAY

02248993 ZOMIG AZC

2.5MG TABLET

02380951 APO-ZOLMITRIPTAN APX

02389525 DOM-ZOLMITRIPTAN DPC

02421623 JAMP-ZOLMITRIPTAN JMP

02399458 MAR-ZOLMITRIPTAN MAR

02419521 MINT-ZOLMITRIPTAN MIN

02421534 NAT-ZOLMITRIPTAN NPH

02324229 PMS-ZOLMITRIPTAN PMS

02401304 RIVA-ZOLMITRIPTAN RIV

02362988 SANDOZ ZOLMITRIPTAN SDZ

02313960 TEVA-ZOLMITRIPTAN TEV

02379929 ZOLMITRIPTAN PDL

02238660 ZOMIG AZC

2.5MG TABLET (ORALLY DISINTEGRATING)

02438453 AG-ZOLMITRIPTAN ODT ANG

02381575 APO-ZOLMITRIPTAN RAPID APX

02428237 JAMP-ZOLMITRIPTAN ODT JMP

02324768 PMS-ZOLMITRIPTAN ODT PMS

02362996 SANDOZ ZOLMITRIPTAN ODT SDZ

02428474 SEPTA-ZOLMITRIPTAN-ODT SPT

02342545 TEVA-ZOLMITRIPTAN OD TEV

02438763 VAN-ZOLMITRIPTAN ODT VAN

02379988 ZOLMITRIPTAN ODT PDL

02243045 ZOMIG RAPIMELT AZC

28:32.92 MISCELLANEOUS ANTIMIGRANE AGENTS

FLUNARIZINE HYDROCHLORIDE

ST **5MG CAPSULE**

02246082 FLUNARIZINE AAP

PIZOTIFEN MALATE

0.5MG TABLET

00329320 SANDOMIGRAN PAL

1MG TABLET

00511552 SANDOMIGRAN DS PAL

28:36.08 ANTIPARKINSONIAN AGENTS - ANTICHOLINERGIC AGENTS

BENZTROPINE MESYLATE

1MG/ML LIQUID

02238903 BENZTROPINE OMEGA OMG

ST **1MG TABLET**

00706531 PDP-BENZTROPINE PED

ST **2MG TABLET**

00426857 PDP-BENZTROPINE PED

00587265 PMS-BENZTROPINE PMS

28:36.08 ANTIPARKINSONIAN AGENTS - ANTICHOLINERGIC AGENTS

ETHOPROPAZINE HYDROCHLORIDE

50MG TABLET

01927744 PARSITAN ERF

PROCYCLIDINE HCL

5MG CAPSULE

99101405 PROCYCLIDINE (PQ) UNK

PROCYCLIDINE HYDROCHLORIDE

0.5MG/ML ELIXIR

00587362 PDP-PROCYCLIDINE PED

2.5MG TABLET

00649392 PDP-PROCYCLIDINE PED

5MG TABLET

00587354 PDP-PROCYCLIDINE PED

TRIHEXYPHENIDYL HYDROCHLORIDE

0.4MG/ML ELIXIR

00885398 PMS-TRIHEXYPHENIDYL PMS

2MG TABLET

00545058 TRIHEXYPHENIDYL AAP

5MG TABLET

00545074 TRIHEXYPHENIDYL AAP

28:36.12 ANTIPARKINSONIAN AGENTS - CATECHOL-O-METHYLTRANSFERASE (COMT) INHIBITORS

ENTACAPONE

ST **200MG TABLET**

02243763 COMTAN NVR

02380005 SANDOZ ENTACAPONE SDZ

02375559 TEVA-ENTACAPONE TEV

28:36.16 ANTIPARKINSONIAN AGENTS - DOPAMINE PRECURSORS

LEVODOPA, BENSERAZIDE HYDROCHLORIDE

ST **50MG & 12.5MG CAPSULE**

00522597 PROLOPA HLR

ST **100MG & 25MG CAPSULE**

00386464 PROLOPA HLR

ST **200MG & 50MG CAPSULE**

00386472 PROLOPA HLR

LEVODOPA, CARBIDOPA

ST **100MG & 10MG TABLET**

02195933 APO-LEVOCARB APX

02457954 MINT-LEVOCARB MIN

02244494 TEVA-LEVOCARBIDOPA TEV

ST **100MG & 25MG TABLET**

02195941 APO-LEVOCARB APX

02457962 MINT-LEVOCARB MIN

02421488 PMS-LEVOCARB PMS

02311178 PRO-LEVOCARB PDL

00513997 SINEMET FRS

02244495 TEVA-LEVOCARBIDOPA TEV

**28:36.16 ANTIPARKINSONIAN AGENTS -
DOPAMINE PRECURSORS**

LEVODOPA, CARBIDOPA

ST 250MG & 25MG TABLET

02195968	APO-LEVOCARB	APX
02457970	MINT-LEVOCARB	MIN
00328219	SINEMET	FRS
02244496	TEVA-LEVOCARBIDOPA	TEV

ST 100MG & 25MG TABLET (EXTENDED RELEASE)

02272873	APO-LEVOCARB	APX
02028786	SINEMET	FRS

ST 200MG & 50MG TABLET (EXTENDED RELEASE)

02245211	APO-LEVOCARB	APX
02421496	PMS-LEVOCARB	PMS

LEVODOPA, CARBIDOPA, ENTACAPONE

ST 50MG & 12.5MG & 200MG TABLET

02305933	STALEVO	NVR
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ST 75MG & 18.75MG & 200MG TABLET

02337827	STALEVO	NVR
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ST 100MG & 25MG & 200MG TABLET

02305941	STALEVO	NVR
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ST 125MG & 31.25MG & 200MG TABLET

02337835	STALEVO	NVR
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ST 150MG & 37.5MG & 200MG TABLET

02305968	STALEVO	NVR
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**28:36.20 ANTIPARKINSONIAN AGENTS -
DOPAMINE RECEPTOR
AGONISTS**

BROMOCRIPTINE MESYLATE

ST 5MG CAPSULE

02230454	BROMOCRIPTINE	AAP
02238637	DOM-BROMOCRIPTINE	DPC
02236949	PMS-BROMOCRIPTINE	PMS

ST 2.5MG TABLET

02087324	BROMOCRIPTINE	AAP
02238636	DOM-BROMOCRIPTINE	DPC
02231702	PMS-BROMOCRIPTINE	PMS

CABERGOLINE

Limited use benefit (prior approval required).

For treatment of hyperprolactinemia in patients who have failed therapy with or are intolerant to bromocriptine.

0.5MG TABLET

02301407	ACT CABERGOLINE	ACG
02455897	APO-CABERGOLINE	APX
02242471	DOSTINEX	PFI

PRAMIPEXOLE DIHYDROCHLORIDE

ST 0.25MG TABLET

02297302	ACT PRAMIPEXOLE	ACG
02292378	APO-PRAMIPEXOLE	APX
02424061	AURO-PRAMIPEXOLE	AUR
02309017	DOM-PRAMIPEXOLE	DPC
02237145	MIRAPEX	BOE
09857268	MIRAPEX (ON)	BOE
02290111	PMS-PRAMIPEXOLE	PMS

**28:36.20 ANTIPARKINSONIAN AGENTS -
DOPAMINE RECEPTOR
AGONISTS**

PRAMIPEXOLE DIHYDROCHLORIDE

ST 0.25MG TABLET

02309122	PRAMIPEXOLE	SIV
02325802	PRAMIPEXOLE	PDL
02315262	SANDOZ PRAMIPEXOLE	SDZ

ST 0.5MG TABLET

02297310	ACT PRAMIPEXOLE	ACG
02292386	APO-PRAMIPEXOLE	APX
02424088	AURO-PRAMIPEXOLE	AUR
02290138	PMS-PRAMIPEXOLE	PMS
02309130	PRAMIPEXOLE	SIV
02325810	PRAMIPEXOLE	PDL
02315270	SANDOZ PRAMIPEXOLE	SDZ

ST 1MG TABLET

02297329	ACT PRAMIPEXOLE	ACG
02292394	APO-PRAMIPEXOLE	APX
02424096	AURO-PRAMIPEXOLE	AUR
02290146	PMS-PRAMIPEXOLE	PMS
02309149	PRAMIPEXOLE	SIV
02325829	PRAMIPEXOLE	PDL
02315289	SANDOZ PRAMIPEXOLE	SDZ

ST 1.5MG TABLET

02297337	ACT PRAMIPEXOLE	ACG
02292408	APO-PRAMIPEXOLE	APX
02424118	AURO-PRAMIPEXOLE	AUR
02290154	PMS-PRAMIPEXOLE	PMS
02309157	PRAMIPEXOLE	SIV
02325837	PRAMIPEXOLE	PDL
02315297	SANDOZ PRAMIPEXOLE	SDZ

ROPINIROLE HYDROCHLORIDE

ST 0.25MG TABLET

02316846	ACT ROPINIROLE	ACG
02337746	APO-ROPINIROLE	APX
02352338	JAMP-ROPINIROLE	JMP
02326590	PMS-ROPINIROLE	PMS
02314037	RAN-ROPINIROLE	RBV
02353040	ROPINIROLE	SAN

ST 1MG TABLET

02316854	ACT ROPINIROLE	ACG
02337762	APO-ROPINIROLE	APX
02352346	JAMP-ROPINIROLE	JMP
02326612	PMS-ROPINIROLE	PMS
02314053	RAN-ROPINIROLE	RBV
02353059	ROPINIROLE	SAN

ST 2MG TABLET

02316862	ACT ROPINIROLE	ACG
02337770	APO-ROPINIROLE	APX
02352354	JAMP-ROPINIROLE	JMP
02326620	PMS-ROPINIROLE	PMS
02314061	RAN-ROPINIROLE	RBV

ST 5MG TABLET

02316870	ACT ROPINIROLE	ACG
02337800	APO-ROPINIROLE	APX

**28:36.20 ANTIPARKINSONIAN AGENTS -
DOPAMINE RECEPTOR
AGONISTS**

ROPINIROLE HYDROCHLORIDE

ST **5MG TABLET**

02352362	JAMP-ROPINIROLE	JMP
02326639	PMS-ROPINIROLE	PMS
02314088	RAN-ROPINIROLE	RBV

ROTIGOTINE

Limited use benefit (prior approval required).

As an adjunct to levodopa for the treatment of patients with advanced stage Parkinson's disease; AND

Patient is currently receiving treatment with levodopa.

2MG PATCH

02403900	NEUPRO	UCB
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4MG PATCH

02403927	NEUPRO	UCB
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6MG PATCH

02403935	NEUPRO	UCB
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8MG PATCH

02403943	NEUPRO	UCB
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**28:36.32 ANTIPARKINSONIAN AGENTS -
MONOAMINE OXIDASE B
INHIBITORS**

SELEGILINE HYDROCHLORIDE

ST **5MG TABLET**

02230641	APO-SELEGILINE	APX
02068087	TEVA-SELEGILINE	TEV

**28:92.00 MISCELLANEOUS CENTRAL
NERVOUS SYSTEM AGENTS**

ACAMPROSATE CALCIUM

Limited use benefit (prior approval required).

For patients who have been abstinent from alcohol for at least four days and where available, are currently enrolled in an alcohol addiction treatment program.

333MG TABLET (DELAYED RELEASE)

02293269	CAMPRAL	MYL
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ATOMOXETINE HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with Attention Deficit Hyperactivity Disorder (ADHD) who meet one of the following criteria:

- Failure or intolerance to methylphenidate or amphetamine; OR
- Contraindication to stimulant medication; OR
- Potential risk of stimulant misuse or diversion; OR
- Prescribed or recommended by a pediatrician or a psychiatrist.

10MG CAPSULE

02318024	APO-ATOMOXETINE	APX
02358190	ATOMOXETINE	AAP
02396904	ATOMOXETINE	PDL
02445883	ATOMOXETINE	SIV

**28:92.00 MISCELLANEOUS CENTRAL
NERVOUS SYSTEM AGENTS**

ATOMOXETINE HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with Attention Deficit Hyperactivity Disorder (ADHD) who meet one of the following criteria:

- Failure or intolerance to methylphenidate or amphetamine; OR
- Contraindication to stimulant medication; OR
- Potential risk of stimulant misuse or diversion; OR
- Prescribed or recommended by a pediatrician or a psychiatrist.

10MG CAPSULE

02467747	ATOMOXETINE	SAN
02390469	DOM-ATOMOXETINE	DPC
02381028	PMS-ATOMOXETINE	PMS
02405962	RIVA-ATOMOXETINE	RIV
02386410	SANDOZ ATOMOXETINE	SDZ
02262800	STRATTERA	LIL
02314541	TEVA-ATOMOXETINE	TEV

18MG CAPSULE

02318032	APO-ATOMOXETINE	APX
02358204	ATOMOXETINE	AAP
02396912	ATOMOXETINE	PDL
02445905	ATOMOXETINE	SIV
02467755	ATOMOXETINE	SAN
02390477	DOM-ATOMOXETINE	DPC
02381036	PMS-ATOMOXETINE	PMS
02405970	RIVA-ATOMOXETINE	RIV
02386429	SANDOZ ATOMOXETINE	SDZ
02262819	STRATTERA	LIL
02314568	TEVA-ATOMOXETINE	TEV

25MG CAPSULE

02318040	APO-ATOMOXETINE	APX
02358212	ATOMOXETINE	AAP
02396920	ATOMOXETINE	PDL
02445913	ATOMOXETINE	SIV
02467763	ATOMOXETINE	SAN
02390485	DOM-ATOMOXETINE	DPC
02381044	PMS-ATOMOXETINE	PMS
02405989	RIVA-ATOMOXETINE	RIV
02386437	SANDOZ ATOMOXETINE	SDZ
02262827	STRATTERA	LIL
02314576	TEVA-ATOMOXETINE	TEV

40MG CAPSULE

02318059	APO-ATOMOXETINE	APX
02358220	ATOMOXETINE	AAP
02396939	ATOMOXETINE	PDL
02445948	ATOMOXETINE	SIV
02467771	ATOMOXETINE	SAN
02390493	DOM-ATOMOXETINE	DPC
02381052	PMS-ATOMOXETINE	PMS
02405997	RIVA-ATOMOXETINE	RIV
02386445	SANDOZ ATOMOXETINE	SDZ
02262835	STRATTERA	LIL
02314584	TEVA-ATOMOXETINE	TEV

28:92.00 MISCELLANEOUS CENTRAL NERVOUS SYSTEM AGENTS

ATOMOXETINE HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with Attention Deficit Hyperactivity Disorder (ADHD) who meet one of the following criteria:

- Failure or intolerance to methylphenidate or amphetamine; OR
- Contraindication to stimulant medication; OR
- Potential risk of stimulant misuse or diversion; OR
- Prescribed or recommended by a pediatrician or a psychiatrist.

60MG CAPSULE

02318067	APO-ATOMOXETINE	APX
02358239	ATOMOXETINE	AAP
02396947	ATOMOXETINE	PDL
02445956	ATOMOXETINE	SIV
02467798	ATOMOXETINE	SAN
02390515	DOM-ATOMOXETINE	DPC
02381060	PMS-ATOMOXETINE	PMS
02406004	RIVA-ATOMOXETINE	RIV
02386453	SANDOZ ATOMOXETINE	SDZ
02262843	STRATTERA	LIL
02314592	TEVA-ATOMOXETINE	TEV

80MG CAPSULE

02318075	APO-ATOMOXETINE	APX
02358247	ATOMOXETINE	AAP
02467801	ATOMOXETINE	SAN
02404664	PMS-ATOMOXETINE	PMS
02422824	RIVA-ATOMOXETINE	RIV
02386461	SANDOZ ATOMOXETINE	SDZ
02279347	STRATTERA	LIL
02362511	TEVA-ATOMOXETINE	TEV

100MG CAPSULE

02318083	APO-ATOMOXETINE	APX
02358255	ATOMOXETINE	AAP
02467828	ATOMOXETINE	SAN
02404672	PMS-ATOMOXETINE	PMS
02422832	RIVA-ATOMOXETINE	RIV
02386488	SANDOZ ATOMOXETINE	SDZ
02279355	STRATTERA	LIL
02362538	TEVA-ATOMOXETINE	TEV

BETAHISTINE HYDROCHLORIDE

8MG TABLET

02449145	AURO-BETAHISTINE	AUR
02280183	TEVA-BETAHISTINE	TEV

16MG TABLET

02449153	AURO-BETAHISTINE	AUR
02466449	BETAHISTINE	SAN
02330210	PMS-BETAHISTINE	PMS
02243878	SERC	BGP
02280191	TEVA-BETAHISTINE	TEV

24MG TABLET

02449161	AURO-BETAHISTINE	AUR
02466457	BETAHISTINE	SAN
02330237	PMS-BETAHISTINE	PMS

28:92.00 MISCELLANEOUS CENTRAL NERVOUS SYSTEM AGENTS

BETAHISTINE HYDROCHLORIDE

24MG TABLET

02247998	SERC	BGP
02280205	TEVA-BETAHISTINE	TEV

DIMETHYL FUMARATE

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

120MG CAPSULE (DELAYED RELEASE)

02404508	TECFIDERA	UNK
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240MG CAPSULE (DELAYED RELEASE)

02420201	TECFIDERA	UNK
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TETRABENAZINE

25MG TABLET

02407590	APO-TETRABENAZINE	APX
02199270	NITOMAN	VAE
02402424	PMS-TETRABENAZINE	PMS
02410338	TETRABENAZINE	RAX

32:00 CONTRACEPTIVES (NON-ORAL)

32:00.00 CONTRACEPTIVES (NON-ORAL)

CONDOM

DEVICE

99400527	CONDOM, LATEX, LUBRICATED	UNK
99400486	CONDOM, LATEX, NON-LUBRICATED	UNK
99400786	CONDOM, NON-LATEX, LUBRICATED	UNK
09991648	FC2 FEMALE CONDOMS	UNK

CONTRACEPTIVE

DEVICE

09991647	TODAY SPONGE VAGINAL CONTRACEPTIVE	UNK
09991646	VCF VAGINAL CONTRACEPTIVE FILM	UNK

FOAM

09991645	VCF FOAM VAGINAL CONTRACEPTIVE	UNK
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CONTRACEPTIVE DEVICE

DEVICE

00970905	CAYA CONTOURED DIAPHRAGM	TSN
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FEMCAP

DEVICE

09991642	CERVICAL	UNK
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INTRAUTERINE DEVICE

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 1 device every 12 months.

DEVICE

00970328	FLEXI-T +300 IUD	TSN
00970336	FLEXI-T +380 IUD	TSN
98099999	FLEXI-TD	TSN
99401085	LIBERTE UT380 SHORT IUD	MSF
99401086	LIBERTE UT380 STANDARD IUD	MSF
00970379	MONA LISA 10	SEA
00970387	MONA LISA 5	SEA
00970395	MONA LISA N	SEA
99400482	NOVA-T	BEX

36:00 DIAGNOSTIC AGENTS (DX)

36:26.00 DX - DIABETES MELLITUS

GLUCOSE OXIDASE, PEROXIDASE

Limited use benefit (prior approval not required).

The number of test strips that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 500 test strips per 100 days. A client can test up to five times per day.
- Clients managing diabetes with diabetes medication with a high risk of causing low blood sugar will be allowed 400 test strips per 365 days. A client can test once daily.
- Clients managing diabetes with diabetes medication with a low risk of causing low blood sugar will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Clients managing diabetes with diet/lifestyle therapy only (no insulin or diabetes medications) will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Non-diabetic clients with rare conditions leading to symptomatic hypo or hyperglycemia will be allowed 500 test strips per 100 days.

STRIP

09857563 ACCU-CHEK GUIDE (ON) ROD
 97799177 ACCU-CHEK GUIDE (SK) ROD

ACCU-CHEK ADVANTAGE STRIP

09853626 ACCU-CHEK ADVANTAGE ROD
 97799824 ACCU-CHEK ADVANTAGE ROD

ACCU-CHEK AVIVA STRIP

09857178 ACCU-CHEK AVIVA ROD
 97799814 ACCU-CHEK AVIVA ROD

ACCU-CHEK COMPACT STRIP

09854282 ACCU-CHEK COMPACT ROD
 97799962 ACCU-CHEK COMPACT ROD

ACCU-CHEK MOBILE STRIP

09857452 ACCU-CHEK MOBILE BG ROD
 97799497 ACCU-CHEK MOBILE CASSETT ROD

ACCUTREND STRIP

09853162 ACCUTREND ROD
 97799959 ACCUTREND ROD

ASCENSIA BREEZE 2 STRIP

97799748 ASCENSIA BREEZE 2 BAY
 09857293 BREEZE 2 BG (ON) BAY

ASCENSIA CONTOUR STRIP

97799702 ASCENSIA CONTOUR BAY
 09857127 CONTOUR BG (ON) BAY

BG STAR STRIP

97799465 BG STAR SAC
 09857422 BG STAR (ON) SAC

CONTOUR NEXT STRIP

97799459 CONTOUR NEXT BAY
 09857453 CONTOUR NEXT (ON) BAY

EZ HEALTH STRIP

09857357 EZ HEALTH ORACLE TRE
 97799564 EZ HEALTH ORACLE TRE

FREESTYLE STRIP

97799829 FREESTYLE ABB
 09857141 FREESTYLE (ON) ABB

36:26.00 DX - DIABETES MELLITUS

GLUCOSE OXIDASE, PEROXIDASE

Limited use benefit (prior approval not required).

The number of test strips that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 500 test strips per 100 days. A client can test up to five times per day.
- Clients managing diabetes with diabetes medication with a high risk of causing low blood sugar will be allowed 400 test strips per 365 days. A client can test once daily.
- Clients managing diabetes with diabetes medication with a low risk of causing low blood sugar will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Clients managing diabetes with diet/lifestyle therapy only (no insulin or diabetes medications) will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Non-diabetic clients with rare conditions leading to symptomatic hypo or hyperglycemia will be allowed 500 test strips per 100 days.

FREESTYLE LITE STRIP

97799597 FREESTYLE LITE ABB
 09857297 FREESTYLE LITE (ON) ABB

FREESTYLE PRECISION STRIP

97799346 FREESTYLE PRECISION ABB
 09857502 FREESTYLE PRECISION (ON) ABB

GE200 STRIP

97799373 GE200 AUC
 09857525 GE200 (ON) AUC

ITEST STRIP

09857348 ITEST AUC
 97799692 ITEST AUC

MEDI+SURE STRIP

97799403 MEDI+SURE MEC
 09857432 MEDI+SURE (ON) MEC

NOVA MAX STRIP

09857313 NOVA MAX NCA

ONE TOUCH ULTRA STRIP

09854290 ONE TOUCH ULTRA JAJ
 97799985 ONE TOUCH ULTRA JAJ

ONE TOUCH VERIO STRIP

97799475 ONETOUCH VERIO JAJ
 09857392 ONETOUCH VERIO (ON) JAJ

PRECISION XTRA STRIP

09854070 PRECISION XTRA ABB
 97799840 PRECISION XTRA AUC

SIDEKICK STRIP

97799601 SIDEKICK HOD

SPIRIT STRIP

97799291 FIRST CANHEALTH SPIRIT ARA
 09857547 SPIRIT TEST STRIP (ON) ARA

SURE STEP STRIP

97799355 SURE STEP SKY

SURETEST STRIP

09857522 SURETEST (ON) SKY

TRUETEST STRIP

97799532 TRUETEST HOD

36:26.00 DX - DIABETES MELLITUS

GLUCOSE OXIDASE, PEROXIDASE

Limited use benefit (prior approval not required).

The number of test strips that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 500 test strips per 100 days. A client can test up to five times per day.
- Clients managing diabetes with diabetes medication with a high risk of causing low blood sugar will be allowed 400 test strips per 365 days. A client can test once daily.
- Clients managing diabetes with diabetes medication with a low risk of causing low blood sugar will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Clients managing diabetes with diet/lifestyle therapy only (no insulin or diabetes medications) will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Non-diabetic clients with rare conditions leading to symptomatic hypo or hyperglycemia will be allowed 500 test strips per 100 days.

TRUETRACK STRIP

09857283	TRUE TRACK	AUC
97799602	TRUE TRACK	HOD

36:60.00 DX - THYROID FUNCTION

THYROTROPIN ALFA

0.9MG/ML POWDER FOR SOLUTION

02246016	THYROGEN	GEE
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36:88.00 DX - URINE AND FECES

CONTENTS

URINE TEST STRIP

STRIP

97799914	DIASTIX	BAY
97799913	KETOSTIX	BAY

**40:00 ELECTROLYTIC, CALORIC,
AND WATER BALANCE**

40:08.00 ALKALINIZING AGENTS

CITRIC ACID, SODIUM CITRATE

66.8MG & 100MG/ML SOLUTION

00721344 DICITRATE PMS

POTASSIUM CITRATE

1080MG TABLET

02243768 KCITRA 10 UNK

SODIUM BICARBONATE

325MG TABLET

00481912 XENEX SODIUM BICARBONATE XEN

40:10.00 AMMONIA DETOXICANTS

LACTULOSE

667MG SOLUTION

02469391 PMS-LACTULOSE-PHARMA PMS

ST **667MG/ML SYRUP**

02242814 APO-LACTULOSE APX

02295881 JAMP-LACTULOSE JMP

02412268 LACTULOSE SAN

02247383 PHARMA-LACTULOSE PMS

00703486 PMS-LACTULOSE PMS

00854409 RATIO-LACTULOSE TEV

02331551 TEVA-LACTULOSE TEV

40:12.00 REPLACEMENT PREPARATIONS

CALCIUM

ST **500MG CAPLET**

80001408 OYSTER SHELL CALCIUM NUR

80001122 PHARMA-CAL PED

ST **5ML LIQUID**

80004123 CARBOCAL EUR

ST **20MG/ML LIQUID**

80054754 M-CAL MAN

80002626 SOLUCAL JMP

80006877 WAMPOLE MINERAL CALCIUM WAM

ST **100MG LIQUID**

80043628 NU-CAL ODN

80025527 SOLUCAL GREEN APPLE JMP

80025523 SOLUCAL RASPBERRY JMP

ST **500MG TABLET**

00682039 APOCAL APX

80017732 CAL500 PDL

02240240 CALCIUM PMT

02246040 CALCIUM JMP

80003658 CALCIUM WNP

80076097 CALCIUM UNK

80003773 CALCIUM 500 TRI

80062015 CALCIUM CARBONATE SAN

02237352 EUROCAL EUR

80055526 M-CAL MAN

00618098 NU-CAL ODN

00622443 O-CALCIUM VTH

80079608 PROCAL 500 PDL

40:12.00 REPLACEMENT PREPARATIONS

CALCIUM

ST **500MG TABLET**

00705373 WAMPOLE CALCIUM WAM

02239356 WAMPOLE CALCIUM WAM

ST **500MG TABLET (CHEWABLE)**

80027026 JAMP-CALCIUM CARBONATE JMP

500MG TABLET (FILM COATED)

80066648 BIOCALCIUM BMI

CALCIUM GLUCONATE, VIT D

ST **25MCG LIQUID**

80068920 SOLUCAL D FORT CITRUS JMP

80069353 SOLUCAL D FORT GREEN APPLE JMP

CALCIUM, VITAMIN D

ST **10MG CAPLET**

80008566 PROCALD 400 PDL

ST **500MG & 400IU CAPLET**

80012594 BIOCALD FORTE BMI

ST **500MG LIQUID**

80025543 SOLUCAL D CITRUS JMP

80025541 SOLUCAL D RASPBERRY JMP

ST **500MG & 1,000IU LIQUID**

80025038 SOLUCAL D FORT JMP

ST **500MG & 400IU LIQUID**

80061575 CALCITE LIQUIDE D 400 RIV

80054755 M-CAL D MAN

80008126 SOLUCAL D JMP

ST **500MG & 800IU LIQUID**

80025722 JAMP CALCIUM LACTOGLUCONATE VITAMIN D JMP

500MG & 1,000IU TABLET

80066093 CALCIUM 500 VITAMINE D1000 UNK

80018540 JAMP CALCIUM CARBONATE VITAMIN D JMP

80019536 M CALCIUM VITAMINE D MAN

ST **500MG & 400IU TABLET**

80004963 CALCITE 500 D 400 RIV

80004969 CALCIUM 500 D 400 TRI

80066082 CALCIUM 500 VITAMINE D400 UNK

80066089 CALCIUM 500 VITAMINE D400 UNK

80002623 CALCIUM VITAMIN D LEMON JMP

FLAVOUR

80017190 CALD 400 PDL

80009628 CALODAN D 400 ODN

02245511 CARBOCAL D EUR

80002901 CARBOCAL D EUR

99100832 JAMP-CALCIUM + VITAMIN D JMP

80002122 J-CAL+D JMP

80025360 J-CAL+D JMP

80013329 M-CAL D MAN

80002703 NU-CAL D ODN

80020974 OPUS CAL D OPU

80065914 RIVA-CAL D RIV

80006794 WAMPOLE CALCIUM VITAMIN D WAM

ST **500MG & 800IU TABLET**

80019533 M CALCIUM VITAMINE D MAN

40:12.00 REPLACEMENT PREPARATIONS

CALCIUM, VITAMIN D

ST 500MG & 1,000IU TABLET (CHEWABLE)			
80029083	JAMP CALCIUM CITRATE VITAMIN D	JMP	
80027787	JAMP-CALCIUM VITAMIN D	JMP	
80050701	M-CAL D	MAN	
ST 500MG & 400IU TABLET (CHEWABLE)			
80009412	CALCIUM CARBONATE VITAMINE D	MAN	
ST 600MG & 400IU TABLET (CHEWABLE)			
80021716	WAMPOLE CALCIUM AND D	WAM	
500MG & 400IU TABLET (FILM COATED)			
80066647	BIOCALCIUMD	BMI	

ELECTROLYTES

ST MISCELLANEOUS			
80023410	HYDRALYTE ELECTROLYTE	HYD	
ST 3.56G & 300MG & 470MG & 530MG POWDER			
01931563	GASTROLYTE REGULAR	SAC	
ST POWDER FOR SOLUTION			
80026860	HYDRALYTE ELECTROLYTE	HYD	
80027403	JAMP REHYDRALYTE	JMP	
ST 0.856MG/ML SOLUTION			
80026861	HYDRALYTE ELECTROLYTE	HYD	
ST 25MG & 2.2MG & 2.2MG & 0.9MG/ML SOLUTION			
00630365	PEDIALYTE	ABB	
02219883	PEDIATRIC ELECTROLYTE	PMS	

MAGNESIUM

25MG CAPLET			
80005079	MAGNESIUM COMPLEX	JAM	
100MG TABLET			
80041590	JAMP-MAGNESIUM	JMP	
02068400	MAGNESIUM	JAM	

MAGNESIUM GLUCOHEPTONATE

ST 25MG LIQUID			
80009357	MAGNESIUM	JMP	
ST 100MG/ML ORAL LIQUID			
00026697	ROUGIER-MAGNESIUM	TEV	
ST 100MG/ML SOLUTION			
80004109	MAGNESIUM-ODAN	ODN	

MAGNESIUM GLUCONATE

29MG TABLET			
80062929	MMAGNESIUM GLUCONATE	MAN	
ST 500MG TABLET			
80009539	JAMP MAGNESIUM GLUCONATE	JMP	
00555126	MAGLUCATE	PED	

POTASSIUM CHLORIDE

ST 600MG CAPSULE			
80062704	JAMP POTASSIUM CHLORIDE ER	JMP	
02042304	MICRO K	PAL	
ST 1,500MG LIQUID			
80024835	JAMP-POTASSIUM CHLORIDE	JMP	
ST 1.33MEQ/ML SOLUTION			
02238604	PMS-POTASSIUM	PMS	

40:12.00 REPLACEMENT PREPARATIONS

POTASSIUM CHLORIDE

ST 8MMOL TABLET			
00602884	APO-K	APX	
02246734	EURO K	EUR	
80035346	MK 8	MAN	
02244068	RIVA-K 8	RIV	
ST 20MMOL TABLET			
80026265	BIO K-20 POTASSIUM	BMI	
02242261	EURO K	EUR	
80013007	JAMP K	JMP	
80004415	ODAN K20	ODN	
02243975	RIVA-K 20	RIV	
ST 780MG TABLET			
80025624	MK 20	MAN	
ST 8MMOL TABLET (EXTENDED RELEASE)			
80013005	JAMP-K 8	JMP	
ST 600MG TABLET (EXTENDED RELEASE)			
80008214	ODAN K8	ODN	
20MEQ TABLET (FILM COATED), EXTENDED RELEASE			
80071412	MK20 SOLUBLE	MAN	
ST 600MG TABLET (SUGAR COATED)			
80040226	SLOWK	NVR	
ST 780MG TABLET (TIME RELEASE)			
80040412	K20 POTASSIUM	UNK	
ST 1,500MG TABLET (TIME RELEASE)			
80040416	PHARMA-K20	PMS	
80053887	PRO-K 20	PDL	

POTASSIUM CITRATE

1080MG LIQUID			
80011529	POTASSIUM CITRATE	UNK	
10MEQ TABLET			
80023817	JAMPKCITRATE	JMP	
ST 10MMOL TABLET			
80026332	MK 10	MAN	
ST 25MEQ TABLET (EFFERVESCENT)			
80033602	JAMP-K EFFERVESCENT	JMP	
02085992	K LYTE	WPC	
ST 25MMOL TABLET (EFFERVESCENT)			
80011428	EURO K	EUR	

SODIUM CHLORIDE

1G CAPSULE			
90726364	SODIUM CHLORIDE 1G	MDS	
0.9% INJECTION			
99002329	SODIUM CHLORIDE (SMALL VOL.)	UNK	
0.9% SOLUTION			
00037818	BACTERIOSTATIC SODIUM CHLORIDE	PFI	
00037796	SODIUM CHLORIDE	PFI	
00060208	SODIUM CHLORIDE	BAX	
00402249	SODIUM CHLORIDE	OMG	
02150204	SODIUM CHLORIDE	OMG	
SYRINGE			
09991564	NACL SALINE PF	UNK	

**40:18.00 ION-REMOVING AGENTS
SODIUM POLYSTYRENE SULFONATE**

ORAL LIQUID

01902776 KAYEXALATE SAC

40:18.18 POTASSIUM - REMOVING AGENTS

CALCIUM POLYSTYRENE SULFONATE

1G POWDER FOR SOLUTION

02017741 RESONIUM CALCIUM SAC

SODIUM POLYSTYRENE SULFONATE

1G POWDER

02026961 KAYEXALATE SAC

00765252 K-EXIT OMG

00755338 SOLYSTAT PED

250MG/ML SUSPENSION

00769533 SOLYSTAT PED

00769541 SOLYSTAT PED

40:18.19 PHOSPHATE - REMOVING AGENTS

LANTHANUM CARBONATE HYDRATE

Limited use benefit (prior approval required).

For patients with elevated phosphate levels OR elevated phosphate X calcium product despite dietary restriction of phosphate and use of calcium-based phosphate binders (short term elevations should be managed with aluminium based binders).

For patients with elevated calcium levels despite discontinuation of calcium binder, and Vitamin D analogue and/or modification of dialysate calcium.

For patients with adynamic bone disease and low PTH levels (<100 pg/ml or <0.9 pmol/L) with normal or elevated calcium levels.

250MG TABLET (CHEWABLE)
02287145 FOSRENOL UNK

500MG TABLET (CHEWABLE)
02287153 FOSRENOL UNK

750MG TABLET (CHEWABLE)
02287161 FOSRENOL UNK

1000MG TABLET (CHEWABLE)
02287188 FOSRENOL UNK

40:18.19 PHOSPHATE - REMOVING AGENTS

SEVELAMER CARBONATE

Limited use benefit (prior approval required).

For patients with elevated phosphate levels OR elevated phosphate X calcium product despite dietary restriction of phosphate and use of calcium-based phosphate binders (short term elevations should be managed with aluminium based binders).

For patients with elevated calcium levels despite discontinuation of calcium binder, and Vitamin D analogue and/or modification of dialysate calcium.

For patients with adynamic bone disease and low PTH levels (<100 pg/ml or <0.9 pmol/L) with normal or elevated calcium levels.

800MG TABLET

02461501 ACCEL-SEVELAMER ACP

02354586 RENVELA SAC

SEVELAMER HYDROCHLORIDE

Limited use benefit (prior approval required).

For patients with elevated phosphate levels OR elevated phosphate X calcium product despite dietary restriction of phosphate and use of calcium-based phosphate binders (short term elevations should be managed with aluminium based binders).

For patients with elevated calcium levels despite discontinuation of calcium binder, and Vitamin D analogue and/or modification of dialysate calcium.

For patients with adynamic bone disease and low PTH levels (<100 pg/ml or <0.9 pmol/L) with normal or elevated calcium levels.

800MG TABLET

02244310 RENAGEL SAC

40:20.00 CALORIC AGENTS

LEVOCARNITINE

Limited use benefit (prior approval required).

For treatment of carnitine deficiency.

100MG/ML SOLUTION

02144336 CARNITOR UNK

200MG/ML SOLUTION

02144344 CARNITOR UNK

330MG TABLET

02144328 CARNITOR UNK

40:28.08 LOOP DIURETICS

ETHACRYNIC ACID

ST **25MG TABLET**

02258528 EDECRIN VAE

FUROSEMIDE

ST **10MG/ML SOLUTION**

02224720 LASIX SAC

ST **20MG TABLET**

00396788 APO FUROSEMIDE APX

02247371 BIO-FUROSEMIDE BMI

40:28.08 LOOP DIURETICS

FUROSEMIDE

ST **20MG TABLET**

00496723	FUROSEMIDE	PDL
02351420	FUROSEMIDE	SAN
02466759	MINT-FUROSEMIDE	MIN
02247493	PMS-FUROSEMIDE	PMS
00337730	TEVA-FUROSEMIDE	TEV

ST **40MG TABLET**

00362166	APO FUROSEMIDE	APX
02247372	BIO-FUROSEMIDE	BMI
00397792	FUROSEMIDE	PDL
02351439	FUROSEMIDE	SAN
02466767	MINT-FUROSEMIDE	MIN
02247494	PMS-FUROSEMIDE	PMS
00337749	TEVA-FUROSEMIDE	TEV

ST **80MG TABLET**

00707570	APO FUROSEMIDE	APX
00667080	FUROSEMIDE	PDL
02351447	FUROSEMIDE	SAN
02466775	MINT-FUROSEMIDE	MIN
00765953	TEVA-FUROSEMIDE	TEV

ST **500MG TABLET**

02224755	LASIX SPECIAL	SAC
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40:28.16 POTASSIUM SPARING DIURETICS

AMILORIDE

ST **5MG TABLET**

02249510	MIDAMOR	AAP
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AMILORIDE, HYDROCHLOROTHIAZIDE

ST **5MG & 50MG TABLET**

00784400	AA-AMILZIDE	APX
00870943	AMI-HYDRO	PDL
01937219	NOVAMILOR	TEV

TRIAMTERENE, HYDROCHLOROTHIAZIDE

ST **50MG & 25MG TABLET**

00441775	APO TRIAZIDE	APX
00532657	TEVA-TRIAMTERENE/HCTZ	TEV

40:28.20 TIAZIDE DIURETICS

HYDROCHLOROTHIAZIDE

ST **12.5MG TABLET**

02327856	APO-HYDRO	APX
02425947	MINT-HYDROCHLOROTHIAZIDE	MIN
02274086	PMS-HYDROCHLOROTHIAZIDE	PMS

ST **25MG TABLET**

00326844	APO HYDRO	APX
02247170	BIO-HYDROCHLOROTHIAZIDE	BMI
02360594	HYDROCHLOROTHIAZIDE	SAN
02426196	MINT-HYDROCHLOROTHIAZIDE	MIN
02247386	PMS-HYDROCHLOROTHIAZIDE	PMS
00021474	TEVA-HYDROCHLOROTHIAZIDE	TEV

ST **50MG TABLET**

00312800	APO HYDRO	APX
02247171	BIO-HYDROCHLOROTHIAZIDE	BMI
02360608	HYDROCHLOROTHIAZIDE	SAN

40:28.20 TIAZIDE DIURETICS

HYDROCHLOROTHIAZIDE

ST **50MG TABLET**

02247387	PMS-HYDROCHLOROTHIAZIDE	PMS
00021482	TEVA-HYDROCHLOROTHIAZIDE	TEV

ST **100MG TABLET**

00644552	APO HYDRO	APX
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ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503000	HYDROCHLOROTHIAZIDE ORAL LIQUID	UNK
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SPIRONOLACTONE, HYDROCHLOROTHIAZIDE

ST **25MG & 25MG TABLET**

00180408	ALDACTAZIDE	PFI
00613231	TEVA-SPIRONOLACTONE/HCTZ	TEV

ST **50MG & 50MG TABLET**

00594377	ALDACTAZIDE	PFI
00657182	TEVA-SPIRONOLACTONE/HCTZ	TEV

40:28.24 THIAZIDE LIKE DIURETICS

CHLORTHALIDONE

ST **50MG TABLET**

00360279	CHLORTHALIDONE	AAP
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ST **100MG TABLET**

00360287	APO CHLORTHALIDONE	APX
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INDAPAMIDE

ST **1.25MG TABLET**

02245246	APO-INDAPAMIDE	APX
02373904	JAMP-INDAPAMIDE	JMP
02179709	LOZIDE	SEV
02240067	MYLAN-INDAPAMIDE	MYL
02247245	RIVA-INDAPAMIDE	RIV

ST **2.5MG TABLET**

02223678	APO-INDAPAMIDE	APX
02373912	JAMP-INDAPAMIDE	JMP
00564966	LOZIDE	SEV
02153483	MYLAN-INDAPAMIDE	MYL
02312549	PRO-INDAPAMIDE	PDL
02242125	RIVA-INDAPAMIDE	RIV

METOLAZONE

ST **2.5MG TABLET**

00888400	ZAROXOLYN	SAC
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40:36.00 IRRIGATING SOLUTIONS

SODIUM CHLORIDE

0.9% SOLUTION

00801267	SODIUM CHLORIDE	UNK
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40:40.00 URICOSURIC AGENTS

SULFINPYRAZONE

200MG TABLET

00441767	SULFINPYRAZONE	AAP
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40:50.00 IRRIGATING SOLUTIONS

WATER

100% SOLUTION

00038202	BACTERIOSTATIC WATER	PFI
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**40:50.00 IRRIGATING SOLUTIONS
WATER**

100% SOLUTION

00402257	STERILE WATER	OMG
02142546	STERILE WATER	PFI

48:00 RESPIRATORY TRACT AGENTS

48:02.00 ANTIFIBROTIC AGENTS

NINTEDANIB ESILATE

Limited use benefit (prior approval required).

Initial Request - Coverage is provided for a period of 7 months (6 months plus a 4 week allowance for repeat pulmonary function tests):

For the treatment of adult patients with a diagnosis of mild to moderate idiopathic pulmonary fibrosis (IPF) who meet the following criteria:

- Diagnosis confirmed by a respirologist and a high-resolution CT scan within the previous 24 months; AND
- All other causes of restrictive lung disease (e.g. collagen vascular disorder or hypersensitivity pneumonitis) should be excluded; AND
- Mild to moderate IPF is defined as forced vital capacity (FVC) greater than or equal to 50% of predicted; AND
- Patient is under the care of a physician with experience in IPF.

Renewal at 6 months - Coverage is provided for a period of 6 months:

- Patients must NOT demonstrate progression of disease defined as an absolute decline in percent predicted 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 Renewals at 12 months and thereafter - Coverage is provided for a period of 12 months:

- 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.

Combination use of Ofev (nintedanib) and Esbriet (pirfenidone) will not be provided.

100MG CAPSULE

02443066 OFEV

BOE

150MG CAPSULE

02443074 OFEV

BOE

48:02.00 ANTIFIBROTIC AGENTS

PIRFENIDONE

Limited use benefit (prior approval required).

Initial Request - Coverage is provided for a period of 7 months (6 months plus a 4 weeks allowance for repeat pulmonary function tests):

For the treatment of adult patients with a diagnosis of mild to moderate idiopathic pulmonary fibrosis (IPF) who meet the following criteria:

- Diagnosis confirmed by a respirologist and a high-resolution CT scan within the previous 24 months; AND
- All other causes of restrictive lung disease (e.g. collagen vascular disorder or hypersensitivity pneumonitis) should be excluded; AND
- Mild to moderate IPF is defined as forced vital capacity (FVC) greater than or equal to 50% of predicted; AND
- Patient is under the care of a physician with experience in IPF.

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- Patients must NOT demonstrate progression of disease defined as an absolute decline in percent predicted 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.

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- 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.

Combination use of Ofev (nintedanib) and Esbriet (pirfenidone) will not be provided.

267MG CAPSULE

02393751 ESBRIET

HLR

267MG TABLET

02464489 ESBRIET

HLR

801MG TABLET

02464500 ESBRIET

HLR

48:10.24 LEUKOTRIENE MODIFIERS

MONTELUKAST SODIUM

Limited use benefit (prior approval required).

For treatment of:

- asthma when used in patients on concurrent steroid therapy; OR
- asthma patients not well controlled with or intolerant to inhaled corticosteroids.

Montelukast is open benefit for children up to 17 years of age.

ST **4MG GRANULES**

02358611 SANDOZ MONTELUKAST

SDZ

02247997 SINGULAIR

FRS

ST **10MG TABLET**

02374609 APO-MONTELUKAST

APX

02401274 AURO-MONTELUKAST

AUR

02445735 BIO-MONTELUKAST

UNK

48:10.24 LEUKOTRIENE MODIFIERS

MONTELUKAST SODIUM

Limited use benefit (prior approval required).

For treatment of:

- asthma when used in patients on concurrent steroid therapy; OR
- asthma patients not well controlled with or intolerant to inhaled corticosteroids.

Montelukast is open benefit for children up to 17 years of age.

ST **10MG TABLET**

02376695	DOM-MONTELUKAST	DPC
02391422	JAMP-MONTELUKAST	JMP
02399997	MAR-MONTELUKAST	MAR
02408643	MINT-MONTELUKAST	MIN
02379333	MONTELUKAST	SAN
02379856	MONTELUKAST	PDL
02382474	MONTELUKAST	SIV
02379236	MONTELUKAST SODIUM	ACC
02373947	PMS-MONTELUKAST	PMS
02389517	RAN-MONTELUKAST	RBV
02398826	RIVA-MONTELUKAST	RIV
02328593	SANDOZ MONTELUKAST	SDZ
02238217	SINGULAIR	FRS
02355523	TEVA-MONTELUKAST	TEV

4MG TABLET (CHEWABLE)

02377608	APO-MONTELUKAST	APX
02422867	AURO-MONTELUKAST	AUR
02442353	JAMP-MONTELUKAST	JMP
02399865	MAR-MONTELUKAST	MAR
02408627	MINT-MONTELUKAST	MIN
02379317	MONTELUKAST	SAN
02379821	MONTELUKAST	PDL
02382458	MONTELUKAST	SIV
02354977	PMS-MONTELUKAST	PMS
02402793	RAN-MONTELUKAST	RBV
02330385	SANDOZ MONTELUKAST	SDZ
02243602	SINGULAIR	FRS
02355507	TEVA-MONTELUKAST	TEV

ST **5MG TABLET (CHEWABLE)**

02377616	APO-MONTELUKAST	APX
02422875	AURO-MONTELUKAST	AUR
02442361	JAMP-MONTELUKAST	JMP
02399873	MAR-MONTELUKAST	MAR
02408635	MINT-MONTELUKAST	MIN
02379325	MONTELUKAST	SAN
02379848	MONTELUKAST	PDL
02382466	MONTELUKAST	SIV
02354985	PMS-MONTELUKAST	PMS
02402807	RAN-MONTELUKAST	RBV
02330393	SANDOZ MONTELUKAST	SDZ
02238216	SINGULAIR	FRS
02355515	TEVA-MONTELUKAST	TEV

48:10.32 MAST CELL STABILIZERS

CROMOLYN SODIUM

100MG CAPSULE

00500895	NALCROM	SAC
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48:10.32 MAST CELL STABILIZERS

CROMOLYN SODIUM

2% NASAL SPRAY

02231390	APO-CROMOLYN	APX
01950541	RHINARIS-CS	PED

10MG/ML SOLUTION

02046113	PMS-SODIUM CROMOGLYCATE	PMS
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48:48.00 VASODILATING AGENTS

AMBRISENTAN

Limited use benefit (prior approval required).

Maximum dose covered is 10 mg once daily.

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST **5MG TABLET**

02475375	APO-AMBRISENTAN	APX
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ST **10MG TABLET**

02475383	APO-AMBRISENTAN	APX
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BOSENTAN MONOHYDRATE

Limited use benefit (prior approval required).

Maximum dose covered is 125 mg twice daily

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST **125MG TABLET**

02399210	APO-BOSENTAN	APX
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RIOCIGUAT

Limited use benefit (prior approval required).

For the treatment of patients 18 years of age or older with chronic thromboembolic pulmonary hypertension (CTEPH) with World Health Organization (WHO) Functional Class 2 or 3 pulmonary hypertension with:

- Inoperable CTEPH, World Health Organization (WHO) Group 4;
- OR
- Persistent or recurrent CTEPH after surgical treatment; AND
- Prescriber experienced in the diagnosis and treatment of CTEPH.

0.5MG TABLET

02412764	ADEMPAS	BAY
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1MG TABLET

02412772	ADEMPAS	BAY
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1.5MG TABLET

02412799	ADEMPAS	BAY
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2MG TABLET

02412802	ADEMPAS	BAY
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2.5MG TABLET

02412810	ADEMPAS	BAY
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48:48.00 VASODILATING AGENTS

SELEXIPAG

Limited use benefit (prior approval required).

For the treatment of adult patients with World Health Organization (WHO) functional class (FC) II to III pulmonary arterial hypertension (PAH), including idiopathic PAH, heritable PAH, PAH associated with connective tissue disorders or PAH associated with congenital heart disease:

- Patient is under the care of a physician with experience in the diagnosis and treatment of PAH; AND
- Patient has failed to respond to first- and second-line PAH therapies; OR
- Patient has contraindications/intolerance to first- and second-line PAH therapies.

200MCG TABLET

02451158 UPTRAVI JSO

400MCG TABLET

02451166 UPTRAVI JSO

600MCG TABLET

02451174 UPTRAVI JSO

800MCG TABLET

02451182 UPTRAVI JSO

1000MCG TABLET

02451190 UPTRAVI JSO

1200MCG TABLET

02451204 UPTRAVI JSO

1400MCG TABLET

02451212 UPTRAVI JSO

1600MCG TABLET

02451220 UPTRAVI JSO

48:92.00 MISCELLANEOUS RESPIRATORY TRACT AGENTS

OMALIZUMAB

Limited use benefit (prior approval required).

Coverage is provided for an initial period of 24 weeks at a maximum dose of 300 mg every 4 weeks (6 injections over a 24 week period).

1. For the treatment of adults and adolescents (12 years of age or older) with moderate to severe chronic idiopathic urticaria (CIU) who remain symptomatic (presence of hives and/or associated itching) despite optimum management with H1 antihistamines; AND

Prescriber is experienced in the treatment of CIU (Allergist, Dermatologist, Immunologist, OR other authorized prescriber experienced in the treatment of CIU).

Treatment cessation could be considered for patients who experience complete symptom control (UAS-7 = 0) for at least 12 consecutive weeks at the end of a 24-week treatment period.

Renewal coverage is provided for 24 weeks at a maximum dose of 300 mg every 4 weeks (6 injections/24 weeks).

2. For the treatment of adults and adolescents (12 years of age or older) with moderate to severe chronic idiopathic urticaria (CIU); AND

Patient stopped omalizumab after achieving complete symptom control (UAS-7 = 0) for at least 12 weeks while on treatment, but has experienced symptom relapse; OR Patient achieved complete symptom control, but for a period of less than 12 consecutive weeks; OR Patient achieved a partial response to treatment, defined as a ≥ 9.5 -point reduction in baseline urticaria activity score over 7 days (UAS-7).

In patients where treatment is discontinued due to temporary symptom control, treatment re-initiation may be considered should CIU symptoms reappear.

150MG POWDER FOR SOLUTION

02260565 XOLAIR NVR

52:00 EYE, EAR, NOSE AND THROAT (EENT)

52:02.00 EENT - ANTIALLERGIC AGENTS

CROMOLYN SODIUM

2% OPHTHALMIC SOLUTION

02009277 CROMOLYN PED
02230621 OPTICROM ALL

KETOTIFEN FUMARATE

0.25MG SOLUTION

02400871 KETOTIFEN RAX

LEVOCABASTINE HYDROCHLORIDE

0.05% NASAL SPRAY

02020017 LIVOSTIN JSO

LODOXAMIDE TROMETHAMINE

0.1% SOLUTION

00893560 ALOMIDE NVR

OLOPATADINE HYDROCHLORIDE

0.1% OPHTHALMIC SOLUTION

02403986 ACT OLOPATADINE ACG
02305054 APO-OLOPATADINE APX
02422727 MINT-OLOPATADINE MIN
02233143 PATANOL NVR
02358913 SANDOZ OLOPATADINE SDZ

0.2% OPHTHALMIC SOLUTION

02404095 ACT OLOPATADINE ACG
02402823 APO-OLOPATADINE APX
02420171 SANDOZ OLOPATADINE SDZ

0.1% SOLUTION

02458411 JAMP-OLOPATADINE JMP

52:04.04 EENT - ANTIBACTERIALS

CIPROFLOXACIN HYDROCHLORIDE

0.3% OINTMENT

02200864 CILOXAN NVR

0.3% SOLUTION

02263130 APO-CIPROFLOX APX
01945270 CILOXAN NVR
02387131 SANDOZ CIPROFLOXACIN SDZ

CIPROFLOXACIN HYDROCHLORIDE, DEXAMETHASONE

0.3%/0.1% SUSPENSION

02252716 CIPRODEX NVR

ERYTHROMYCIN

5MG OINTMENT

00641324 ODAN-ERYTHROMYCIN ODN

5MG/G OINTMENT

02326663 ERYTHROMYCIN STG
01912755 PDP-ERYTHROMYCIN PED

FUSIDIC ACID

1% DROP

02243862 FUCITHALMIC AMD

52:04.04 EENT - ANTIBACTERIALS

GATIFLOXACIN

0.3% SOLUTION

02257270 ZYMAR ALL

MOXIFLOXACIN HYDROCHLORIDE (OPHTHALMIC)

0.5% SOLUTION

02404656 ACT MOXIFLOXACIN ACG
02406373 APO-MOXIFLOXACIN APX
02432218 PMS-MOXIFLOXACIN PMS
02411520 SANDOZ MOXIFLOXACIN SDZ
02252260 VIGAMOX NVR

OFLOXACIN

0.3% SOLUTION

02248398 APO-OFLOXACIN APX
02143291 OCUFLOX ALL
02247189 SANDOZ OFLOXACIN SDZ

POLYMYXIN B SULFATE, BACITRACIN ZINC

500IU & 10,000IU/G OINTMENT

02160889 OPTIMYXIN SDZ
02239157 POLYSPORIN JAJ

POLYMYXIN B SULFATE, GRAMICIDIN

0.025MG & 10,000U/ML DROP

00701785 OPTIMYXIN SDZ
02239156 POLYSPORIN EYE AND EAR JAJ

POLYMYXIN B SULFATE, TRIMETHOPRIM SULFATE

10,000U & 1MG/ML SOLUTION

02240363 PMS-POLYTRIMETHOPRIM PMS
02011956 POLYTRIM ALL
02239234 SANDOZ POLYTRIMETHOPRIM SDZ

TOBRAMYCIN (OPHTHALMIC)

0.3% OINTMENT

00614254 TOBREX NVR

0.3% SOLUTION

02241755 SANDOZ TOBRAMYCIN SDZ
00513962 TOBREX NVR

52:04.20 EENT - ANTIVIRALS

TRIFLURIDINE

1% SOLUTION

00687456 VIROPTIC VAE

52:04.92 EENT - MISCELLANEOUS ANTI-INFECTIVES

CHLORHEXIDINE GLUCONATE

0.12% MOUTHWASH

02462842 CHLORHEXIDINE EUR
02384272 GUM PAROEX SUS
02240433 PERICHLOR PED
02237452 PERIDEX MAK

52:08.00

FLUTICASONE PROPIONATE

50MCG SPRAY

02248307 FLONASE ALLERGY RELIEF GSK

52:08.08 EENT - CORTICOSTEROIDS

BECLOMETHASONE DIPROPIONATE

50MCG/DOSE NASAL SPRAY

02238796 APO-BECLOMETHASONE APX

02172712 MYLAN-BECLO AQ MYL

02228300 RIVANASE AQ RIV

BUDESONIDE

100MCG/DOSE POWDER

02035324 RHINOCORT TURBUHALER AZC

64MCG/DOSE SPRAY

02241003 MYLAN-BUDESONIDE AQ MYL

02231923 RHINOCORT AQUA MCL

100MCG/DOSE SPRAY

02230648 MYLAN-BUDESONIDE AQ MYL

DEXAMETHASONE

0.1% OINTMENT

00042579 MAXIDEX NVR

0.1% SUSPENSION

00042560 MAXIDEX NVR

DEXAMETHASONE PHOSPHATE

0.1% SOLUTION

02023865 DEXAMETHASONE UNK

00785261 PMS-DEXAMETHASONE PMS

DEXAMETHASONE, TOBRAMYCIN

0.1% & 0.3% OINTMENT

00778915 TOBRADEX NVR

0.1% & 0.3% SUSPENSION

00778907 TOBRADEX NVR

FLUMETHASONE PIVALATE, CLIOQUINOL

0.02% & 1% DROP

00074454 LOCACORTEN VIOFORM PAL

FLUOROMETHOLONE

0.1% DROP

00247855 FML ALL

0.1% SUSPENSION

00756784 FLAREX NVR

00432814 SANDOZ FLUOROMETHOLONE SDZ

FLUTICASONE FUROATE

100MCG POWDER

02446561 ARNUITY ELLIPTA GSK

200MCG POWDER

02446588 ARNUITY ELLIPTA GSK

FLUTICASONE PROPIONATE

50MCG PUMP

02453738 TEVA-FLUTICASONE TEV

50MCG/DOSE SPRAY

02294745 APO-FLUTICASONE APX

02296071 RATIO-FLUTICASONE TEV

52:08.08 EENT - CORTICOSTEROIDS

FRAMYCETIN SULFATE, GRAMICIDIN, DEXAMETHASONE

5MG & 0.05MG/ML & 0.5MG DROP

02224623 SOFRACORT EAR/EYE SAC

MOMETASONE FUROATE

50MCG SPRAY

02403587 APO-MOMETASONE APX

02238465 NASONEX FRS

02475863 TEVA-MOMETASONE TEV

500MCG/ML SPRAY

02449811 SANDOZ MOMETASONE SDZ

PREDNISOLONE ACETATE

0.12% DROP

00299405 PRED MILD ALL

1% DROP

00301175 PRED FORTE ALL

1% SUSPENSION

01916203 SANDOZ PREDNISOLONE SDZ

00700401 TEVA-PREDNISOLONE TEV

PREDNISOLONE ACETATE, SULFACETAMIDE SODIUM

0.2% & 10% DROP

00807788 BLEPHAMIDE ALL

0.2% & 10% OINTMENT

00307246 BLEPHAMIDE ALL

0.5% & 10% SUSPENSION

02023814 PREDNISOLONE/SULFACETAMIDE UNK

PREDNISOLONE SODIUM PHOSPHATE

0.5% DROP

02148498 MINIMS PREDNISOLONE VAE

TRIAMCINOLONE ACETONIDE

55MCG SPRAY

02437635 APO-TRIAMCINOLONE AQ APX

55MCG/DOSE SPRAY

02213834 NASACORT AQ SAC

52:08.20 EENT - NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

DICLOFENAC SODIUM

0.1% SOLUTION

01940414 VOLTAREN OPHTHA NVR

DICLOFENAC SODIUM (TOPICAL)

Limited use benefit (prior approval required).

For the treatment of osteoarthritis when:

- pain is inadequately controlled with acetaminophen AND a non-steroidal anti-inflammatory (NSAID); OR
- there is contraindication to acetaminophen and NSAID; OR
- there is intolerance to acetaminophen and NSAID.

0.1% SOLUTION

02441020 APO-DICLOFENAC APX

02454807 SANDOZ DICLOFENAC OPHTHA SDZ

52:08.20 EENT - NONSTEROIDAL ANTI-INFLAMMATORY AGENTS

KETOROLAC TROMETHAMINE

0.45% SOLUTION

02369362 ACUVAIL ALL

0.5% SOLUTION

01968300 ACULAR ALL

02245821 APO-KETOROLAC AAP

NEPAFENAC

0.1% SUSPENSION

02308983 NEVANAC NVR

0.3% SUSPENSION

02411393 ILEVRO NVR

52:12.00 EENT - CONTACT LENS SOLUTION

HYDROXYPROPYLMETHYLCELLULOSE

3MG SOLUTION

02231289 GENTEAL NVR

52:16.00 EENT - LOCAL ANESTHETICS

LIDOCAINE HYDROCHLORIDE

2% SOLUTION

00001686 XYLOCAINE VISCOUS UNK

52:24.00 EENT - MYDRIATICS

ATROPINE SULFATE

1% SOLUTION

02023695 ATROPINE UNK

00035017 ISOPTO ATROPINE ALC

02148358 MINIMS ATROPINE VAE

CYCLOPENTOLATE HYDROCHLORIDE

0.5% DROP

02148331 MINIMS CYCLOPENTOLATE VAE

1% DROP

00252506 CYCLOGYL ALC

02023644 CYCLOPENTOLATE UNK

02148382 MINIMS CYCLOPENTOLATE VAE

DIPIVEFRIN HYDROCHLORIDE

0.1% LIQUID

02242232 APO-DIPIVEFRIN APX

PHENYLEPHRINE HYDROCHLORIDE

2.5% DROP

02148447 MINIMS PHENYLEPHRINE VAE

00465763 MYDFRIN ALC

02027100 PHENYLEPHRINE UNK

10% DROP

02148455 MINIMS PHENYLEPHRINE VAE

TROPICAMIDE

0.5% SOLUTION

00000981 MYDRIACYL ALC

1% SOLUTION

00001007 MYDRIACYL ALC

52:28.00 EENT - MOUTHWASHES AND GARGLES

BENZYDAMINE HYDROCHLORIDE

Limited use benefit (prior approval required).

• For the treatment of radiation mucositis and oral ulcerative complications of chemotherapy.

• For use in immunocompromised patients who are at risk of mucosal breakdown.

0.15% MOUTHWASH

02239044 APO-BENZYDAMINE APX

02229777 PHARIXIA PED

02239537 PMS-BENZYDAMINE PMS

52:32.00 EENT - VASOCONSTRICTORS

EPINEPHRINE

1MG/ML SOLUTION

00155365 ADRENALIN ERF

NAPHAZOLINE HYDROCHLORIDE

0.1% DROP

00001147 ALBALON ALL

52:40.04 EENT - ALPHA-ADRENERGIC AGONISTS

BRIMONIDINE TARTRATE

0.15% SOLUTION

02248151 ALPHAGAN P ALL

02301334 BRIMONIDINE P AAP

0.2% SOLUTION

02236876 ALPHAGAN ALL

02260077 APO-BRIMONIDINE APX

02246284 PMS-BRIMONIDINE PMS

02305429 SANDOZ BRIMONIDINE SDZ

TIMOLOL MALEATE, BRIMONIDINE TARTRATE

0.2% & 0.5% SOLUTION

02248347 COMBIGAN ALL

52:40.08 EENT - BETA-ADRENERGIC BLOCKING AGENTS

BETAXOLOL HYDROCHLORIDE

0.25% OPHTHALMIC SOLUTION

01908448 BETOPTIC S NVR

LEVOBUNOLOL HYDROCHLORIDE

0.25% OPHTHALMIC SOLUTION

02241575 APO-LEVOBUNOLOL APX

0.5% OPHTHALMIC SOLUTION

00637661 BETAGAN ALL

TIMOLOL MALEATE

0.25% OPHTHALMIC GEL SOLUTION

02242275 TIMOLOL MALEATE-EX SDZ

0.5% OPHTHALMIC GEL SOLUTION

02290812 APO-TIMOP APX

02242276 TIMOLOL MALEATE-EX SDZ

00451207 TIMOPTIC PFR

0.25% OPHTHALMIC SOLUTION

00755826 APO-TIMOP APX

52:40.08 EENT - BETA-ADRENERGIC BLOCKING AGENTS

TIMOLOL MALEATE

0.25% OPHTHALMIC SOLUTION

02238770 DOM-TIMOLOL DPC
02083353 PMS-TIMOLOL PMS

0.5% OPHTHALMIC SOLUTION

00755834 APO-TIMOP APX
02238771 DOM-TIMOLOL DPC
02447800 JAMP-TIMOLOL JMP
02083345 PMS-TIMOLOL PMS
02166720 SANDOZ TIMOLOL SDZ

0.25% SOLUTION (EXTENDED RELEASE)

02171880 TIMOPTIC-XE PFR

0.5% SOLUTION (EXTENDED RELEASE)

02171899 TIMOPTIC-XE PFR

52:40.12 EENT - CARBONIC ANHYDRASE INHIBITORS

ACETAZOLAMIDE

250MG TABLET

00545015 ACETAZOLAMIDE AAP

BRINZOLAMIDE

1% SUSPENSION

02238873 AZOPT NVR

BRINZOLAMIDE, BRIMONIDINE TARTRATE

1% & 0.2% SUSPENSION

02435411 SIMBRINZA NVR

BRINZOLAMIDE, TIMOLOL MALEATE

1%/0.5% SUSPENSION

02331624 AZARGA NVR

DORZOLAMIDE HYDROCHLORIDE

2% OPHTHALMIC SOLUTION

02216205 TRUSOPT FRS
02269090 TRUSOPT FRS

20MG/ML OPHTHALMIC SOLUTION

02316307 SANDOZ DORZOLAMIDE SDZ

DORZOLAMIDE HYDROCHLORIDE, TIMOLOL MALEATE

20MG & 5MG OPHTHALMIC SOLUTION

02437686 MED-DORZOLAMIDE-TIMOLOL GMP

20MG & 5MG/ML OPHTHALMIC SOLUTION

02404389 ACT DORZOTIMOLOL TEV
02299615 APO-DORZO-TIMOP APX
02240113 COSOPT FRS
02442426 PMS-DORZOLAMIDE-TIMOLOL PMS
02441659 RIVA-DORZOLAMIDE/TIMOLOL RIV
02344351 SANDOZ DORZOLAMIDE/TIMOLOL SDZ

200MG & 5MG OPHTHALMIC SOLUTION

02443090 MINT-DORZOLAMIDE/TIMOLOL MIN

METHAZOLAMIDE

50MG TABLET

02245882 METHAZOLAMIDE AAP

52:40.20 EENT - MIOTICS CARBACHOL

0.01% OPHTHALMIC SOLUTION

00042544 MIOSTAT ALC

PILOCARPINE HYDROCHLORIDE

2% OPHTHALMIC SOLUTION

00000868 ISOPTO CARPINE NVR

4% OPHTHALMIC SOLUTION

00000884 ISOPTO CARPINE NVR
02023733 PILOCARPINE UNK

PILOCARPINE NITRATE

2% DROP

02148463 MINIMS PILOCARPINE VAE

52:40.28 EENT - PROSTAGLANDIN AGENTS

BIMATOPROST

0.01% OPHTHALMIC SOLUTION

02324997 LUMIGAN RC ALL
09857368 LUMIGAN RC (ON) ALL
09857398 LUMIGAN RC (ON) ALL

0.03% OPHTHALMIC SOLUTION

02429063 VISTITAN SDZ

LATANOPROST

0.005% SOLUTION

02296527 APO-LATANOPROST APX
02373041 GD-LATANOPROST PFI
02426935 MED-LATANOPROST GMP
02317125 PMS-LATANOPROST PMS
02341085 RIVA-LATANOPROST RIV
02367335 SANDOZ LATANOPROST SDZ
02254786 TEVA-LATANOPROST TEV
02231493 XALATAN PFI

LATANOPROST, TIMOLOL MALEATE

0.005% & 0.5% SOLUTION

02436256 ACT LATANOPROST/TIMOLOL ACG
02414155 APO-LATANOPROST-TIMOP APX
02373068 GD-LATANOPROST/TIMOLOL PFI
02404591 PMS-LATANOPROST-TIMOLOL PMS
02394685 SANDOZ LATANOPROST/TIMOLOL SDZ
02246619 XALACOM PFI

TIMOLOL MALEATE, TRAVOPROST

0.5% & 0.004% SOLUTION

02415305 APO-TRAVOPROST-TIMOP APX
02278251 DUOTRAV PQ NVR
02413817 SANDOZ TRAVOPROST / TIMOLOL SDZ
PQ

TRAVOPROST

0.004% SOLUTION

02415739 APO-TRAVOPROST Z APX
02413167 SANDOZ TRAVOPROST SDZ
02412063 TEVA-TRAVOPROST Z TEV
02318008 TRAVATAN Z NVR

52:92.00 MISCELLANEOUS EENT DRUGS

AFLIBERCEPT

Limited use benefit (prior approval required).

For the treatment of:
Diabetic Macular Edema (DME)
Wet Age-Related Macular Degeneration (w-AMD)
Retinal Vein Occlusion (RVO)

(Please refer to Appendix A).

40MG SOLUTION

02415992 EYLEA BAY

ANETHOLE TRITHIONE

ST **25MG TABLET**

02240344 SIALOR PMS

APRACLONIDINE HYDROCHLORIDE

0.5% OPHTHALMIC SOLUTION

02076306 IOPIDINE NVR

**DEXTRAN 70,
HYDROXYPROPYLMETHYLCELLULOSE**

0.1% & 0.3% DROP

01943308 TEARS NATURALE FREE NVR

00743445 TEARS NATURALE II NVR

HYDROXYPROPYL CELLULOSE

5MG INSERT

02250624 LACRISERT ATO

HYDROXYPROPYLMETHYLCELLULOSE

0.5% DROP

00000809 ISOPTO TEARS NVR

1% DROP

00000817 ISOPTO TEARS NVR

MACROGOL, PROPYLENE GLYCOL

15% & 20% GEL

02220806 LUBRICATING PMS

02352699 RHINARIS NASAL PED

00551805 SECARIS PED

15% & 20% SPRAY

00732230 LUBRICATING NASAL MIST PMS

02354551 RHINARIS NASAL MIST PED

MINERAL OIL, WHITE PETROLATUM

55.5% & 42.5% OINTMENT

00210889 REFRESH LACRI-LUBE ALL

PETROLATUM, MINERAL OIL

80% & 20% OINTMENT

02125706 SOOTHE NIGHT TIME BSH

POLYVINYL ALCOHOL

1.4% OPHTHALMIC SOLUTION

02229570 ARTIFICIAL TEARS PED

00579408 TEARS PLUS ALL

52:92.00 MISCELLANEOUS EENT DRUGS

RANIBIZUMAB

Limited use benefit (prior approval required).

For the treatment of:
Diabetic Macular Edema (DME)
Wet Age-Related Macular Degeneration (w-AMD)
Retinal Vein Occlusion (RVO)
Choroidal Neovascularization secondary to pathologic myopia (mCNV)

(Please refer to Appendix A).

10MG/ML SOLUTION

02296810 LUCENTIS NVR

02425629 LUCENTIS PFS NVR

SODIUM CARBOXYMETHYL CELLULOSE

0.5% DROP

02049260 REFRESH PLUS ALL

02231008 REFRESH TEARS ALL

1% DROP

00870153 REFRESH CELLUVISC ALL

10MG/ML SOLUTION

02244650 REFRESH LIQUIGEL ALL

SODIUM CHLORIDE

9MG/ML NASAL DROPS

80024901 SALINEX SDZ

5% OINTMENT

00750816 MURO 128 BSH

5% OPHTHALMIC OINTMENT

80046696 ODAN SODIUM CHLORIDE ODN

5% SOLUTION

00750824 MURO 128 BSH

80046737 ODAN-SODIUM CHLORIDE ODN

9MG/ML SPRAY

80024381 SALINEX SDZ

VERTEPORFIN

Limited use benefit (prior approval required).

For treatment of age related macular degeneration for patients with this diagnosis who are being treated by a certified ophthalmologist.

15MG/VIAL POWDER FOR SOLUTION

02242367 VISUDYNE VAE

WHITE PETROLATUM, LANOLIN, MINERAL OIL

94% & 3% & 3% OINTMENT

02444062 SYSTANE NVR

56:00 GASTROINTESTINAL DRUGS

56:04.00 ANTACIDS AND ADSORBENTS

BISMUTH SUBSALICYLATE

Limited use benefit (prior approval not required).

Coverage will be limited to 8 tablets a day every 14 days, followed by a 28 day lockout;
OR

Coverage will be limited to 120mL a day every 14 days, followed by a 28 day lockout.

262MG CAPLET

00245730 BISMUTH JMP

17.6MG/ML SUSPENSION

02097079 PEPTO BISMOL PGI

262MG TABLET

02326582 BISMUTH SUBSALICYLATE UNK

02177994 PEPTO BISMOL PGI

MAGNESIUM OXIDE

420MG TABLET

00299448 MAGNESIUM OXIDE VAE

835MG TABLET

00689785 HI POTENCY MAGNESIUM OXIDE SWS

SODIUM BICARBONATE

325MG TABLET

80072247 SODIUM BICARBONATE MDS

56:08.00 ANTIDIARRHEA AGENTS

LOPERAMIDE HYDROCHLORIDE

ST **0.2MG/ML SOLUTION**

02192667 DIARR-EZE PMS

02016095 PMS-LOPERAMIDE PMS

ST **2MG/15ML SOLUTION**

02291800 IMODIUM CALMING MCL

ST **2MG TABLET**

02212005 APO-LOPERAMIDE APX

02229552 DIARR-EZE PMS

02248994 DIARRHEA RELIEF PMS

02256452 DIARRHEA RELIEF VTH

02239535 DOM-LOPERAMIDE DPC

02225182 LOPERAMIDE PDL

02228351 PMS-LOPERAMIDE PMS

02238211 RIVA-LOPERAMIDE RIV

02132591 TEVA-LOPERAMIDE TEV

56:12.00 CATHARTICS AND LAXATIVES

BISACODYL

5MG SUPPOSITORY

02410893 BISACODYL JMP

10MG SUPPOSITORY

02361450 BISACODYL JMP

00003875 DULCOLAX BOE

00582883 PMS-BISACODYL PMS

02241091 THE MAGIC BULLET DCM

ST **5MG TABLET**

00254142 DULCOLAX BOE

02246039 JAMP-BISACODYL JMP

00587273 PMS-BISACODYL PMS

56:12.00 CATHARTICS AND LAXATIVES

BISACODYL

ST **5MG TABLET (DELAYED RELEASE)**

00545023 APO-BISACODYL APX

02273411 BISACODYL-ODAN ODN

CITRIC ACID, MAGNESIUM OXIDE, SODIUM PICOSULFATE

ST **12G & 3.5G & 10MG POWDER FOR SOLUTION**

02254794 PICO-SALAX FEI

02317966 PURG-ODAN ODN

GLYCERINE

ADULT SUPPOSITORY

00873462 GLYCERIN TEV

01926039 GLYCERIN WPC

02020394 GLYCERIN TEV

80029765 JAMP GLYCERIN JMP

PEDIATRIC SUPPOSITORY

02020815 GLYCERIN TEV

01926047 GLYCERIN FOR INFANTS WPC

CHILDREN

MACROGOL, POTASSIUM CHLORIDE, SODIUM BICARBONATE, SODIUM CHLORIDE, SODIUM SULFATE

ST **60G & 750MG & 1.68G & 1.46G & 5.68G/L SOLUTION**

00652512 GOLYTELY BTU

00777838 PEGLYTE PED

MAGNESIUM CITRATE

ST **5.40% SOLUTION**

00262609 CITRO MAG TEV

ST **50MG/ML SOLUTION**

80001809 CITRODAN ODN

MAGNESIUM HYDROXIDE

ST **80MG/ML LIQUID**

02245289 MILK OF MAGNESIA PMS

02150646 PHILLIPS MILK OF MAGNESIA BAY

ST **311MG TABLET (CHEWABLE)**

02150638 PHILIPS MAGNESIA BAY

MINERAL OIL

ST **78% GEL**

00608734 LANSOYL AUP

02186926 LANSOYL SUGAR FREE AUP

ST **100% LIQUID**

01935348 MINERAL OIL (HEAVY) RBW

POLYETHYLENE GLYCOL 3350

POWDER

09991007 POLYETHYLENE GLYCOL MDS

09991054 POLYETHYLENE GLYCOL 3350 MDS

ST **100% POWDER FOR SOLUTION**

02324989 CLEARLAX PER

02374137 EMOLAX JMP

02450070 M-PEG 3350 MAN

ST **1G POWDER FOR SOLUTION**

02317680 LAX-A-DAY PED

56:12.00 CATHARTICS AND LAXATIVES

POLYETHYLENE GLYCOL 3350

ST 1G POWDER FOR SOLUTION

02453193	LAX-A-DAY PHARMA	PMS
02358034	PEG 3350	MDS
02346672	RELAXA	RLI
02318164	RESTORALAX	BAY

POLYETHYLENE GLYCOL 3350, SODIUM SULFATE, SODIUM BICARBONATE, SODIUM CHLORIDE, POTASSIUM CHLORIDE

ST 60G & 750MG & 1.68G & 1.46G & 5.68G/L POWDER

00677442	COLYTE	PED
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POLYETHYLENE GLYCOL 3350, SODIUM SULFATE, SODIUM BICARBONATE, SODIUM CHLORIDE, POTASSIUM CHLORIDE, BISACODYL

ST 59.55G & 5.74G & 1.69G & 1.46G & 0.76G & 5MG LIQUID

02326302	BI-PEGLYTE	PED
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PSYLLIUM MUCILLOID

ST 50% POWDER

00599875	MUCILLIUM	PMS
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ST 680MG/G POWDER

02174812	METAMUCIL FIBRE THERAPY ORIGINAL TEXTURE UNFLAVOURED	PGI
02174790	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE ORANGE FLAVOUR	PGI
02174782	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE ORANGE FLAVOUR (SUGAR-FREE)	PGI
02174804	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE UNFLAVOURED	PGI

SENNOSIDES

ST 1.7MG/ML LIQUID

80024394	JAMP SENNAQUIL	JMP
02144379	SENNALAX	PMS
02084651	SENNAPREP	PMS
00367729	SEKOKOT	PFR

ST 8.6MG TABLET

80043280	M SENNOSIDES	MAN
80047592	OPUS SENNOSIDES	OPU
01949292	RIVA SENNA	RIV

ST 9MG TABLET

80019511	BIOSENNOSIDES	BMI
02247389	EURO SENNA	EUR
80054498	M SENNOSIDES	MAN
00896411	PMS-SENNOSIDES	PMS
80009595	SENNA	JMP
02237105	SENNA LAXATIVE	VTH
02068109	SENNA SENNOSIDES	PMS
80009182	SENNOSIDES	JMP
00026158	SEKOKOT	PFR

ST 12MG TABLET

80055641	M-SENNOSIDES	MAN
00896403	PMS-SENNOSIDES	PMS

56:12.00 CATHARTICS AND LAXATIVES

SENNOSIDES

ST 12MG TABLET

80009183	SENNOSIDES	JMP
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ST 15MG TABLET

02226030	EXLAX CHOCOLATED	NVC
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43MG TABLET

80061813	SENNACE	VAN
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8.6MG TABLET (FILM COATED)

80064362	SENNA SENNOSIDES NATURALS	UNK
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15MG TABLET (FILM COATED)

80054167	SENNOSIDES	UNK
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SODIUM PHOSPHATE

ST 0.9G ORAL SOLUTION

80000689	PHOSLAX	ODN
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ST 60MG & 160MG/ML RECTAL LIQUID

02096900	ENEMOL SODIUM PHOSPHATE	DPC
00009911	FLEET ENEMA	KIM
00108065	FLEET ENEMA PEDIATRIC	KIM

ST 180MG & 480MG/ML SOLUTION

02230399	PHOSPHATES	PMS
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ST 2.4G SOLUTION

80034416	JAMP-SODIUM PHOSPHATE	JMP
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123MG TABLET (EFFERVESCENT)

80047562	JAMP-SODIUM PHOSPHATE	JMP
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SORBITOL, SODIUM CITRATE, SODIUM LAURYL SULFOACETATE

ST 90MG & 9MG & 625MG ENEMA

02063905	MICROLAX	MCL
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56:14.00 CHOLELITHOLYTIC AGENTS

URSODIOL

ST 250MG TABLET

02472392	JAMP-URSODIOL	JMP
02273497	PMS-URSODIOL	PMS
02238984	URSO	APC
02426900	URSODIOL	GLK

ST 500MG TABLET

02472406	JAMP-URSODIOL	JMP
02273500	PMS-URSODIOL	PMS
02245894	URSO DS	APC
02426919	URSODIOL	GLK

ST PDIN FOR EXTEMPORANEOUS MIXTURE

99503024	UROSODIOL ORAL LIQUID	UNK
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56:16.00 DIGESTANTS

LACTASE

ST 3,000U CAPLET

02239139	DAIRY DIGESTIVE	VTH
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ST 4,500U CAPLET

02239140	DAIRY DIGESTIVE	VTH
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ST ORAL LIQUID

99100157	LACTEEZE DROPS	AUP
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ST 300MG TABLET

80070358	JAMPLACTASE ENZYME	JMP
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56:16.00 DIGESTANTS

LACTASE

ST **3,000U TABLET**

01951637	DAIRY AID	TAN
02230653	LACTAID	KIM
02017512	LACTOMAX	STE

ST **4,500U TABLET**

02230654	LACTAID EXTRA STRENGTH	KIM
02224909	LACTOMAX EXTRA	STE

ST **9,000U TABLET**

02231507	LACTAID ULTRA	KIM
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LIPASE, AMYLASE, PROTEASE

ST **8,000U & 30,000U & 30,000U CAPSULE**

00263818	COTAZYM	FRS
00502790	COTAZYM ECS 8	FRS

ST **20,000U & 55,000U & 55,000U CAPSULE**

00821373	COTAZYM ECS 20	FRS
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ST **10000U & 11200U & 730U CAPSULE (DELAYED RELEASE)**

02200104	CREON MINIMICROSPHERES 10	ABB
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ST **25000U & 25500U & 1600U CAPSULE (DELAYED RELEASE)**

01985205	CREON MINIMICROSPHERES 25	ABB
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ST **5000U & 5100U & 320U GRANULES FOR SUSPENSION, DELAYED RELEASE**

02445158	CREON MINIMICROSPHERES MICRO	BGP
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56:20.00 EMETICS

IPECAC

14MG/ML LIQUID

00378801	XENEX IPECAC	XEN
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56:22.08 ANTIHISTAMINES

DIMENHYDRINATE

Limited use benefit (prior approval not required).

The NIHB Program implemented a dose coverage limit for DIMENHYDRINATE in June 2017 as part of a strategy to address safety concerns and potential misuse.

The dimenhydrinate dose limit is currently 400 mg per day for a total of 12,000 mg of dimenhydrinate in a 30-day period.

This limit applies only to the 15 mg and 50 mg tablets. Dimenhydrinate in liquid, suppository and injectable forms are not included in this limit.

50MG/ML INJECTION

00392537	DIMENHYDRINATE	SDZ
00013579	GRAVOL	CHU

10MG LIQUID

00392731	DIMENHYDRINATE	SDZ
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25MG SUPPOSITORY

00783595	GRAVOL	CHU
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50MG SUPPOSITORY

00392553	SANDOZ DIMENHYDRINATE	SDZ
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100MG SUPPOSITORY

00013609	GRAVOL	CHU
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56:22.08 ANTIHISTAMINES

DIMENHYDRINATE

Limited use benefit (prior approval not required).

The NIHB Program implemented a dose coverage limit for DIMENHYDRINATE in June 2017 as part of a strategy to address safety concerns and potential misuse.

The dimenhydrinate dose limit is currently 400 mg per day for a total of 12,000 mg of dimenhydrinate in a 30-day period.

This limit applies only to the 15 mg and 50 mg tablets. Dimenhydrinate in liquid, suppository and injectable forms are not included in this limit.

ST **3MG/ML SYRUP**

00230197	GRAVOL	CHU
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50MG TABLET

02241532	ANTI-NAUSEANT	VTH
00363766	APO DIMENHYDRINATE	APX
00013803	GRAVOL	CHU
02245416	JAMP-DIMENHYDRINATE	JMP
02377179	MOTION SICKNESS	APX
00586331	PMS-DIMENHYDRINATE	PMS
00605786	TRAVEL	VTH
00021423	TRAVEL ON	NOP

DOXYLAMINE SUCCINATE, PYRIDOXINE HYDROCHLORIDE

ST **10MG & 10MG TABLET (DELAYED RELEASE)**

00609129	DICLECTIN	DUI
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56:22.20 5-HT3 RECEPTOR ANTAGONISTS

GRANISETRON HYDROCHLORIDE

ST **1MG TABLET**

02308894	APO-GRANISETRON	APX
02452359	NAT-GRANISETRON	NPH

ONDANSETRON HYDROCHLORIDE

ST **4MG FILM**

02389983	ONDISSOLVE ODF	TAK
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ST **8MG FILM**

02389991	ONDISSOLVE ODF	TAK
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ST **0.8MG/ML SOLUTION**

02291967	ONDANSETRON	AAP
02229639	ZOFAN	NVR

ST **4MG TABLET**

02296349	ACT ONDANSETRON	ACG
02288184	APO-ONDANSETRON	APX
02313685	JAMP-ONDANSETRON	JMP
02371731	MAR-ONDANSETRON	MAR
02305259	MINT-ONDANSETRON	MIN
02297868	MYLAN-ONDANSETRON	MYL
02417839	NAT-ONDANSETRON	NPH
02421402	ONDANSETRON	SAN
02258188	PMS-ONDANSETRON	PMS
02312247	RAN-ONDANSETRON	RBV
02274310	SANDOZ ONDANSETRON	SDZ
02376091	SEPTA-ONDANSETRON	SPT
02448440	VAN-ONDANSETRON	VAN
02213567	ZOFAN	NVR

56:22.20 5-HT3 RECEPTOR ANTAGONISTS

ONDANSETRON HYDROCHLORIDE

ST 8MG TABLET

02296357	ACT ONDANSETRON	ACG
02288192	APO-ONDANSETRON	APX
02313693	JAMP-ONDANSETRON	JMP
02371758	MAR-ONDANSETRON	MAR
02305267	MINT-ONDANSETRON	MIN
02297876	MYLAN-ONDANSETRON	MYL
02417847	NAT-ONDANSETRON	NPH
02325160	ONDANSETRON	PDL
02421410	ONDANSETRON	SAN
02258196	PMS-ONDANSETRON	PMS
02312255	RAN-ONDANSETRON	RBY
02274329	SANDOZ ONDANSETRON	SDZ
02376105	SEPTA-ONDANSETRON	SPT
02448467	VAN-ONDANSETRON	VAN
02213575	ZOFRAN	NVR

ST 4MG TABLET (ORALLY DISINTEGRATING)

02444674	VPI-ONDANSETRON ODT	UNK
02239372	ZOFRAN ODT	NVR

ST 8MG TABLET (ORALLY DISINTEGRATING)

02444682	VPI-ONDANSETRON ODT	UNK
02239373	ZOFRAN ODT	NVR

56:22.32 MISCELLANEOUS ANTIEMETICS

APREPITANT

Limited use benefit (prior approval required).

When used in combination with a 5-HT3 antagonist and dexamethasone for the prevention of acute and delayed nausea and vomiting due to highly emetogenic cancer chemotherapy (e.g. Cisplatin > 70mg/m2).

ST 80MG CAPSULE

02298791	EMEND	FRS
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ST 125MG CAPSULE

02298805	EMEND	FRS
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ST 125MG & 80MG CAPSULE

02298813	EMEND TRI-PACK	FRS
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56:22.92 MISCELLANEOUS ANTIEMETICS

NABILONE

Limited use benefit (prior approval required).

For patients who are experiencing nausea and vomiting due to cancer chemotherapy or radiation;
OR

Patient is palliative (diagnosed with a terminal illness or disease which is expected to be the primary cause of death within six months or less).

0.25MG CAPSULE

02312263	CESAMET	VAE
02358077	RAN-NABILONE	RBY
02392925	TEVA-NABILONE	TEV

0.5MG CAPSULE

02393581	ACT NABILONE	ACG
02256193	CESAMET	VAE
02380900	PMS-NABILONE	PMS
02358085	RAN-NABILONE	RBY

56:22.92 MISCELLANEOUS ANTIEMETICS

NABILONE

Limited use benefit (prior approval required).

For patients who are experiencing nausea and vomiting due to cancer chemotherapy or radiation;
OR

Patient is palliative (diagnosed with a terminal illness or disease which is expected to be the primary cause of death within six months or less).

0.5MG CAPSULE

02384884	TEVA-NABILONE	TEV
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1MG CAPSULE

02393603	ACT NABILONE	ACG
00548375	CESAMET	VAE
02380919	PMS-NABILONE	PMS
02358093	RAN-NABILONE	RBY
02384892	TEVA-NABILONE	TEV

56:28.12 HISTAMINE H2-ANTAGONISTS

CIMETIDINE

ST 200MG TABLET

00584215	CIMETIDINE	APX
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ST 300MG TABLET

00487872	CIMETIDINE	APX
02227444	MYLAN-CIMETIDINE	MYL

ST 400MG TABLET

00600059	CIMETIDINE	APX
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ST 600MG TABLET

00600067	CIMETIDINE	APX
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ST 800MG TABLET

00749494	CIMETIDINE	APX
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FAMOTIDINE

ST 20MG TABLET

01953842	APO-FAMOTIDINE	APX
02351102	FAMOTIDINE	SAN
02022133	TEVA-FAMOTIDINE	TEV

ST 40MG TABLET

01953834	APO-FAMOTIDINE	APX
02351110	FAMOTIDINE	SAN
02022141	TEVA-FAMOTIDINE	TEV

NIZATIDINE

ST 150MG CAPSULE

00778338	AXID	PED
02177714	PMS-NIZATIDINE	PMS

ST 300MG CAPSULE

00778346	AXID	PED
02177722	PMS-NIZATIDINE	PMS

RANITIDINE HYDROCHLORIDE

ST 15MG/ML SOLUTION

02280833	APO-RANITIDINE	APX
02242940	TEVA-RANITIDINE	TEV

ST 150MG TABLET

02248570	ACT RANITIDINE	TEV
00733059	APO-RANITIDINE	APX
02463717	JAMP-RANITIDINE	JMP
02443708	MAR-RANITIDINE	MAR

56:28.12 HISTAMINE H2-ANTAGONISTS

RANITIDINE HYDROCHLORIDE

ST **150MG TABLET**

02293471	MAXIMUM STRENGTH ACID REDUCER	PMS
02473534	M-RANITIDINE	MAN
02242453	PMS-RANITIDINE	PMS
00740748	RANITIDINE	PDL
02353016	RANITIDINE	SAN
02385953	RANITIDINE	SIV
02336480	RAN-RANITIDINE	RBV
02247814	RIVA-RANITIDINE	RIV
02243229	SANDOZ RANITIDINE	SDZ

ST **300MG TABLET**

02248571	ACT RANITIDINE	TEV
00733067	APO-RANITIDINE	APX
02463725	JAMP-RANITIDINE	JMP
02443716	MAR-RANITIDINE	MAR
02473542	M-RANITIDINE	MAN
02242454	PMS-RANITIDINE	PMS
00740756	RANITIDINE	PDL
02353024	RANITIDINE	SAN
02385961	RANITIDINE	SIV
02336502	RAN-RANITIDINE	RBV
02247815	RIVA-RANITIDINE	RIV
02243230	SANDOZ RANITIDINE	SDZ

56:28.28 PROSTAGLANDINS

MISOPROSTOL

ST **100MCG TABLET**

02244022	MISOPROSTOL	AAP
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ST **200MCG TABLET**

02244023	MISOPROSTOL	AAP
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56:28.32 PROTECTANTS

SUCRALFATE

ST **200MG/ML SUSPENSION**

02103567	SULCRATE PLUS	APC
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ST **1G TABLET**

02125250	APO-SUCRALFATE	APX
02100622	SULCRATE	APC
02045702	TEVA-SUCRALFATE	TEV

56:28.36 PROTON-PUMP INHIBITORS

AMOXICILLIN, CLARITHROMYCIN, LANSOPRAZOLE

ST **500MG & 500MG & 30MG KIT**

02470780	APO-LANSOPRAZOLE-AMOXICILLIN-CLARITHROMYCIN	APX
02238525	HP-PAC	TAK

56:28.36 PROTON-PUMP INHIBITORS

LANSOPRAZOLE

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

(Please refer to Appendix A).

ST **15MG CAPSULE (DELAYED RELEASE)**

02293811	APO-LANSOPRAZOLE	APX
02357682	LANSOPRAZOLE	SAN
02385767	LANSOPRAZOLE	SIV
02433001	LANSOPRAZOLE	PMS
02353830	MYLAN-LANSOPRAZOLE	MYL
02395258	PMS-LANSOPRAZOLE	PMS
02165503	PREVACID	TAK
02402610	RAN-LANSOPRAZOLE	RBV
02422808	RIVA-LANSOPRAZOLE	RIV
02385643	SANDOZ LANSOPRAZOLE	SDZ
02280515	TEVA-LANSOPRAZOLE	TEV

ST **30MG CAPSULE (DELAYED RELEASE)**

02293838	APO-LANSOPRAZOLE	APX
02414775	DOM-LANSOPRAZOLE	DPC
02357690	LANSOPRAZOLE	SAN
02366282	LANSOPRAZOLE	PDL
02410389	LANSOPRAZOLE	SIV
02433028	LANSOPRAZOLE	PMS
02353849	MYLAN-LANSOPRAZOLE	MYL
02395266	PMS-LANSOPRAZOLE	PMS
02165511	PREVACID	TAK
02402629	RAN-LANSOPRAZOLE	RBV
02422816	RIVA-LANSOPRAZOLE	RIV
02280523	TEVA-LANSOPRAZOLE	TEV

ST **30MG TABLET (DELAYED RELEASE)**

02385651	SANDOZ LANSOPRAZOLE	SDZ
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PDIN FOR EXTEMPORANEOUS MIXTURE

99503010	LANSOPRAZOLE ORAL LIQUID	UNK
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LANSOPRAZOLE ODT

Limited use benefit (prior approval required).

Coverage will be limited to 400 tablets/capsules every 180 days.

For children 12 years of age or under who are unable to swallow the capsule formulation; OR
For patients with dysphagia or a feeding tube when the use of the capsule formulation is not possible.

(Please refer to Appendix A).

ST **15MG TABLET (DELAYED RELEASE)**

02249464	PREVACID FASTAB	TAK
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ST **30MG TABLET (DELAYED RELEASE)**

02249472	PREVACID FASTAB	TAK
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56:28.36 PROTON-PUMP INHIBITORS

OMEPRAZOLE MAGNESIUM

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

(Please refer to Appendix A).

ST **20MG CAPSULE (DELAYED RELEASE)**

02245058	APO-OMEPRAZOLE	APX
00846503	LOSEC	AZC
02339927	OMEPRAZOLE	PDL
02348691	OMEPRAZOLE	SAN
02411857	OMEPRAZOLE-20	SIV
02320851	PMS-OMEPRAZOLE	PMS
02403617	RAN-OMEPRAZOLE	RBV
02296446	SANDOZ OMEPRAZOLE	SDZ

20MG TABLET (DELAYED RELEASE)

02449927	BIO-OMEPRAZOLE	BMI
02420198	JAMP-OMEPRAZOLE DR	JMP
02190915	LOSEC	AZC
02439549	NAT-OMEPRAZOLE DR	NPH
02416549	OMEPRAZOLE	ACC
02374870	RAN-OMEPRAZOLE	RBV
02402416	RIVA-OMEPRAZOLE DR	RIV
02295415	TEVA-OMEPRAZOLE	TEV
02432404	VAN-OMEPRAZOLE	VAN

PDIN FOR EXTEMPOREANEOUS MIXTURE

99503002	OMEPRAZOLE ORAL LIQUID	UNK
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PANTOPRAZOLE MAGNESIUM

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

(Please refer to Appendix A).

ST **40MG TABLET (DELAYED RELEASE)**

02466147	PANTOPRAZOLE T	SAN
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ST **40MG TABLET (ENTERIC COATED)**

02408570	MYLAN-PANTOPRAZOLE T	MYL
02441853	PANTOPRAZOLE MAGNESIUM	UNK
02267233	TECTA	TAK
02440628	TEVA-PANTOPRAZOLE MAGNESIUM	TEV

PANTOPRAZOLE SODIUM

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

(Please refer to Appendix A).

ST **40MG TABLET (DELAYED RELEASE)**

02292920	APO-PANTOPRAZOLE	APX
02415208	AURO-PANTOPRAZOLE	AUR
02445867	BIO-PANTOPRAZOLE	BMI
02310007	DOM-PANTOPRAZOLE	DPC
02357054	JAMP-PANTOPRAZOLE	JMP
02416565	MAR-PANTOPRAZOLE	MAR
02417448	MINT-PANTOPRAZOLE	MIN

56:28.36 PROTON-PUMP INHIBITORS

PANTOPRAZOLE SODIUM

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

(Please refer to Appendix A).

ST **40MG TABLET (DELAYED RELEASE)**

02467372	M-PANTOPRAZOLE	MAN
02471825	NRA-PANTOPRAZOLE	UNK
02229453	PANTOLOC	TAK
02318695	PANTOPRAZOLE	PDL
02370808	PANTOPRAZOLE	SAN
02431327	PANTOPRAZOLE	RIV
02437945	PANTOPRAZOLE	PMS
02439107	PANTOPRAZOLE	DPC
02428180	PANTOPRAZOLE-40	SIV
02307871	PMS-PANTOPRAZOLE	PMS
02425378	PRIVA-PANTOPRAZOLE	PHA
02305046	RAN-PANTOPRAZOLE	RBV
02316463	RIVA-PANTOPRAZOLE	RIV
02301083	SANDOZ PANTOPRAZOLE	SDZ
02285487	TEVA-PANTOPRAZOLE	TEV

RABEPRAZOLE SODIUM

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

(Please refer to Appendix A).

ST **10MG TABLET (ENTERIC COATED)**

02345579	APO-RABEPRAZOLE	APX
02243796	PARIET	JSO
02310805	PMS-RABEPRAZOLE	PMS
02315181	PRO-RABEPRAZOLE	PDL
02385449	RABEPRAZOLE	SIV
02356511	RABEPRAZOLE EC	SAN
02298074	RAN-RABEPRAZOLE	RBV
02330083	RIVA-RABEPRAZOLE EC	RIV
02314177	SANDOZ RABEPRAZOLE	SDZ
02296632	TEVA-RABEPRAZOLE	TEV

ST **20MG TABLET (ENTERIC COATED)**

02345587	APO-RABEPRAZOLE	APX
02320460	DOM-RABEPRAZOLE EC	DPC
02243797	PARIET	JSO
02310813	PMS-RABEPRAZOLE	PMS
02315203	PRO-RABEPRAZOLE	PDL
02385457	RABEPRAZOLE	SIV
02356538	RABEPRAZOLE EC	SAN
02298082	RAN-RABEPRAZOLE	RBV
02330091	RIVA-RABEPRAZOLE	RIV
02314185	SANDOZ RABEPRAZOLE	SDZ
02296640	TEVA-RABEPRAZOLE	TEV

56:32.00 PROKINETIC AGENTS

DOMPERIDONE MALEATE

ST **10MG TABLET**

02103613	APO-DOMPERIDONE	APX
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56:32.00 PROKINETIC AGENTS

DOMPERIDONE MALEATE

ST 10MG TABLET

02445034	BIO-DOMPERIDONE	BMI
02238315	DOM-DOMPERIDONE	DPC
02236857	DOMPERIDONE	PDL
02238341	DOMPERIDONE	SIV
02350440	DOMPERIDONE	SAN
02369206	JAMP-DOMPERIDONE	JMP
02403870	MAR-DOMPERIDONE	MAR
02236466	PMS-DOMPERIDONE	PMS
02268078	RAN-DOMPERIDONE	RBY
01912070	TEVA-DOMPERIDONE	TEV

PDIN FOR EXTEMPORANEOUS MIXTURE

99503005	DOMPERIDONE ORAL LIQUID	UNK
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METOCLOPRAMIDE HYDROCHLORIDE

ST 1MG/ML SOLUTION

02230433	METONIA	PED
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ST 5MG TABLET

00842826	APO-METOCLOP	APX
02230431	METONIA	PED

ST 10MG TABLET

00842834	APO-METOCLOP	APX
02230432	METONIA	PED

56:36.00 ANTI-INFLAMMATORY AGENTS

BETAMETHASONE SODIUM PHOSPHATE

0.05MG/ML ENEMA

02060884	BETNESOL	PAL
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HYDROCORTISONE ACETATE

10% AEROSOL

00579335	CORTIFOAM	PAL
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100MG/60ML ENEMA

02112736	CORTENEMA	APC
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MESALAZINE

500MG SUPPOSITORY

02112760	SALOFALK	APC
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1G SUPPOSITORY

02474018	MEZERA	UNK
02153564	PENTASA	FEI
02242146	SALOFALK	APC

1G/100ML SUSPENSION

02153521	PENTASA	FEI
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2G/60G SUSPENSION

02112795	SALOFALK	APC
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4G/100ML SUSPENSION

02153556	PENTASA	FEI
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4G/60G SUSPENSION

02112809	SALOFALK	APC
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ST 500MG TABLET (DELAYED RELEASE)

02112787	SALOFALK	APC
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ST 800MG TABLET (DELAYED RELEASE)

02267217	ASACOL	ALL
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ST 400MG TABLET (ENTERIC COATED)

01997580	ASACOL	ALL
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56:36.00 ANTI-INFLAMMATORY AGENTS

MESALAZINE

ST 400MG TABLET (ENTERIC COATED)

02171929	TEVA-5 ASA	TEV
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ST 500MG TABLET (EXTENDED RELEASE)

02099683	PENTASA	FEI
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ST 1G TABLET (EXTENDED RELEASE)

02399466	PENTASA	FEI
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ST 1.2G TABLET (EXTENDED RELEASE)

02297558	MEZAVANT	SHI
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OLSALAZINE SODIUM

ST 250MG CAPSULE

02063808	DIPENTUM	APU
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56:92.00 MISCELLANEOUS GI DRUGS

OBETICHOLIC ACID

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

The patient has a confirmed diagnosis of primary biliary cholangitis (PBC), defined as:

- Positive antimitochondrial antibodies (AMA); OR
- Liver biopsy results consistent with PBC.

AND

The patient is under the care of a gastroenterologist, hepatologist or internal medicine specialist with experience in the treatment of PBC.

AND

The patient has received ursodeoxycholic acid (UDCA) for a minimum of 12 months and has experienced an inadequate response to UDCA and can benefit from the addition of obeticholic acid. An inadequate response is defined as:

- Alkaline phosphatase (ALP) ≥ 1.67 x upper limit of normal (ULN); AND/OR
- Bilirubin $>$ ULN and < 2 x ULN; AND/OR
- Evidence of compensated cirrhosis by fibroscan or biopsy.

OR

The patient has experienced documented and unmanageable intolerance to UDCA.

Criteria for renewal every 12 months:

The patient continues to benefit from treatment with obeticholic acid as evidenced by:

- A reduction in the ALP level to less than 1.67 x ULN; OR
- A 15% reduction in the ALP level compared with values before beginning treatment with obeticholic acid.

5MG TABLET

02463121	OCALIVA	UNK
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10MG TABLET

02463148	OCALIVA	UNK
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PINAVERIUM BROMIDE

Limited use benefit (prior approval required).

For the treatment and relief of symptoms associated with functional bowel disorders including Irritable Bowel Syndrome (IBS), spastic colon, spastic colitis and mucous colitis; OR In postoperative paralytic ileus in order to accelerate the resumption of the intestinal transit following abdominal surgery.

50MG CAPSULE

00465240	DICETEL	SPH
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50MG TABLET

02469677	APO-PINAVERIUM	APX
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56:92.00 MISCELLANEOUS GI DRUGS

PINAVERIUM BROMIDE

Limited use benefit (prior approval required).

For the treatment and relief of symptoms associated with functional bowel disorders including Irritable Bowel Syndrome (IBS), spastic colon, spastic colitis and mucous colitis; OR In postoperative paralytic ileus in order to accelerate the resumption of the intestinal transit following abdominal surgery.

50MG TABLET

01950592	DICETEL	BGP
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100MG TABLET

02469685	APO-PINAVERIUM	APX
02230684	DICETEL	BGP

60:00 GOLD COMPOUNDS

60:00.00 GOLD COMPOUNDS

AURANOFIN

3MG CAPSULE

01916823 RIDAURA XED

SODIUM AUROTHIOMALATE

10MG/ML SOLUTION

01927620 MYOCHRYSSINE SAC

02245456 SODIUM AUROTHIOMALATE SDZ

25MG/ML SOLUTION

01927612 MYOCHRYSSINE SAC

50MG/ML SOLUTION

01927604 MYOCHRYSSINE SAC

02245458 SODIUM AUROTHIOMALATE SDZ

64:00 HEAVY METAL ANTAGONISTS

64:00.00 HEAVY METAL ANTAGONISTS

PENICILLAMINE

250MG CAPSULE

00016055 CUPRIMINE

VAE

**68:00 HORMONES AND SYNTHETIC
SUBSTITUTES**

68:04.00 ADRENALS

BECLOMETHASONE DIPROPIONATE

50MCG AEROSOL

02242029 QVAR VAE

100MCG AEROSOL

02242030 QVAR VAE

BUDESONIDE

3MG CAPSULE (SUSTAINED RELEASE)

02229293 ENTOCORT TIL

100MCG POWDER

00852074 PULMICORT TURBUHALER AZC

200MCG POWDER

00851752 PULMICORT TURBUHALER AZC

400MCG POWDER

00851760 PULMICORT TURBUHALER AZC

0.125MG SUSPENSION

02465949 TEVA-BUDESONIDE TEV

0.125MG/ML SUSPENSION

02229099 PULMICORT NEBUAMP AZC

0.25MG/ML SUSPENSION

01978918 PULMICORT NEBUAMP AZC

0.5MG SUSPENSION

02465957 TEVA-BUDESONIDE TEV

0.5MG/ML SUSPENSION

01978926 PULMICORT NEBUAMP AZC

CICLESONIDE

100MG/INHALATION AEROSOL

02285606 ALVESCO AZC

200MG/INHALATION AEROSOL

02285614 ALVESCO AZC

CORTISONE ACETATE

25MG TABLET

00280437 CORTISONE VAE

DEXAMETHASONE

0.1MG/ML LIQUID

01946897 PMS DEXAMETHASONE PMS

0.5MG TABLET

02261081 APO-DEXAMETHASONE APX

01964976 PMS DEXAMETHASONE PMS

0.75MG TABLET

01964968 PMS DEXAMETHASONE PMS

2MG TABLET

02279363 PMS-DEXAMETHASONE PMS

4MG TABLET

02250055 APO-DEXAMETHASONE APX

01964070 PMS DEXAMETHASONE PMS

02311267 PRO-DEXAMETHASONE PDL

PDIN FOR EXTEMPORANEOUS MIXTURE

99503007 DEXAMETHASONE ORAL LIQUID UNK

68:04.00 ADRENALS

DEXAMETHASONE PHOSPHATE

4MG/ML LIQUID

00664227 DEXAMETHASONE SDZ

01977547 DEXAMETHASONE RAX

02204266 DEXAMETHASONE-OMEGA OMG

10MG/ML LIQUID

00874582 DEXAMETHASONE SDZ

02204274 DEXAMETHASONE-OMEGA OMG

00783900 PMS-DEXAMETHASONE PMS

FLUDROCORTISONE ACETATE

0.1MG TABLET

02086026 FLORINEF PAL

FLUTICASONE PROPIONATE

50MCG/INHALATION AEROSOL

02244291 FLOVENT HFA GSK

125MCG/INHALATION AEROSOL

02244292 FLOVENT HFA GSK

250MCG/INHALATION AEROSOL

02244293 FLOVENT HFA GSK

100MCG/DOSE POWDER

02237245 FLOVENT DISKUS GSK

250MCG/DOSE POWDER

02237246 FLOVENT DISKUS GSK

500MCG/DOSE POWDER

02237247 FLOVENT DISKUS GSK

**HYDROCORTISONE (HYDROCORTISONE
SODIUM SUCCINATE)**

100MG POWDER FOR SOLUTION

00030600 SOLU-CORTEF ACT-O-VIAL PFI

250MG POWDER FOR SOLUTION

00030619 SOLU-CORTEF ACT-O-VIAL PFI

1G POWDER FOR SOLUTION

00030635 SOLU-CORTEF ACT-O-VIAL PFI

HYDROCORTISONE ACETATE

10MG TABLET

00030910 CORTEF PFI

20MG TABLET

00030929 CORTEF PFI

METHYLPREDNISOLONE

4MG TABLET

00030988 MEDROL PFI

16MG TABLET

00036129 MEDROL PFI

**METHYLPREDNISOLONE
(METHYLPREDNISOLONE SODIUM SUCCINATE)**

40MG INJECTION

02367947 SOLU-MEDROL PFI

125MG INJECTION

02367955 SOLU-MEDROL PFI

500MG INJECTION

00030678 SOLU-MEDROL PFI

1G INJECTION

00036137 SOLU-MEDROL PFI

68:04.00 ADRENALS

**METHYLPREDNISOLONE
(METHYLPREDNISOLONE SODIUM SUCCINATE)**

1G INJECTION

02367971 SOLU-MEDROL PFI

500MG POWDER FOR SOLUTION

02231895 METHYLPREDNISOLONE SODIUM SUCCINATE TEV

1G POWDER FOR SOLUTION

02241229 METHYLPREDNISOLONE SODIUM SUCCINATE TEV

METHYLPREDNISOLONE ACETATE

20MG/ML SUSPENSION

01934325 DEPO-MEDROL PFI

40MG/ML SUSPENSION

00030759 DEPO-MEDROL PFI

01934333 DEPO-MEDROL PFI

02245400 METHYLPREDNISOLONE SDZ

02245407 METHYLPREDNISOLONE SDZ

80MG/ML SUSPENSION

00030767 DEPO-MEDROL PFI

01934341 DEPO-MEDROL PFI

02245406 METHYLPREDNISOLONE SDZ

02245408 METHYLPREDNISOLONE SDZ

**METHYLPREDNISOLONE ACETATE, LIDOCAINE
HYDROCHLORIDE**

40MG & 10MG SUSPENSION

00260428 DEPO-MEDROL WITH LIDOCAINE PFI

MOMETASONE FUROATE

200MCG POWDER

02243595 ASMANEX TWISTHALER FRS

400MCG POWDER

02243596 ASMANEX TWISTHALER FRS

PREDNISOLONE SODIUM PHOSPHATE

1MG/ML SOLUTION

02230619 PEDIAPRED SAC

02245532 PMS-PREDNISOLONE PMS

PREDNISONONE

1MG TABLET

00598194 APO PREDNISONONE APX

00271373 WINPRED AAP

5MG TABLET

00312770 APO PREDNISONONE APX

00021695 TEVA-PREDNISONONE TEV

50MG TABLET

00550957 APO PREDNISONONE APX

00232378 TEVA-PREDNISONONE TEV

PDIN FOR EXTEMPORANEOUS MIXTURE

99503008 PREDNISONONE ORAL LIQUID UNK

TRIAMCINOLONE ACETONIDE

40MG/ML INJECTION

00990876 KENALOG-40 BMS

10MG/ML SUSPENSION

01999761 KENALOG-10 BMS

68:04.00 ADRENALS

TRIAMCINOLONE ACETONIDE

10MG/ML SUSPENSION

02229540 TRIAMCINOLONE SDZ

40MG/ML SUSPENSION

01999869 KENALOG-40 BMS

01977563 TRIAMCINOLONE RAX

02229550 TRIAMCINOLONE SDZ

09857128 TRIAMCINOLONE UNK

TRIAMCINOLONE DIACETATE

40MG/ML SUSPENSION

01977555 TRIAMCINOLONE RAX

68:08.00 ANDROGENS

DANAZOL

50MG CAPSULE

02018144 CYCLOMEN SAC

100MG CAPSULE

02018152 CYCLOMEN SAC

200MG CAPSULE

02018160 CYCLOMEN SAC

TESTOSTERONE (TOPICAL)

Limited use benefit (prior approval required).

The NIH Program covers topical testosterone for the treatment of the following in adult males above 18 years old.

- Orchiectomy, undescended testes, Klinefelter's; OR
- Pituitary tumour or post-pituitary surgery with low testosterone; OR
- AIDS-wasting syndrome with low testosterone; OR
- Gender affirming hormone therapy.

Note: Older males with non-specific symptoms such as, but not limited to, fatigue, malaise, or depression who have a low random testosterone level do not meet coverage criteria.

1% GEL

02245345 ANDROGEL BGP

02245346 ANDROGEL BGP

02463792 TARO-TESTOSTERONE TAR

02463806 TARO-TESTOSTERONE TAR

02280248 TESTIM PAL

12.5MG GEL

02249499 ANDROGEL BGP

2.5MG PATCH

02239653 ANDRODERM ALL

5MG PATCH

02245972 ANDRODERM ALL

TESTOSTERONE CYPIONATE

100MG/ML SOLUTION

00030783 DEPO-TESTOSTERONE PFI

02246063 TESTOSTERONE CYPIONATE SDZ

TESTOSTERONE ENANTHATE

200MG/ML SOLUTION

00029246 DELATESTRYL VAE

TESTOSTERONE UNDECANOATE

40MG CAPSULE

02322498 PMS-TESTOSTERONE PMS

68:08.00 ANDROGENS

TESTOSTERONE UNDECANOATE

40MG CAPSULE

02421186 TARO-TESTOSTERONE TAR

68:12.00 CONTRACEPTIVES

DESOGESTREL, ETHINYL ESTRADIOL

ST 25MCG & 150MCG, 125MCG, 100MCG TABLET

02272903 LINESSA 21 ASP

02257238 LINESSA 28 ASP

ETHINYL ESTRADIOL, DESOGESTREL

ST 30MCG & 150MCG TABLET

02317192 APRI 21 TEV

02317206 APRI 28 TEV

02396491 FREYA 21 MYL

02396610 FREYA 28 MYL

02042487 MARVELON 21 FRS

02042479 MARVELON 28 FRS

02410249 MIRVALA 21 APX

02410257 MIRVALA 28 APX

ETHINYL ESTRADIOL, DROSPIRENONE

ST 0.02MG & 3MG TABLET

02415380 MYA APX

02321157 YAZ BAY

ST 0.03MG & 3MG TABLET

02261723 YASMIN 21 BAY

02261731 YASMIN 28 BAY

02410788 ZAMINE 21 APX

02410796 ZAMINE 28 APX

ETHINYL ESTRADIOL, ETHYNODIOL DIACETATE

ST 30MCG & 2MG TABLET

00469327 DEMULEN 30 (21 DAY PACK) PFI

00471526 DEMULEN 30 (28 DAY PACK) PFI

ETHINYL ESTRADIOL, ETONOGESTREL

ST 2.6MG & 11.4MG RING (SLOW-RELEASE)

02253186 NUVARING FRS

ETHINYL ESTRADIOL, LEVONORGESTREL

0.03MG & 0.15MG TABLET

02398869 INDAYO MYL

ST 0.15MG & 0.03MG TABLET

02296659 SEASONALE TEV

ST 20MCG & 100MCG TABLET

02236974 ALESSE 21 PFI

02236975 ALESSE 28 PFI

02387875 ALYSENA 21 APX

02387883 ALYSENA 28 APX

02298538 AVIANE 21 TEV

02298546 AVIANE 28 TEV

ST 30MCG & 0.05MG, 40MCG & 0.075MG, 30MCG & 0.125MG TABLET

00707600 TRIQUILAR 21 BAY

00707503 TRIQUILAR 28 BAY

ST 30MCG & 150MCG TABLET

02042320 MIN-OVRAL 21 PFI

02042339 MIN-OVRAL 28 PFI

68:12.00 CONTRACEPTIVES

ETHINYL ESTRADIOL, LEVONORGESTREL

ST 30MCG & 150MCG TABLET

02387085 OVIMA 21 APX

02387093 OVIMA 28 APX

02295946 PORTIA 21 TEV

02295954 PORTIA 28 TEV

ETHINYL ESTRADIOL, NORELGESTROMIN

ST 6MG & 0.6MG PATCH (EXTENDED RELEASE)

02248297 EVRA JSO

ETHINYL ESTRADIOL, NORETHINDRONE

35MCG & 0.5MG TABLET

02187086 BREVICON 0.5/35 (21-DAY PACK) PFI

02187094 BREVICON 0.5/35 (28-DAY PACK) PFI

ST 35MCG & 1MG TABLET

02189054 BREVICON 1/35 (21-DAY PACK) PFI

02189062 BREVICON 1/35 (28-DAY PACK) PFI

02197502 SELECT 1/35 (21-DAY) PFI

02199297 SELECT 1/35 (28-DAY) PFI

ETHINYL ESTRADIOL, NORETHINDRONE ACETATE

ST 10MCG & 1MG TABLET

02417456 LOLO ALL

ST 20MCG & 1MG TABLET

00315966 MINESTRIN 1/20 (21-DAY) ALL

00343838 MINESTRIN 1/20 (28-DAY) ALL

ST 30MCG & 1.5MG TABLET

00297143 LOESTRIN ALL

00353027 LOESTRIN ALL

ETHINYL ESTRADIOL, NORGESTIMATE

ST 35MCG & 0.25MG TABLET

01968440 CYCLEN (21 DAY) JSO

01992872 CYCLEN (28 DAY) JSO

LEVONORGESTREL

19.5MG INSERT (EXTENDED-RELEASE)

02459523 KYLEENA BAY

0.75MG TABLET

02371189 OPTION 2 PER

1.5MG TABLET

02433532 BACKUP PLAN ONESTEP APX

02425009 CONTINGENCY ONE MYL

02293854 PLAN B UNK

LEVONORGESTREL INTRAUTERINE INSERT

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 1 device every 2 years.

52MG INSERT (EXTENDED-RELEASE)

02243005 MIRENA BAY

LEVONORGESTREL, ETHINYL ESTRADIOL

ST 0.15MG & 0.03MG & 0.01MG TABLET

02346176 SEASONIQUE TEV

68:12.00 CONTRACEPTIVES

NORETHINDRONE

ST **0.35MG TABLET**

02441306	JENCYCLA	LUP
00037605	MICRONOR 28-DAY	JSO
02410303	MOVISSE	MYL

NORETHINDRONE, ETHINYL ESTRADIOL

35MCG & 0.5MG, 35MCG & 1MG TABLET

02187108	SYNPHASIC 21	PFI
02187116	SYNPHASIC 28	PFI

NORGESTIMATE, ETHINYL ESTRADIOL

ST **25MCG & 0.180MG, 25MCG & 0.215MG, 25MCG & 0.25MG TABLET**

02401967	TRICIRA LO 21	APX
02401975	TRICIRA LO 28	APX
02258560	TRI-CYCLEN LO (21 DAY)	JSO
02258587	TRI-CYCLEN LO (28 DAY)	JSO

ST **35MCG & 0.180MG, 35MCG & 0.215MG, 35MCG & 0.25MG TABLET**

02028700	TRI-CYCLEN 21-DAY	JSO
02029421	TRI-CYCLEN 28-DAY	JSO

ULIPRISTAL ACETATE

Limited use benefit (prior approval not required).

Coverage will be limited to 90 tablets, benefits only for women age 18 to 60 years.

ST **5MG TABLET**

02408163	FIBRISTAL	ALL
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68:16.04 ESTROGENS

CONJUGATED ESTROGENS

ST **0.625MG/G CREAM**

02043440	PREMARIN	PFI
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ST **0.3MG TABLET (EXTENDED RELEASE)**

02414678	PREMARIN	PFI
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ST **0.625MG TABLET (EXTENDED RELEASE)**

02414686	PREMARIN	PFI
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ST **1.25MG TABLET (EXTENDED RELEASE)**

02414694	PREMARIN	PFI
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CONJUGATED ESTROGENS, MEDROXYPROGESTERONE ACETATE

ST **0.625MG & 2.5MG TABLET**

02242878	PREMPLUS	PFI
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ST **0.625MG & 5MG TABLET**

02242879	PREMPLUS	PFI
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ESTRADIOL

ST **0.25MG GEL**

02424924	DIVIGEL	SEA
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ST **0.5MG GEL**

02424835	DIVIGEL	SEA
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ST **1MG GEL**

02424843	DIVIGEL	SEA
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ST **25MCG PATCH**

02245676	ESTRADOT 25	NVR
02243722	OESCLIM	SEA

68:16.04 ESTROGENS

ESTRADIOL

ST **37.5MCG PATCH**

02243999	ESTRADOT 37.5	NVR
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ST **50MCG PATCH**

02244000	ESTRADOT 50	NVR
02243724	OESCLIM	SEA

ST **75MCG PATCH**

02244001	ESTRADOT 75	NVR
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ST **100MCG PATCH**

02244002	ESTRADOT 100	NVR
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ST **2MG RING (SLOW-RELEASE)**

02168898	ESTRING	PFI
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ST **0.5MG TABLET**

02225190	ESTRACE	TRM
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ST **1MG TABLET**

02148587	ESTRACE	TRM
----------	---------	-----

ST **2MG TABLET**

02148595	ESTRACE	TRM
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ESTRADIOL HEMIHYDRATE

ST **0.06% GEL**

02238704	ESTROGEL	FRS
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ST **25MCG PATCH**

02247499	CLIMARA 25	BAY
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ST **50MCG PATCH**

02231509	CLIMARA 50	BAY
02246967	SANDOZ ESTRADIOL DERM	SDZ

ST **75MCG PATCH**

02247500	CLIMARA 75	BAY
02246968	SANDOZ ESTRADIOL DERM	SDZ

ST **100MCG PATCH**

02246969	SANDOZ ESTRADIOL DERM	SDZ
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ST **0.5MG TABLET**

02449048	LUPIN-ESTRADIOL	LUP
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ST **1MG TABLET**

02449056	LUPIN-ESTRADIOL	LUP
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ST **2MG TABLET**

02449064	LUPIN-ESTRADIOL	LUP
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ST **10MCG VAGINAL TABLET**

02325462	VAGIFEM 10	NOO
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ESTRADIOL, NORETHINDRONE ACETATE

ST **50MCG & 140MCG PATCH**

02241835	ESTALIS	NVR
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ST **50MCG & 250MCG PATCH**

02241837	ESTALIS	NVR
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ESTRONE

ST **1MG/G CREAM**

00727369	ESTRAGYN	SEA
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68:16.12 ESTROGEN AGONISTS-ANTAGONISTS

RALOXIFENE HYDROCHLORIDE

Limited use benefit (prior approval required).

For secondary prevention of osteoporosis in women who experience failure on bisphosphonates.
For secondary prevention of osteoporosis in women who have a personal history or a first degree relative with a history of breast cancer.

60MG TABLET

02358840	ACT RALOXIFENE	ACG
02279215	APO-RALOXIFENE	APX
02239028	EVISTA	LIL
02358921	PMS-RALOXIFENE	PMS

68:18.00 GONADOTROPINS

GOSERELIN ACETATE

3.6MG/DEPOT IMPLANT

02049325	ZOLADEX	UNK
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NAFARELIN ACETATE

2MG/ML AEROSOL

02188783	SYNAREL	PFI
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68:18.04

DEGARELIX ACETATE

80MG POWDER FOR SOLUTION

02337029	FIRMAGON	FEI
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120MG POWDER FOR SOLUTION

02337037	FIRMAGON	FEI
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68:18.08

LEUPROLIDE ACETATE

3.75MG/VIAL POWDER FOR SUSPENSION

00884502	LUPRON DEPOT	ABV
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7.5MG/VIAL POWDER FOR SUSPENSION

00836273	LUPRON DEPOT	ABV
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11.25MG/VIAL POWDER FOR SUSPENSION

02239834	LUPRON DEPOT	ABV
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22.5MG/VIAL POWDER FOR SUSPENSION

02230248	LUPRON DEPOT	ABV
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30MG/VIAL POWDER FOR SUSPENSION

02239833	LUPRON DEPOT	ABV
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68:20.02 ALPHA-GLUCOSIDASE INHIBITORS

ACARBOSE

ST **50MG TABLET**

02190885	GLUCOBAY	BAY
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ST **100MG TABLET**

02190893	GLUCOBAY	BAY
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68:20.04 BIGUANIDES

METFORMIN HYDROCHLORIDE

ST **500MG TABLET**

02257726	ACT METFORMIN	TEV
02167786	APO-METFORMIN	APX
02438275	AURO-METFORMIN	AUR

68:20.04 BIGUANIDES

METFORMIN HYDROCHLORIDE

ST **500MG TABLET**

02229994	DOM-METFORMIN	DPC
02099233	GLUCOPHAGE	SAC
02229516	GLYCON	VAE
02380196	JAMP-METFORMIN	JMP
02380722	JAMP-METFORMIN BLACKBERRY	JMP
02353377	METFORMIN	SAN
02378841	METFORMIN	MAR
02385341	METFORMIN FC	SIV
02388766	MINT-METFORMIN	MIN
02223562	PMS-METFORMIN	PMS
02314908	PRO-METFORMIN	PDL
02269031	RAN-METFORMIN	RBV
02242974	RATIO-METFORMIN	TEV
02239081	RIVA-METFORMIN	RIV
02246820	SANDOZ METFORMIN FC	SDZ
02379767	SEPTA-METFORMIN	SPT

ST **850MG TABLET**

02257734	ACT METFORMIN	TEV
02229785	APO-METFORMIN	APX
02438283	AURO-METFORMIN	AUR
02242726	DOM-METFORMIN	DPC
02162849	GLUCOPHAGE	SAC
02239214	GLYCON	VAE
02380218	JAMP-METFORMIN	JMP
02380730	JAMP-METFORMIN BLACKBERRY	JMP
02353385	METFORMIN	SAN
02378868	METFORMIN	MAR
02385368	METFORMIN FC	SIV
02388774	MINT-METFORMIN	MIN
02242589	PMS-METFORMIN	PMS
02314894	PRO-METFORMIN	PDL
02269058	RAN-METFORMIN	RBV
02242931	RATIO-METFORMIN	TEV
02242783	RIVA-METFORMIN	RIV
02246821	SANDOZ METFORMIN	SDZ
02379775	SEPTA-METFORMIN	SPT

68:20.05 DIPEPTIDYL PEPTIDASE-4 (DPP-4) INHIBITORS

LINAGLIPTIN

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **5MG TABLET**

02370921	TRAJENTA	BOE
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68:20.05 DIPEPTIDYL PEPTIDASE-4 (DPP-4) INHIBITORS

LINAGLIPTIN, METFORMIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST 2.5MG & 1000MG TABLET		
02403277	JENTADUETO	BOE
ST 2.5MG & 500MG TABLET		
02403250	JENTADUETO	BOE
ST 2.5MG & 850MG TABLET		
02403269	JENTADUETO	BOE

SAXAGLIPTIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who: did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST 2.5MG TABLET		
02375842	ONGLYZA	AZC
ST 5MG TABLET		
02333554	ONGLYZA	AZC

SAXAGLIPTIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST 2.5MG & 1000MG TABLET		
02389185	KOMBOGLYZE	AZC
ST 2.5MG & 500MG TABLET		
02389169	KOMBOGLYZE	AZC
ST 2.5MG & 850MG TABLET		
02389177	KOMBOGLYZE	AZC

SITAGLIPTIN PHOSPHATE MONOHYDRATE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST 25MG TABLET		
02388839	JANUVIA	FRS
ST 50MG TABLET		
02388847	JANUVIA	FRS
ST 100MG TABLET		
02303922	JANUVIA	FRS

68:20.05 DIPEPTIDYL PEPTIDASE-4 (DPP-4) INHIBITORS

SITAGLIPTIN PHOSPHATE MONOHYDRATE, METFORMIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST 50MG & 1000MG TABLET		
02333872	JANUMET	FRS
ST 50MG & 500MG TABLET		
02333856	JANUMET	FRS
ST 50MG & 850MG TABLET		
02333864	JANUMET	FRS
ST 50MG & 1000MG TABLET (EXTENDED RELEASE)		
02416794	JANUMET XR	FRS
ST 50MG & 500MG TABLET (EXTENDED RELEASE)		
02416786	JANUMET XR	FRS
ST 100MG & 1000MG TABLET (EXTENDED RELEASE)		
02416808	JANUMET XR	FRS

68:20.08 INSULINS

INSULIN (30% NEUTRAL & 70% ISOPHANE) HUMAN BIOSYNTHETIC

100U/ML INJECTION		
00795879	HUMULIN 30/70	LIL
01959212	HUMULIN 30/70 CARTRIDGE	LIL
09853855	HUMULIN 30/70 CARTRIDGE	LIL
02024217	NOVOLIN GE 30/70	NOO
02025248	NOVOLIN GE 30/70 PENFILL	NOO
09853812	NOVOLIN GE 30/70 PENFILL	NOO

INSULIN (40% NEUTRAL & 60% ISOPHANE) HUMAN BIOSYNTHETIC

100U/ML INJECTION		
02024314	NOVOLIN GE 40/60 PENFILL	NOO

INSULIN (50% NEUTRAL & 50% ISOPHANE) HUMAN BIOSYNTHETIC

100U/ML INJECTION		
02024322	NOVOLIN GE 50/50 PENFILL	NOO

INSULIN (ISOPHANE) HUMAN BIOSYNTHETIC

100U/ML INJECTION		
00587737	HUMULIN N	LIL
01959239	HUMULIN N (CARTRIDGE)	LIL
02403447	HUMULIN N (KWIKPEN)	LIL
09853804	HUMULIN N 100U/ML (CARTRIDGE)	LIL
02024225	NOVOLIN GE NPH	NOO
09853782	NOVOLIN GE NPH 100U/ML PENFILL	NOO
02024268	NOVOLIN GE NPH PENFILL	NOO

INSULIN (ZINC CRYSTALLINE) HUMAN BIOSYNTHETIC (RDNA ORIGIN)

100U/ML INJECTION		
00586714	HUMULIN R	LIL
09853766	HUMULIN R 100U/ML (CARTRIDGE)	LIL

68:20.08 INSULINS

INSULIN (ZINC CRYSTALLINE) HUMAN BIOSYNTHETIC (RDNA ORIGIN)

100U/ML INJECTION

01959220 HUMULIN R CARTRIDGE LIL

INSULIN ASPART

100U/ML INJECTION

02244353 NOVORAPID NOO

02245397 NOVORAPID NOO

02377209 NOVORAPID NOO

INSULIN BIOSYNTHETIC HUMAN BR

100U SOLUTION

02415089 HUMULIN R (KWIKPEN) LIL

INSULIN DEGLUDEC

100U SOLUTION

02467879 TRESIBA NOO

200U SOLUTION

02467887 TRESIBA NOO

INSULIN DETEMIR

100U/ML INJECTION

02412829 LEVEMIR FLEXTOUCH NOO

02271842 LEVEMIR PENFILL NOO

INSULIN GLARGINE

100U/ML INJECTION

02245689 LANTUS SAC

02251930 LANTUS SAC

02294338 LANTUS SOLOSTAR SAC

100U SOLUTION

02444844 BASAGLAR LIL

02444852 BASAGLAR LIL

02461528 BASAGLAR LIL

300U SOLUTION

02441829 TOUJEO SOLOSTAR SAC

INSULIN GLULISINE

100U/ML INJECTION

02279479 APIDRA CARTRIDGE SAC

02294346 APIDRA SOLOSTAR SAC

02279460 APIDRA VIAL SAC

INSULIN HUMAN BIOSYNTHETIC

100U/ML INJECTION

02024233 NOVOLIN GE TORONTO NOO

02024284 NOVOLIN GE TORONTO PENFILL NOO

09853774 NOVOLIN GE TORONTO PENFILL NOO

INSULIN LISPRO

100U/ML INJECTION

02229704 HUMALOG LIL

02229705 HUMALOG (CARTRIDGE) LIL

02403412 HUMALOG (KWIKPEN) LIL

09853715 HUMALOG 100U/ML CARTRIDGE LIL

200U/ML INJECTION

02439611 HUMALOG 200U/ML KWIKPEN LIL

68:20.08 INSULINS

INSULIN LISPRO, INSULIN LISPRO PROTAMINE

100U/ML INJECTION

02240294 HUMALOG MIX 25 (CARTRIDGE) LIL

02403420 HUMALOG MIX 25 (KWIKPEN) LIL

02240297 HUMALOG MIX 50 (CARTRIDGE) LIL

02403439 HUMALOG MIX 50 (KWIKPEN) LIL

68:20.16 MEGLITINIDES

REPAGLINIDE

ST **0.5MG TABLET**

02321475 ACT REPAGLINIDE ACG

02355663 APO-REPAGLINIDE APX

02424258 AURO-REPAGLINIDE AUR

02239924 GLUCONORM NOO

02354926 JAMP REPAGLINIDE JMP

02415968 REPAGLINIDE PDL

02357453 SANDOZ REPAGLINIDE SDZ

ST **1MG TABLET**

02321483 ACT REPAGLINIDE ACG

02355671 APO-REPAGLINIDE APX

02424266 AURO-REPAGLINIDE AUR

02239925 GLUCONORM NOO

02354934 JAMP REPAGLINIDE JMP

02415976 REPAGLINIDE PDL

02357461 SANDOZ REPAGLINIDE SDZ

ST **2MG TABLET**

02321491 ACT REPAGLINIDE ACG

02355698 APO-REPAGLINIDE APX

02424274 AURO-REPAGLINIDE AUR

02239926 GLUCONORM NOO

02354942 JAMP REPAGLINIDE JMP

02415984 REPAGLINIDE PDL

02357488 SANDOZ REPAGLINIDE SDZ

68:20.18 SODIUM-GLUCOSE CONTRANSPORTER 2 (SGLT2) INHIBITORS

CANAGLIFLOZIN

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **100MG TABLET**

02425483 INVOKANA JSO

ST **300MG TABLET**

02425491 INVOKANA JSO

**68:20.18 SODIUM-GLUCOSE
CONTRANSPORTER 2 (SGLT2)
INHIBITORS**

**DAPAGLIFLOZIN PROPANEDIOL
MONOHYDRATE**

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **5MG TABLET**

02435462 FORXIGA AZC

ST **10MG TABLET**

02435470 FORXIGA AZC

EMPAGLIFLOZIN

Open benefit with therapeutic notes (prior approval is not required).

For the treatment of type 2 diabetes mellitus:
- in patients who did not achieve glycemic control with an adequate trial of metformin AND a sulfonylurea
OR
- to reduce the incidence of cardiovascular death in patients with established cardiovascular disease who did not achieve adequate glycemic control despite an appropriate trial of metformin

Established cardiovascular disease is defined as one of the following:

- history of myocardial infarction
- 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 either a positive non-invasive stress test or discharged from hospital with a documented diagnosis of unstable angina
- unstable angina with confirmed evidence of coronary multi-vessel or single-vessel disease
- history of ischemic or hemorrhagic stroke
- occlusive peripheral artery disease.

ST **10MG TABLET**

02443937 JARDIANCE BOE

ST **25MG TABLET**

02443945 JARDIANCE BOE

**METFORMIN HYDROCHLORIDE,
DAPAGLIFLOZIN**

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **850MG & 5MG TABLET**

02449935 XIGDUO AZC

ST **1000MG & 5MG TABLET**

02449943 XIGDUO AZC

**68:20.18 SODIUM-GLUCOSE
CONTRANSPORTER 2 (SGLT2)
INHIBITORS**

**METFORMIN HYDROCHLORIDE,
EMPAGLIFLOZIN**

Open benefit with therapeutic notes (prior approval is not required).

For the treatment of patients with type 2 diabetes mellitus in patients who are eligible to receive metformin and empagliflozin, to replace the individual components.

500MG & 12.5MG TABLET

02456605 SYNJARDY BOE

500MG & 5MG TABLET

02456575 SYNJARDY BOE

850MG & 12.5MG TABLET

02456613 SYNJARDY BOE

850MG & 5MG TABLET

02456583 SYNJARDY BOE

1000MG & 12.5MG TABLET

02456621 SYNJARDY BOE

1000MG & 5MG TABLET

02456591 SYNJARDY BOE

**68:20.20 ANTIDIABETIC AGENTS -
SULFONYLUREAS**

GLICLAZIDE

ST **80MG TABLET**

02245247 APO-GLICLAZIDE APX

00765996 DIAMICRON SEV

02248453 GLICLAZIDE PDL

02287072 GLICLAZIDE SAN

02238103 TEVA-GLICLAZIDE TEV

ST **30MG TABLET (EXTENDED RELEASE)**

02297795 APO-GLICLAZIDE MR APX

02242987 DIAMICRON MR SEV

02429764 JAMP GLICLAZIDE-MR JMP

02423286 MINT-GLICLAZIDE MR MIN

02438658 MYLAN-GLICLAZIDE MR MYL

02463571 RAN-GLICLAZIDE MR RBY

02461323 SANDOZ GLICLAZIDE MR SDZ

ST **60MG TABLET (EXTENDED RELEASE)**

02407124 APO-GLICLAZIDE MR APX

02356422 DIAMICRON MR SEV

02423294 MINT-GLICLAZIDE MR MIN

02439328 RAN-GLICLAZIDE RBY

02461331 SANDOZ GLICLAZIDE MR SDZ

GLYBURIDE

ST **2.5MG TABLET**

01913654 APO GLYBURIDE APX

02224550 DIABETA SAC

01959352 GLYBURIDE PDL

02350459 GLYBURIDE SAN

01913670 TEVA-GLYBURIDE TEV

ST **5MG TABLET**

01913662 APO GLYBURIDE APX

02224569 DIABETA SAC

**68:20.20 ANTIDIABETIC AGENTS -
SULFONYLUREAS**

GLYBURIDE

ST **5MG TABLET**

02234514	DOM-GLYBURIDE	DPC
00720941	EUGLUCON	PMS
02350467	GLYBURIDE	SAN
02236734	PMS-GLYBURIDE	PMS
01913689	TEVA-GLYBURIDE	TEV

**68:20.28 THIAZOLIDINEDIONES
PIOGLITAZONE HYDROCHLORIDE**

ST **15MG TABLET**

02303442	ACCEL PIOGLITAZONE	ACP
02391600	ACH-PIOGLITAZONE	ACC
02302861	ACT PIOGLITAZONE	ACG
02302942	APO-PIOGLITAZONE	APX
02307634	DOM-PIOGLITAZONE	DPC
02397307	JAMP-PIOGLITAZONE	JMP
02326477	MINT-PIOGLITAZONE	MIN
02303124	PMS-PIOGLITAZONE	PMS
02312050	PRO-PIOGLITAZONE	PDL
02375850	RAN-PIOGLITAZONE	RBV
02297906	SANDOZ PIOGLITAZONE	SDZ
02434121	VAN-PIOGLITAZONE	VAN

ST **30MG TABLET**

02303450	ACCEL PIOGLITAZONE	ACP
02339587	ACH-PIOGLITAZONE	ACC
02302888	ACT PIOGLITAZONE	ACG
02302950	APO-PIOGLITAZONE	APX
02307642	DOM-PIOGLITAZONE	DPC
02365529	JAMP-PIOGLITAZONE	JMP
02326485	MINT-PIOGLITAZONE	MIN
02303132	PMS-PIOGLITAZONE	PMS
02312069	PRO-PIOGLITAZONE	PDL
02375869	RAN-PIOGLITAZONE	RBV
02297914	SANDOZ PIOGLITAZONE	SDZ
02434148	VAN-PIOGLITAZONE	VAN

ST **45MG TABLET**

02303469	ACCEL PIOGLITAZONE	ACP
02339595	ACH-PIOGLITAZONE	ACC
02302896	ACT PIOGLITAZONE	ACG
02302977	APO-PIOGLITAZONE	APX
02307650	DOM-PIOGLITAZONE	DPC
02365537	JAMP-PIOGLITAZONE	JMP
02326493	MINT-PIOGLITAZONE	MIN
02303140	PMS-PIOGLITAZONE	PMS
02312077	PRO-PIOGLITAZONE	PDL
02375877	RAN-PIOGLITAZONE	RBV
02297922	SANDOZ PIOGLITAZONE	SDZ
02434156	VAN-PIOGLITAZONE	VAN

**68:22.12 GLYCOGENOLYTIC AGENTS
GLUCAGON RECOMBINANT DNA ORGIN**

1MG/ML INJECTION

02333619	GLUCAGEN	NOO
02333627	GLUCAGEN HYPOKIT	NOO

**68:22.12 GLYCOGENOLYTIC AGENTS
GLUCAGON RECOMBINANT DNA ORGIN**

1MG/ML INJECTION

02243297	GLUCAGON	LIL
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68:24.00 PARATHYROID

CALCITONIN SALMON (SYNTHETIC)

200IU/ML SOLUTION

01926691	CALCIMAR	SAC
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68:28.00 PITUITARY

DESMOPRESSIN ACETATE

4MCG/ML LIQUID

00873993	DDAVP	FEI
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0.1MG/ML NASAL SPRAY

00402516	DDAVP	FEI
00836362	DDAVP	FEI
02242465	DESMOPRESSIN	AAP

ST **0.1MG TABLET**

00824305	DDAVP	FEI
02284030	DESMOPRESSIN	APX
02304368	PMS-DESMOPRESSIN	PMS
02287730	TEVA-DESMOPRESSIN	TEV

ST **0.2MG TABLET**

00824143	DDAVP	FEI
02284049	DESMOPRESSIN	APX
02304376	PMS-DESMOPRESSIN	PMS

ST **60MCG TABLET (ORALLY DISINTEGRATING)**

02284995	DDAVP MELT	FEI
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ST **120MCG TABLET (ORALLY DISINTEGRATING)**

02285002	DDAVP MELT	FEI
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ST **240MCG TABLET (ORALLY DISINTEGRATING)**

02285010	DDAVP MELT	FEI
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68:32.00 PROGESTINS

DIENOGEST

Limited use benefit (prior approval required).

For the management of pelvic pain associated with endometriosis.

ST **2MG TABLET**

02374900	VISANNE	BAY
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MEDROXYPROGESTERONE ACETATE

50MG/ML SUSPENSION

00030848	DEPO-PROVERA	PFI
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150MG/ML SUSPENSION

00585092	DEPO-PROVERA	PFI
02322250	MEDROXYPROGESTERONE	SDZ

ST **2.5MG TABLET**

02244726	APO-MEDROXY	APX
02253550	MEDROXY	PDL
00708917	PROVERA	PFI
02221284	TEVA-MEDROXYPROGESTERONE	TEV

ST **5MG TABLET**

02244727	APO-MEDROXY	APX
02253577	MEDROXY	PDL
00030937	PROVERA	PFI

68:32.00 PROGESTINS

MEDROXYPROGESTERONE ACETATE

ST 5MG TABLET			
02221292	TEVA-MEDROXYPROGESTERONE	TEV	
ST 10MG TABLET			
02277298	APO-MEDROXY	APX	
00729973	PROVERA	PFI	
02221306	TEVA-MEDROXYPROGESTERONE	TEV	
ST 100MG TABLET			
02267640	APO-MEDROXY	APX	

PROGESTERONE

Limited use benefit (prior approval required).

For the treatment of women:

- With postmenopausal symptoms who are intolerant to medroxyprogesterone acetate (MPA); OR
- Who are at risk of preterm birth; OR
- Who are using the medication to prevent miscarriage.

In adults:

- For use as Gender Affirming Hormone Therapy.

100MG CAPSULE			
02476576	PMS-PROGESTERONE	PMS	
02166704	PROMETRIUM	FRS	
02463113	REDDY-PROGESTERONE	REC	
02439913	TEVA-PROGESTERONE	TEV	

68:36.04 THYROID AGENTS

LEVOTHYROXINE SODIUM

ST 0.025MG TABLET			
02264323	EUTHYROX	SRO	
02172062	SYNTHROID	BGP	
ST 0.05MG TABLET			
02213192	ELTROXIN	ASP	
02264331	EUTHYROX	SRO	
02172070	SYNTHROID	BGP	
ST 0.075MG TABLET			
02264358	EUTHYROX	SRO	
02172089	SYNTHROID	BGP	
ST 0.088MG TABLET			
02172097	SYNTHROID	BGP	
ST 0.1MG TABLET			
02213206	ELTROXIN	ASP	
02264374	EUTHYROX	SRO	
02172100	SYNTHROID	BGP	
ST 0.112MG TABLET			
02264390	EUTHYROX	SRO	
02171228	SYNTHROID	BGP	
ST 0.125MG TABLET			
02264404	EUTHYROX	SRO	
02172119	SYNTHROID	BGP	
ST 0.137MG TABLET			
02264412	EUTHYROX	SRO	
02233852	SYNTHROID	BGP	
ST 0.15MG TABLET			
02213214	ELTROXIN	ASP	
02264420	EUTHYROX	SRO	
02172127	SYNTHROID	BGP	

68:36.04 THYROID AGENTS

LEVOTHYROXINE SODIUM

ST 0.175MG TABLET			
02264439	EUTHYROX	SRO	
02172135	SYNTHROID	BGP	
ST 0.2MG TABLET			
02213222	ELTROXIN	ASP	
02264447	EUTHYROX	SRO	
02172143	SYNTHROID	BGP	
ST 0.3MG TABLET			
02264455	EUTHYROX	SRO	
02172151	SYNTHROID	BGP	

LIOthyRONINE SODIUM

ST 5MCG TABLET			
01919458	CYTOMEL	PFI	
ST 25MCG TABLET			
01919466	CYTOMEL	PFI	

THYROID

ST 30MG TABLET			
00023949	THYROID	ERF	
ST 60MG TABLET			
00023957	THYROID	ERF	
ST 125MG TABLET			
00023965	THYROID	ERF	

68:36.08 ANTITHYROID AGENTS

PROPYLTHIOURACIL

ST 50MG TABLET			
00010200	PROPYL-THYRACIL	PAL	
ST 100MG TABLET			
00010219	PROPYL-THYRACIL	PAL	

THIAMAZOLE

ST 5MG TABLET			
02480107	MAR-METHIMAZOLE	MAR	
00015741	TAPAZOLE	PAL	
ST 10MG TABLET			
02480115	MAR-METHIMAZOLE	MAR	
02296039	TAPAZOLE	PAL	

72:00 LOCAL ANESTHETICS

72:00.00 LOCAL ANESTHETICS

LIDOCAINE HYDROCHLORIDE

2% LIQUID

00811874 PMS-LIDOCAINE VISCOUS PMS

2% SOLUTION

01968823 LIDODAN VISCOUS ODN

76:00 OXYTOCICS

76:00.00 OXYTOCICS

MISOPROSTOL, MIFEPRISTONE

200MCG & 200MG TABLET

02444038 MIFEGYMISO

LIP

**84:00 SKIN AND MUCOUS
MEMBRANE AGENTS (SMMA)**

84:04.04 SMMA - ANTIBIOTICS

BACITRACIN ZINC

500IU OINTMENT

00584908	BACITIN	PED
02351714	JAMP-BACITRACINE	JMP

CLINDAMYCIN PHOSPHATE

2% CREAM

02060604	DALACIN	PFI
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1% LIQUID

99101429	CLINDAMYCIN	UNK
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1% SOLUTION

02243659	CLINDA-T	VAE
00582301	DALACIN T	PFI
02266938	TARO-CLINDAMYCIN	TAR

PDIN FOR EXTEMPORANEOUS MIXTURE

99502000	CLINDAMYCIN IN DILUSOL OR DUONALC	UNK
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**CLINDAMYCIN PHOSPHATE, BENZOYL
PEROXIDE**

1% & 3% GEL

02382822	CLINDOXYL ADV	GSK
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1% & 5% GEL

02248472	BENZACLIN	VAE
02243158	CLINDOXYL	GSK
02464519	TARO-BENZOYL PEROXIDE / CLINDAMYCIN KIT	TAR
02440180	TARO-CLINDAMYCIN/BENZOYL PEROXIDE	TAR

ERYTHROMYCIN, BENZOYL PEROXIDE

3% & 5% GEL

02225271	BENZAMYCIN	VAE
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FUSIDATE SODIUM

2% OINTMENT

00586676	FUCIDIN	LEO
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FUSIDIC ACID

2% CREAM

00586668	FUCIDIN	LEO
----------	---------	-----

FUSIDIC ACID, HYDROCORTISONE ACETATE

2% & 1% CREAM

02238578	FUCIDIN H	LEO
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METRONIDAZOLE

1% CREAM

02156091	NORITATE	BSH
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0.75% GEL

02092832	METROGEL	GAC
02125226	NIDAGEL	VAE

1% GEL

02297809	METROGEL	GAC
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0.75% LOTION

02248206	METROLOTION	GAC
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84:04.04 SMMA - ANTIBIOTICS

METRONIDAZOLE, NYSTATIN

500MG & 100,000IU SUPPOSITORY

01926829	FLAGYSTATIN	SAC
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MUPIROICIN

2% OINTMENT

01916947	BACTROBAN	GSK
02279983	TARO-MUPIROICIN	TAR

MUPIROICIN CALCIUM

2% CREAM

02239757	BACTROBAN	GSK
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POLYMYXIN B SULFATE, BACITRACIN ZINC

10,000IU & 500IU OINTMENT

02304473	ANTIBIOTIC OINT	PMS
00876488	BACIMYXIN ONGUENT	PMS
00621366	BIODERM	ODN
02357569	JAMPOLYCIN	JMP
02237227	POLYSPORIN ANTIBIOTIC	JAJ
01942921	POLYTOPIC	SDZ

**POLYMYXIN B SULFATE, BACITRACIN ZINC,
GRAMICIDIN**

10,000U & 500U & 0.25MG OINTMENT

02237226	POLYSPORIN TRIPLE	JAJ
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POLYMYXIN B SULFATE, GRAMICIDIN

0.25MG & 10,000IU CREAM

02230844	POLYSPORIN ANTIBIOTIC	JAJ
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84:04.06 SMMA - ANTIVIRALS

ACYCLOVIR

5% CREAM

02039524	ZOVIRAX	VAE
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5% OINTMENT

02477130	APO-ACYCLOVIR	APX
00569771	ZOVIRAX	VAE

SINECATECHINS

10% OINTMENT

02411849	VEREGEN	PAL
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84:04.08 SMMA - ANTIFUNGALS

**BETAMETHASONE DIPROPIONATE,
CLOTRIMAZOLE**

0.05% & 1% CREAM

00611174	LOTRIDERM	FRS
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CICLOPIROX OLAMINE

1% CREAM

02221802	LOPROX	VAE
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1% LOTION

02221810	LOPROX	VAE
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CLOTRIMAZOLE

1% CREAM

02150867	CANESTEN	BAY
02150891	CANESTEN	BAY
00812366	CLOTRIMADERM	TAR

84:04.08 SMMA - ANTIFUNGALS

CLOTRIMAZOLE

1% CREAM

00812382	CLOTRIMADERM	TAR
02229380	CLOTRIMAZOLE	TAR
00874043	NEO-ZOL	PPI
00874051	NEO-ZOL	PPI

2% CREAM

02150905	CANESTEN	BAY
00812374	CLOTRIMADERM	TAR

1% & 200MG TABLET (CONTROLLED RELEASE)

02264099	CANESTEN COMBI-PAK COMFORTAB 3	BAY
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1% & 500MG TABLET (CONTROLLED RELEASE)

02264102	CANESTEN COMBI-PAK COMFORTAB 1	BAY
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500MG VAGINAL TABLET

02150859	CANESTEN COMFORTAB 1	BAY
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KETOCONAZOLE

2% CREAM

02245662	KETODERM	TPT
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2% SHAMPOO

02182920	NIZORAL	UNK
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MICONAZOLE NITRATE

2% CREAM

02085852	MICATIN	WPC
02231106	MICOZOLE	TAR
02084309	MONISTAT 7	INS
02126567	MONISTAT DERM	INS

2% & 100MG CREAM/VAGINAL SUPPOSITORY

02126257	MONISTAT 7 DUAL-PAK	INS
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2% & 400MG CREAM/VAGINAL SUPPOSITORY

02126249	MONISTAT 3 DUAL-PAK	INS
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400MG OVULE

02126605	MONISTAT 3	INS
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400MG SUPPOSITORY

02171775	MICONAZOLE 3 DAY OVULE TREATMENT	VTH
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NYSTATIN

25,000IU CREAM

00716901	NYADERM	TAR
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100,000IU CREAM

00716871	NYADERM	TAR
02194236	RATIO-NYSTATIN	TEV
02194163	TEVA-NYSTATIN	TEV

100,000IU OINTMENT

02194228	RATIO-NYSTATIN	TEV
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TERBINAFINE HYDROCHLORIDE

1% CREAM

02031094	LAMISIL	NVR
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TERCONAZOLE

0.4% CREAM

02247651	TARO-TERCONAZOLE	TAR
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84:04.08 SMMA - ANTIFUNGALS

TOLNAFTATE

1% AEROSOL

00576050	TINACTIN AEROSOL	BAY
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1% CREAM

00576034	TINACTIN	BAY
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1% POWDER

01919245	DRSCHOLL'S ATHLETE'S FOOT SPRAY	BAY
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00576042	TINACTIN	BAY
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84:04.12 SMMA - SCABICIDES AND PEDICULICIDES

CROTAMITON

10% CREAM

00623377	EURAX	CLC
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DIMETHICONE

50% SOLUTION

02373785	NYDA	GPB
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ISOPROPYL MYRISTATE

50% SOLUTION

02279592	RESULTZ	MDF
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PERMETHRIN

1% CREAM

00771368	NIX	INS
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5% CREAM

02219905	NIX DERMAL	GSK
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1% LIQUID

02231480	KWELLADA-P	MTC
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5% LOTION

02231348	KWELLADA-P	MTC
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PIPERONYL BUTOXIDE, PYRETHRINS

3% & 0.3% SHAMPOO

02125447	R & C SHAMPOO WITH CONDITIONER	MTC
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84:04.92 SMMA - MISCELLANEOUS LOCAL ANTI-INFECTIVES

ISOPROPYL ALCOHOL

70% LIQUID

00426539	DUONALC	ICN
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METRONIDAZOLE

10% CREAM

01926861	FLAGYL	SAC
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POVIDONE-IODINE

10% SOLUTION

00158348	BETADINE	PFR
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SELENIUM SULFIDE

2.5% LOTION

00594601	VERSEL	VAE
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2.5% SHAMPOO

00243000	EXTRA STRENGTH SELSUN	SAC
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**84:04.92 SMMA - MISCELLANEOUS
LOCAL ANTI-INFECTIVES**

SILVER SULFADIAZINE

1% CREAM

00323098	FLAMAZINE	SNE
09854037	FLAMAZINE	SMW

**84:06.00 SMMA - ANTI-INFLAMMATORY
AGENTS**

AMCINONIDE

0.1% CREAM

02246714	TARO-AMCINONIDE	TAR
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0.1% LOTION

02247097	RATIO-AMCINONIDE	TEV
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0.1% OINTMENT

02247096	RATIO-AMCINONIDE	TEV
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BECLOMETHASONE DIPROPIONATE

0.025% CREAM

02089602	PROPADERM	VAE
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BETAMETHASONE DIPROPIONATE

0.05% CREAM

00688622	DIPROLENE	FRS
00323071	DIPROSONE	FRS
02122073	ROLENE	RIV
02122049	ROSONE	RIV
01925350	TARO-SONE	TAR
00849650	TEVA-TOPILENE	TEV
00804991	TEVA-TOPISONE	TEV

0.05% LOTION

00417246	DIPROSONE	FRS
02122065	ROLENE	RIV
02122030	ROSONE	RIV
01927914	TEVA-TOPILENE	TEV
00809187	TEVA-TOPISONE	TEV

0.05% OINTMENT

00629367	DIPROLENE	FRS
00344923	DIPROSONE	FRS
02122081	ROLENE	RIV
02122057	ROSONE	RIV
00849669	TEVA-TOPILENE	TEV
00805009	TEVA-TOPISONE	TEV

**BETAMETHASONE DIPROPIONATE, SALICYLIC
ACID**

0.05% & 2% LOTION

00578428	DIPROSALIC	FRS
02245688	RATIO-TOPISALIC	TEV

0.05% & 3% OINTMENT

00578436	DIPROSALIC	FRS
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PDIN FOR EXTEMPORANEOUS MIXTURE

99500003	SALICYLIC ACID IN CORTICOSTEROID CREAM	UNK
99501001	SALICYLIC ACID IN NON- MEDICATED OINTMENT	UNK

**84:06.00 SMMA - ANTI-INFLAMMATORY
AGENTS**

BETAMETHASONE VALERATE

0.05% CREAM

00716618	BETADERM	TAR
02357860	CELESTODERM V	VAE
00535427	RATIO-ECTOSONE	TEV

0.1% CREAM

00716626	BETADERM	TAR
02357844	CELESTODERM V	VAE
00535435	RATIO-ECTOSONE	TEV

0.05% LOTION

00653209	RATIO-ECTOSONE	TEV
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0.1% LOTION

00716634	BETADERM	TAR
00750050	RATIO-ECTOSONE	TEV
01940112	RIVASONE	RIV
00653217	TEVA-ECTOSONE	TEV
00027944	VALISONE	VAE

0.05% OINTMENT

00716642	BETADERM	TAR
02357879	CELESTODERM V	VAE

0.1% OINTMENT

00716650	BETADERM	TAR
02357852	CELESTODERM V	VAE

BUDESONIDE, SODIUM CHLORIDE

0.02MG/ML ENEMA

02052431	ENTOCORT	TIL
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**CALCIPOTRIOL, BETAMETHASONE
DIPROPIONATE**

50MCG & 0.5MG AEROSOL, FOAM

02457393	ENSTILAR	LEO
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0.5MG & 50MCG GEL

02319012	DOVOBET	LEO
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0.5MG & 50MCG OINTMENT

02244126	DOVOBET	LEO
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CLOBETASOL PROPIONATE

0.05% CREAM

02213265	DERMOVATE	TPT
02024187	MYLAN-CLOBETASOL	MYL
02232191	PMS-CLOBETASOL	PMS
02309521	PMS-CLOBETASOL	PMS
02245523	TARO-CLOBETASOL	TAR
01910272	TEVA-CLOBETASOL	TEV

0.05% LOTION

02213281	DERMOVATE	TPT
02216213	MYLAN-CLOBETASOL	MYL
02232195	PMS-CLOBETASOL	PMS
02245522	TARO-CLOBETASOL	TAR
01910299	TEVA-CLOBETASOL	TEV

0.05% OINTMENT

02213273	DERMOVATE	TPT
02026767	MYLAN-CLOBETASOL	MYL
02309548	PMS-CLOBETASOL	PMS
02245524	TARO-CLOBETASOL	TAR

84:06.00 SMMA - ANTI-INFLAMMATORY AGENTS

CLOBETASOL PROPIONATE

0.05% OINTMENT

01910280 TEVA-CLOBETASOL TEV

CLOBETASONE BUTYRATE

0.05% CREAM

02214415 SPECTRO ECZEMACARE GSK

DESONIDE

0.05% CREAM

02229315 PDP-DESONIDE PED

02154862 TRIDESILON PER

0.05% OINTMENT

02229323 PDP-DESONIDE PED

02154870 TRIDESILON PER

DESOXIMETASONE

0.05% CREAM

02221918 TOPICORT MILD VAE

0.25% CREAM

02221896 TOPICORT VAE

0.05% GEL

02221926 TOPICORT VAE

0.25% OINTMENT

02221934 TOPICORT VAE

ESCULIN, FRAMYCETIN SULFATE, DIBUCAINE HYDROCHLORIDE, HYDROCORTISONE ACETATE

1% & 1% & 0.5% & 0.5% OINTMENT

02247322 PROCTOL ODN

02223252 PROCTOSEDYL APC

02242527 SANDOZ PROCTOMYXIN HC SDZ

10MG & 10MG & 5MG & 5MG OINTMENT

02226383 TEVA-PROCTOSONE TEV

10MG & 10MG & 5MG & 5MG SUPPOSITORY

02247882 PROCTOL ODN

02223260 PROCTOSEDYL APC

02242528 SANDOZ PROCTOMYXIN HC SDZ

02226391 TEVA-PROCTOSONE TEV

FLUOCINONIDE

0.05% CREAM

02163152 LIDEMOL VAE

02161923 LIDEX VAE

00716863 LYDERM TPT

00598933 TIAMOL TPT

0.05% GEL

02161974 LIDEX VAE

02236997 LYDERM TPT

0.01% LOTION

00873292 DERMA-SMOOTHIE HIL

0.025% OINTMENT

02162512 SYNALAR VAE

0.05% OINTMENT

02161966 LIDEX VAE

02236996 LYDERM TPT

84:06.00 SMMA - ANTI-INFLAMMATORY AGENTS

FLUOCINONIDE

0.01% SOLUTION

02162504 SYNALAR VAE

HALOBETASOL PROPIONATE

0.05% CREAM

01962701 ULTRAVATE VAE

0.05% OINTMENT

01962728 ULTRAVATE VAE

HYDROCORTISONE ACETATE

0.5% CREAM

80021088 CORTATE BAY

00716820 HYDERM TAR

02242930 HYDROCORTISONE ACETATE TAR

1% CREAM

00192597 EMOCORT GSK

02412926 EUROHYDROCORTISONE EUR

00716839 HYDERM TAR

00564281 HYDROSONE TEV

80057178 JAMP-HC JMP

80057189 JAMP-HYDROCORTISONE JMP

80066164 M-HC MAN

00804533 PREVEX HC GSK

0.5% LOTION

80021087 CORTATE BAY

1% LOTION

80057191 JAMP-HYDROCORTISONE JMP

80066168 M-HC MAN

00578541 SARNA HC GSK

0.5% OINTMENT

80021085 CORTATE BAY

00716685 CORTODERM TAR

1% OINTMENT

00716693 CORTODERM TAR

HYDROCORTISONE ACETATE, UREA

1% CREAM

80073645 M-HC UREA MAN

1% & 10% CREAM

00681989 DERMAFLEX HC PAL

1% LOTION

80073689 M-HC UREA MAN

1.00% LOTION

00681997 DERMAFLEX HC PAL

HYDROCORTISONE ACETATE, ZINC SULFATE

0.5% & 0.5% OINTMENT

02128446 ANODAN-HC ODN

00505773 ANUSOL HC CHU

02209764 EGOZINC-HC PMS

00607789 RATIO-HEMCORT-HC TEV

02179547 RIVA-HC RIV

02247691 SANDOZ ANUZINC HC SDZ

10MG & 10MG SUPPOSITORY

02236399 ANODAN-HC ODN

00476285 ANUSOL HC CHU

84:06.00 SMMA - ANTI-INFLAMMATORY AGENTS

HYDROCORTISONE ACETATE, ZINC SULFATE

10MG & 10MG SUPPOSITORY

02210517	EGOZINC-HC	PMS
02240112	RIVASOL-HC	RIV
02242798	SANDOZ ANUZINC HC	SDZ

HYDROCORTISONE ACETATE, ZINC SULFATE MONOHYDRATE

0.5% & 0.5% OINTMENT

02387239	JAMP-ZINC-HC	JMP
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HYDROCORTISONE ACETATE, ZINC SULFATE, PRAMOXINE HYDROCHLORIDE

0.5% & 0.5% & 1% OINTMENT

00505781	ANUGESIC HC	MCL
02234466	PROCTODAN-HC	ODN

10MG & 10MG & 20MG SUPPOSITORY

00476242	ANUGESIC HC	MCL
02240851	PROCTODAN-HC	ODN
02242797	SANDOZ ANUZINC HC PLUS	SDZ

HYDROCORTISONE ACETATE-UREA

1% CREAM

80061501	JAMP-HYDROCORTISONE UREA	MAN
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HYDROCORTISONE VALERATE

0.2% CREAM

02242984	HYDROVAL	TPT
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0.2% OINTMENT

02242985	HYDROVAL	TPT
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MOMETASONE FUROATE

0.1% CREAM

00851744	ELOCOM	FRS
02367157	TARO-MOMETASONE	TAR

0.1% LOTION

00871095	ELOCOM	FRS
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0.1% OINTMENT

00851736	ELOCOM	FRS
02244769	PMS-MOMETASONE	PMS
02270862	PMS-MOMETASONE	PMS
02264749	TARO-MOMETASONE	TAR
02266385	TARO-MOMETASONE	TAR
02248130	TEVA-MOMETASONE	TEV

PDIN FOR EXTEMPORANEOUS MIXTURE

99500008	MOMETASONE CREAM	UNK
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TRIAMCINOLONE ACETONIDE

0.1% CREAM

02194058	ARISTOCORT R	VAE
00716960	TRIADERM	TAR

0.5% CREAM

02194066	ARISTOCORT C	VAE
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0.1% OINTMENT

02194031	ARISTOCORT R	VAE
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0.1% PASTE

01964054	ORACORT DENTAL PASTE	TAR
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84:08.00 SMMA - ANTIPRURITICS AND LOCAL ANESTHETICS

LIDOCAINE HYDROCHLORIDE

2% SOLUTION

02427745	JAMPOCAINE VISCOUS	JMP
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LIDOCAINE, PRILOCAINE

2.5% & 2.5% CREAM

00886858	EMLA	UNK
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2.5% & 2.5% PATCH

02057794	EMLA	UNK
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84:16.00 SMMA - CELL STIMULANTS AND PROLIFERANTS

TRETINOIN

0.01% CREAM

00897329	RETIN-A	VAE
00657204	STIEVA-A	GSK

0.025% CREAM

00897310	RETIN-A	VAE
00578576	STIEVA-A	GSK

0.05% CREAM

00443794	RETIN-A	VAE
00518182	STIEVA-A	GSK

0.01% GEL

00870013	RETIN-A	VAE
01926462	VITAMIN A ACID	VAE

0.025% GEL

00443816	RETIN-A	VAE
01926470	VITAMIN A ACID	VAE

0.05% GEL

01926489	VITAMIN A ACID	VAE
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84:24.00 EMOLLIENTS, DEMULCENTS, AND PROTECTANTS

UREA

10% CREAM

80079497	UREMOL 10	ODN
80005397	URISEC10	ODN

20% CREAM

80083394	UREMOL	ODN
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22% CREAM

00396125	URISEC 22	ODN
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10% LOTION

80079498	UREMOL 10	ODN
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12% LOTION

00514896	URISEC 12	ODN
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84:24.12 BASIC OINTMENTS AND PROTECTANTS

DIMETHICONE

20% CREAM

02060841	BARRIERE	WPC
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WHITE PETROLATUM

71.5% OINTMENT

02277778	CRITIC-AID CLEAR	UNK
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84:24.12 BASIC OINTMENTS AND PROTECTANTS

ZINC OXIDE

15% CREAM

02215799 ZINC OXIDE HJS

25% PASTE

00532576 PATE D'IHLE TEV

00886327 PÂTE D'IHLE ATL

ZINC OXIDE, WHITE PETROLATUM

15% & 80.3% CREAM

02337452 DIAPER RASH HJS

40% OINTMENT

02239160 ZINCOFAX EXTRA STRENGTH PAL

84:28.00 KERATOLYTIC AGENTS

BENZOYL PEROXIDE

5% GEL

02162113 BENZAGEL CLC

4% LOTION

02413353 SPECTRO ACNECARE WASH GSK

5% LOTION

02166607 BENZAGEL 5 CLC

5% SOLUTION

02162121 BENZAGEL CLC

CANTHARIDIN

1% LIQUID

80028872 CANTHACUR 07 PAL

CANTHARIDIN, PODOPHYLLIN, SALICYLIC ACID

1% & 2% & 30% LIQUID

00772011 CANTHARONE PLUS DOR

CLINDAMYCIN PHOSPHATE, TRETINOIN

1.2% & 0.025% GEL

02359685 BIACNA TOPICAL VAE

SALICYLIC ACID

170MG/ML GEL

00614246 COMPOUND W GEL UNK

20% LIQUID

00690333 SOLUVER DPT

26% LIQUID

00754951 OCCLUSAL HP VAE

27% LIQUID

00837733 SOLUVER PLUS DPT

40% PLASTER

01967878 DR SCHOLLS CLEAR AWAY PLANTAR WART REMOVER SYSTEM BAY

01974335 DR SCHOLLS CLEAR AWAY WART REMOVER SYSTEM BAY

4% SHAMPOO

00666106 SEBCUR DPT

84:32.00 KERATOPLASTIC AGENTS

COAL TAR

10% GEL

00344508 TARGEL ODN

84:32.00 KERATOPLASTIC AGENTS

COAL TAR

0.5% SHAMPOO

02240645 NEUTROGENA JAJ

1% SHAMPOO

02307146 T/ THERAPEUTIC SHAMPOO EXTRA STRENGTH JAJ

20% SOLUTION

00358495 ODAN LIQUOR CARBONIS DETERGENT ODN

COAL TAR, SALICYLIC ACID

10% & 3% GEL

00510335 TARGEL SA ODN

10% & 4% SHAMPOO

00666114 SEBCUR-T DPT

84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS

ACITRETIN

Open benefit (prior approval not required).

Soriatane should be used with caution in women of childbearing potential due to its teratogenicity. Pregnancy must be excluded. Effective contraception must be used. Manufacturer's literature regarding contraindications and warnings should be consulted prior to prescribing or dispensing this drug.

ST **10MG CAPSULE**

02468840 MINT-ACITRETIN MIN

02070847 SORIATANE ALL

02466074 TARO-ACITRETIN TAR

ST **25MG CAPSULE**

02468859 MINT-ACITRETIN MIN

02070863 SORIATANE ALL

02466082 TARO-ACITRETIN TAR

ADAPALENE

0.1% CREAM

02231592 DIFFERIN GAC

0.1% GEL

02148749 DIFFERIN GAC

AZELAIC ACID

15% GEL

02270811 FINACEA BAY

BRODALUMAB

Limited use benefit (prior approval required).

• Psoriasis according to established criteria.

(Please refer to Appendix A).

210MG SOLUTION

02473623 SILIQ VAE

CALCIPOTRIOL

50MCG/G OINTMENT

01976133 DOVONEX LEO

84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS

CAPSAICIN

0.025% CREAM

02157101	CAPSAICIN	VAE
02244952	ZODERM	EUR
00740306	ZOSTRIX	VAE

0.075% CREAM

02157128	CAPSAISIN	VAE
02004240	ZOSTRIX HP	VAE

COLLAGENASE

250U OINTMENT

02063670	SANTYL	SNE
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FLUOROURACIL

5% CREAM

00330582	EFUDEX	VAE
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IMIQUIMOD

Limited use benefit (prior approval required).

For the treatment of condylomata acuminata (genital warts) in patients who have failed:

- self-applied podophyllotoxin (podofilox 0.5% solution); OR
- provider-applied podophyllum resin (10%-25%).

5% CREAM

02239505	ALDARA P	VAE
02407825	APO-IMIQUIMOD	APX
02482983	TARO-IMIQUIMOD PUMP	TAR

ISOTRETINOIN

Open benefit (prior approval not required).

Accutane should be used with caution in women of childbearing potential due to its teratogenicity. Pregnancy must be excluded. Effective contraception must be used. Manufacturer's literature regarding contraindications and warnings should be consulted prior to prescribing or dispensing this drug.

ST **10MG CAPSULE**

00582344	ACUTANE ROCHE	HLR
02257955	CLARUS	MYL
02396971	EPURIS	CIP

20MG CAPSULE

02396998	EPURIS	CIP
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30MG CAPSULE

02397005	EPURIS	CIP
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ST **40MG CAPSULE**

00582352	ACUTANE ROCHE	HLR
02257963	CLARUS	MYL
02397013	EPURIS	CIP

IXEKIZUMAB

Limited use benefit (prior approval required).

- Psoriatic Arthritis according to established criteria.
- Psoriasis according to established criteria.

(Please refer to Appendix A).

80MG SOLUTION

02455102	TALTZ	LIL
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84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS

IXEKIZUMAB

Limited use benefit (prior approval required).

- Psoriatic Arthritis according to established criteria.
- Psoriasis according to established criteria.

(Please refer to Appendix A).

80MG SOLUTION

02455110	TALTZ	LIL
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LUBRICANT

GEL VAG

09991643	CAYA DIAPHRAGM	TSN
09991644	CONTRAGEL GREEN	TSN

PIMECROLIMUS

Limited use benefit (prior approval required).

For patients who have failed topical corticosteroid therapy or have experienced side effects from such treatment.

Note: Contraindicated in children less than 2 years of age.

1% CREAM

02247238	ELIDEL	VAE
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PODOFILOX

0.5% SOLUTION

01945149	CONDYLINE	SAC
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PODOPHYLLIN

25% LIQUID

00598208	PODOFILM	PAL
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SALICYLIC ACID, FLUOROURACIL

10% & 0.5% SOLUTION

02428946	ACTIKERALL	CIP
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SECUKINUMAB

Limited use benefit (prior approval required).

- Psoriasis according to established criteria.
- Psoriatic Arthritis according to established criteria.
- Ankylosing Spondylitis according to established criteria.

(Please refer to Appendix A).

150MG/ML INJECTION

99101215	COSENTYX (STYLO)	NVC
09857548	COSENTYX PEN (ON)	NVC

150MG SOLUTION

02438070	COSENTYX	NVR
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TACROLIMUS (PROTOPIC)

Limited use benefit (prior approval required).

For patients who have failed topical corticosteroid therapy or have experienced side effects from such treatment.

Note: Contraindicated in children less than 2 years of age.

0.03% OINTMENT

02244149	PROTOPIC	LEO
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0.1% OINTMENT

02244148	PROTOPIC	LEO
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**84:92.00 MISCELLANEOUS SKIN AND
MUCOUS MEMBRANE AGENTS**

TAZAROTENE

0.05% CREAM

02243894 TAZORAC ALL

0.1% CREAM

02243895 TAZORAC ALL

0.05% GEL

02230784 TAZORAC ALL

0.1% GEL

02230785 TAZORAC ALL

86:00 SMOOTH MUSCLE RELAXANTS

86:12.04 ANTIMUSCARINICS

DARIFENACIN HYDROBROMIDE

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:
 • with symptoms of urinary frequency, urgency or urge incontinence; AND
 • who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

7.5MG TABLET (EXTENDED RELEASE)

02273217 ENABLEX UNK

15MG TABLET (EXTENDED RELEASE)

02273225 ENABLEX UNK

FESOTERODINE FUMARATE

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:
 • with symptoms of urinary frequency, urgency or urge incontinence; AND
 • who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

ST 4MG TABLET (EXTENDED RELEASE)

02380021 TOVIAZ PFI

ST 8MG TABLET (EXTENDED RELEASE)

02380048 TOVIAZ PFI

FLAVOXATE HYDROCHLORIDE

ST 200MG TABLET

00728179 URISPAS PAL

OXYBUTYNIN CHLORIDE

ST 1MG/ML SYRUP

02231089 APO-OXYBUTYNIN APX

02223376 PMS-OXYBUTYNIN PMS

ST 2.5MG TABLET

02240549 PMS-OXYBUTYNIN PMS

ST 5MG TABLET

02163543 APO-OXYBUTYNIN APX

02241285 DOM-OXYBUTYNIN DPC

02350238 OXYBUTYNIN SAN

02240550 PMS-OXYBUTYNIN PMS

02299364 RIVA-OXYBUTYNIN RIV

02230394 TEVA-OXYBUTYNIN TEV

PROPIVERINE HYDROCHLORIDE

5MG TABLET

02460289 MICTORYL PEDIATRIC DUI

SOLIFENACIN SUCCINATE

ST 5MG TABLET

02423375 APO-SOLIFENACIN APX

02446375 AURO-SOLIFENACIN AUR

02424339 JAMP-SOLIFENACIN JMP

02428911 MED-SOLIFENACIN GMP

02443171 MINT-SOLIFENACIN MIN

02417723 PMS-SOLIFENACIN PMS

86:12.04 ANTIMUSCARINICS

SOLIFENACIN SUCCINATE

ST 5MG TABLET

02437988 RAN-SOLIFENACIN RBY

02399032 SANDOZ SOLIFENACIN SDZ

02458144 SOLIFENACIN PDL

02458241 SOLIFENACIN SAN

02397900 TEVA-SOLIFENACIN TEV

02277263 VESICARE AST

ST 10MG TABLET

02423383 APO-SOLIFENACIN APX

02446383 AURO-SOLIFENACIN AUR

02424347 JAMP-SOLIFENACIN JMP

02428938 MED-SOLIFENACIN GMP

02443198 MINT-SOLIFENACIN MIN

02417731 PMS-SOLIFENACIN PMS

02437996 RAN-SOLIFENACIN RBY

02399040 SANDOZ SOLIFENACIN SDZ

02458152 SOLIFENACIN PDL

02458268 SOLIFENACIN SAN

02397919 TEVA-SOLIFENACIN TEV

02277271 VESICARE AST

TOLTERODINE TARTRATE

ST 2MG CAPSULE (EXTENDED RELEASE)

02244612 DETROL LA PFI

02404184 MYLAN-TOLTERODINE ER MYL

02413140 SANDOZ TOLTERODINE LA SDZ

02412195 TEVA-TOLTERODINE LA TEV

ST 4MG CAPSULE (EXTENDED RELEASE)

02244613 DETROL LA PFI

02404192 MYLAN-TOLTERODINE ER MYL

02413159 SANDOZ TOLTERODINE LA SDZ

02412209 TEVA-TOLTERODINE LA TEV

ST 1MG TABLET

02369680 APO-TOLTERODINE APX

02239064 DETROL PFI

02423308 MINT-TOLTERODINE MIN

02299593 TEVA-TOLTERODINE TEV

ST 2MG TABLET

02369699 APO-TOLTERODINE APX

02239065 DETROL PFI

02423316 MINT-TOLTERODINE MIN

02299607 TEVA-TOLTERODINE TEV

TROSPIUM CHLORIDE

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:
 • with symptoms of urinary frequency, urgency or urge incontinence; AND
 • who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

ST 20MG TABLET

02275066 TROSEC SPC

86:12.08 BETA-ADRENERGIC AGONISTS**MIRABEGRON**

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:

- with symptoms of urinary frequency, urgency or urge incontinence; AND
- who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

ST **25MG TABLET (EXTENDED RELEASE)**

02402874 MYRBETRIQ AST

ST **50MG TABLET (EXTENDED RELEASE)**

02402882 MYRBETRIQ AST

86:16.00 RESPIRATORY SMOOTH MUSCLE RELAXANTS**OXTRIPHYLLINE**ST **20MG/ML ELIXIR**

00476366 CHOLEDYL ERF

THEOPHYLLINEST **5.33MG/ML ELIXIR**

00466409 PULMOPHYLLINE RIV

01966219 THEOLAIR VAE

00627410 THEOPHYLLINE ATL

ST **100MG TABLET (EXTENDED RELEASE)**

00692689 APO-THEO-LA APX

ST **200MG TABLET (EXTENDED RELEASE)**

00692697 APO-THEO-LA APX

ST **300MG TABLET (EXTENDED RELEASE)**

00692700 APO-THEO-LA APX

ST **400MG TABLET (EXTENDED RELEASE)**

02360101 THEO ER AAP

02014165 UNIPHYL PFR

ST **600MG TABLET (EXTENDED RELEASE)**

02360128 THEO ER AAP

02014181 UNIPHYL PFR

88:00 VITAMINS

88:04.00 VITAMIN A

VITAMIN A

ST **10,000IU CAPSULE**

80054130	JAMP-VITAMIN A	JMP
00297720	VITAMIN A	JAM
00557447	VITAMIN A	VTH

88:08.00 VITAMIN B COMPLEX

CYANOCOBALAMIN

100MCG/ML LIQUID

00497533	VITAMIN B12	HOS
02241500	VITAMIN B12	SDZ

ST **200MCG/ML LIQUID**

80039903	BEDUZIL	ORM
80026092	JAMP-VITAMIN B12	JMP

1,000MCG/ML LIQUID

00626112	B-12	OMG
02052717	CYANOCOBALAMIN	TAR
02413795	CYANOCOBALAMIN	MYL
02420147	JAMP-CYANOCOBALAMIN	JMP
00038830	VITAMIN B12	HOS

1,000MCG/ML SOLUTION

01987003	CYANOCOBALAMIN	RAX
00521515	VITAMIN B12	SDZ

ST **250MCG TABLET**

80015294	JAMP-VITAMIN B12	JMP
80055743	M-B12	MAN
00335940	VITAMIN B12	JAM
02239695	VITAMIN B12	PMT
80004053	VITAMIN B12	WNP

ST **1000MCG TABLET**

80028902	JAMP VITAMIN B12	JMP
80015276	JAMP-VITAMIN B12	JMP
80055741	M-B12	MAN
02237736	VITAMIN B12	VAE
80003575	VITAMIN B12	PMT
80006939	VITAMIN B12	WNP
80012952	VITAMIN B12 SUBLINGUAL	JAM

FOLIC ACID

ST **1MG TABLET**

00318973	FOLIC ACID	JAM
00647039	FOLIC ACID	VTH
02048841	FOLIC ACID	PMT
80000273	FOLIC ACID	WNP
80053274	JAMP FOLIC ACID	JMP
80061488	M-FOLIQUÉ	MAN
02236747	WAMPOLE FOLIC ACID	WAM

ST **5MG TABLET**

00426849	FOLIC ACID	APX
02366061	JAMP-FOLIC ACID	JMP
02285673	SANDOZ FOLIC ACID	SDZ

ST **1000MCG TABLET**

02239882	FOLIC ACID	UNK
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88:08.00 VITAMIN B COMPLEX

NIACIN

ST **500MG CAPLET**

00309737	NIACIN	JAM
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ST **50MG TABLET**

00041084	NIACIN	ADA
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ST **500MG TABLET**

00557412	NIACIN	VTH
01939130	NIACIN	ODN
02247004	NIACIN	PMT

PYRIDOXINE HYDROCHLORIDE

ST **25MG TABLET**

80056458	M-B6	MAN
00122645	VITAMIN B6	JAM
00232475	VITAMIN B6	ADA
01943200	VITAMIN B6	ODN
80002890	VITAMIN B6	JMP

ST **50MG TABLET**

00305227	VITAMIN B6	JAM
00608599	VITAMIN B6	ADA

ST **100MG TABLET**

00450677	B6	VTH
00263958	VITAMIN B6	VAE
00329185	VITAMIN B6	JAM
02239348	VITAMIN B6	PMT

THIAMINE HYDROCHLORIDE

100MG/ML LIQUID

02193221	THIAMJECT	OMG
02243525	THIAMINE	RAX

100MG/ML SOLUTION

00816078	VITAMIN B1	SDZ
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ST **50MG TABLET**

02245506	EURO VITAMIN B1	EUR
80054199	M-B1	MAN
00268631	THIAMINE	VAE
80009633	VITAMIN B1	JMP

ST **100MG TABLET**

80054205	M-B1	MAN
00232467	VITAMIN B1	PED
00407011	VITAMIN B1	JAM
02239350	VITAMIN B1	PMT
80000352	VITAMIN B1	WNP
80009588	VITAMIN B1	JMP

88:12.00 VITAMIN C

ASCORBIC ACID

ST **500MG CAPLET**

02163268	VITAMIN C	JAM
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ST **250MG TABLET**

00162515	VITAMIN C	PMT
00221244	VITAMIN C	ADA
00266051	VITAMIN C	PMT
00557811	VITAMIN C	VTH

ST **500MG TABLET**

00266086	ASCORBIC ACID	PMT
00041114	VITAMIN C	ADA

88:12.00 VITAMIN C

ASCORBIC ACID

ST 500MG TABLET

00322326	VITAMIN C	ADA
00557838	VITAMIN C	VTH
00784591	VITAMIN C	VTH
01922378	VITAMIN C	VAE
02243893	VITAMIN C	PMT
02244469	VITAMIN C	PMT
02245348	VITAMIN C	WNP
02245721	VITAMIN C	PMT
00322997	VITAMINE C	LAL
00036188	WAMPOLE VITAMIN C	WAM
00274240	WAMPOLE VITAMIN C	WAM

VITAMIN C

ST 500MG TABLET

80003328	VITAMIN C	WNP
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88:16.00 VITAMIN D

ALFACALCIDOL

ST 0.25MCG CAPSULE

00474517	ONE ALPHA	LEO
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ST 1MCG CAPSULE

00474525	ONE ALPHA	LEO
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ST 2MCG/ML DROP

02240329	ONE-ALPHA	LEO
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CALCITRIOL

ST 0.25MCG CAPSULE

02431637	CALCITRIOL-ODAN	ODN
00481823	ROCALTROL	HLR

ST 0.5MCG CAPSULE

02431645	CALCITRIOL-ODAN	ODN
00481815	ROCALTROL	HLR

CHOLECALCIFEROL

ST 400IU CAPSULE

80006629	DGEL	JMP
02242651	EURO D	EUR
80005560	RIVA-D	RIV

ST 800IU CAPSULE

80007769	DGEL	JMP
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1,000IU CAPSULE

80027592	DGEL	OPU
80009635	VITAMIN D3	WAM

ST 10,000IU CAPSULE

02253178	EURO D	SDZ
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ST 400IU LIQUID

80001869	BABY DDROPS	DDP
80001792	DDROPS	DDP

ST 400IU/ML LIQUID

00762881	D VI INFANTS	MJO
80003038	JAMP VITAMIN D	JMP
02231624	PEDIAVIT D	EUR

ST 1,000IU LIQUID

80001791	DDROPS	DDP
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88:16.00 VITAMIN D

CHOLECALCIFEROL

ST 400IU TABLET

02238729	VITAMIN D	VTH
02240858	VITAMIN D	PMT
00765384	VITAMINE D	LAL
02240624	WAMPOLE VITAMIN D	WAM

ST 1,000IU TABLET

02245842	VITAMIN D3	PMT
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ST 10,000IU TABLET

00821772	D-TABS	RIV
02417995	VITAMINE D	PDL

ERGOCALCIFEROL

ST 50,000IU CAPSULE

02237450	SANDOZ D-FORTE	SDZ
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ST 8,288IU/ML SOLUTION

80020776	D2-DOL	JMP
80003615	ERDOL	ODN

VITAMIN D

ST 10MCG CAPSULE

80063895	VIT D 400	UNK
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ST 25MCG CAPSULE

80063899	VIT D 1000	UNK
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ST 200U CAPSULE

02442256	VITAMIN D3	ORM
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ST 400IU CAPSULE

80055196	M-D	MAN
80001145	PHARMA-D	PED
80008590	VITAMINE D	BMI

ST 800IU CAPSULE

80003010	EURO D	EUR
80008446	VITAMINE D	BMI

ST 1,000IU CAPSULE

80007766	DGEL	JMP
80003707	EURO-D	EUR
80055204	M-D	MAN
80008496	PHARMA-D	PMS
80043412	VITAMINE D	BMI

ST 10,000IU CAPSULE

02371499	EURO-D	PMS
02449099	JAMP-VITAMIN D	JMP

ST 15MCG LIQUID

80013189	DDROPS BOOSTER	DDP
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ST 400IU LIQUID

80019649	D3-DOL	JMP
80038155	DECAXIL	ORM
80041145	DECAXIL	ORM

ST 800IU LIQUID

80003285	PEDIAVIT D	EUR
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ST 1,000IU LIQUID

80007346	JAMP VITAMIN D	JMP
80028362	JAMP VITAMIN D	JMP
80028371	JAMP VITAMIN D	JMP

ST 25MCG TABLET

80031157	VITAMIN D	WNP
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88:16.00 VITAMIN D

VITAMIN D

ST **400IU TABLET**

80002452	VITAMIN D	WNP
80009578	VITAMIN D	VAE

ST **1,000IU TABLET**

80002169	PHARMA-D	PMS
80051562	RIVA-D	RIV
80000131	VITAMIN D	VTH
80000436	VITAMIN D	JAM
80003663	VITAMIN D	WNP
80009580	VITAMIN D	VAE
80015278	WAMPOLE VITAMIN D	WAM

ST **10,000IU TABLET**

02379007	JAMP-VITAMIN D	JMP
02417685	VIDEXTRA	ORM

88:20.00 VITAMIN E

VITAMIN E

Limited use benefit (prior approval required).

For use in malabsorption

ST **100IU CAPSULE (SOFTGEL)**

00122823	VITAMIN E	JAM
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ST **200IU CAPSULE (SOFTGEL)**

00122831	VITAMIN E	JAM
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ST **400IU CAPSULE (SOFTGEL)**

00122858	VITAMIN E	JAM
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ST **800IU CAPSULE (SOFTGEL)**

00330191	VITAMIN E	JAM
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ST **20U/ML LIQUID**

09991656	AQUA-E/ML	UNK
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ST **75U/ML LIQUID**

09991652	AQUA-E	UNK
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ST **50IU ORAL LIQUID**

00480215	AQUASOL E	NVC
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ST **50IU/ML ORAL LIQUID**

02162075	AQUASOL E VITAMIN E	CLC
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88:24.00 VITAMIN K

PHYTONADIONE

2MG/ML EMULSION

00781878	VITAMIN K1	SDZ
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10MG/ML EMULSION

00804312	VITAMIN K1	SDZ
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88:28.00 MULTIVITAMIN PREPARATIONS

CALCIUM, VITAMIN D

ST **500-400MGU TABLET**

80088060	BIO-CAL DR FORTE	BIO
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MULTIVITAMINS (CHILDREN AND YOUTH)

Limited use benefit (prior approval is not required).

Multivitamins are benefits for children up to 19 years of age.

ST **DROP**

00762946	ENFAMIL POLYVISOL	MJO
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88:28.00 MULTIVITAMIN PREPARATIONS

MULTIVITAMINS (CHILDREN AND YOUTH)

Limited use benefit (prior approval is not required).

Multivitamins are benefits for children up to 19 years of age.

ST **450MG & 10MG & 30MG LIQUID**

80008471	JAMP VITAMIN A, D AND C	JMP
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ST **2,500IU & 666.67IU & 50MG/ML LIQUID**

00762903	ENFAMIL TRIVISOL	MJO
02229790	PEDIAVIT	EUR

0MG TABLET

02246362	CENTRUM	PFI
80021452	CENTRUM	PFI
80024482	CENTRUM FOR WOMEN	PFI

2MG TABLET

80045908	ONE A DAY WOMEN	BAY
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10MG TABLET

80039441	STRESSTABS FOR WOMEN	PFI
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ST **TABLET (CHEWABLE)**

80011134	CENTRUM JUNIOR COMPLETE	PFI
80020794	CENTRUM JUNIOR COMPLETE	PFI
02247995	FLINTSTONES MULTIPLE VITAMINS PLUS IRON	BAY
02247975	FLINTSTONES MULTIPLE VITAMINS WITH EXTRA C	BAY

MULTIVITAMINS (PRENATAL)

Limited use benefit (prior approval is not required.).

Prenatal and postnatal vitamins are benefits only for women of childbearing age (12 to 50 years).

ST **CAPSULE**

80042704	CENTRUM DHA	PFI
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ST **TABLET**

80045822	CENTRUM PRENATAL	PFI
80080882	MATERNA	NES
80082297	MATERNA	NES
80001842	NESTL MATERNA	NES
02241235	PRENATAL AND POSTPARTUM VITAMINS AND MINERALS	VTH
80005770	PRENATAL AND POSTPARTUM VITAMINS AND MINERALS	PMT
02229535	WAMPOLE COMPLETE MULT-PRE AND POST NATAL WITH FOLIC ACID	WAM

2MG TABLET

80004919	NATURES BOUNTY PRENATAL VITAMINS	VTH
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92:00 UNCLASSIFIED THERAPEUTIC AGENTS

92:00.00 UNCLASSIFIED THERAPEUTIC AGENTS

EXTEMPORANEOUS MIXTURE

CAPSULE

99505003 PHENAZOPYRIDINE COMPOUNDED UNK

CREAM

99500000 HYDROCORTISONE POWDER AND CLOTRIMAZOLE CREAM UNK

99500010 LCD IN CORTICOSTEROID CREAM UNK

99500009 LCD IN NON-MEDICATED CREAM UNK

99500002 MENTHOL &/OR CAMPHOR IN STEROID UNK

99500004 MISCELLANEOUS COMPOUNDED TOPICAL CREAM UNK

99500001 STEROID AND ANTIFUNGAL CREAM UNK

99500006 SULFUR IN NON-MEDICATED CREAM UNK

LOTION

99502001 MENTHOL & CAMPHOR IN CORTICOSTEROID LOTION UNK

99502002 MISCELLANEOUS COMPOUNDED EXTERNAL LOTION UNK

MISCELLANEOUS

00915000 STERILE EXTEMPORANEOUS MIXTURE (QC) UNK

OINTMENT

99501006 ALL PURPOSE NIPPLE OINTMENT UNK

99501003 CALCIUM CHANNEL BLOCKER IN OINTMENT UNK

99501008 DILTIAZEM IN OINTMENT UNK

99501000 LCD IN CORTICOSTEROID OINTMENT UNK

99501005 LCD IN NON-MEDICATED OINTMENT UNK

99501004 MISCELLANEOUS COMPOUNDED TOPICAL OINTMENT UNK

99501002 SULFUR IN NON-MEDICATED OINTMENT UNK

OPHTHALMIC SOLUTION

99507002 ANTIBIOTIC DROPS UNK

99507001 ANTIFUNGAL DROPS UNK

99507003 ANTIVIRAL DROPS UNK

ORAL LIQUID

99503028 ANTACID AND LIDOCAINE ORAL LIQUID UNK

99503029 MAGIC MOUTHWASH UNK

99503025 MISCELLANEOUS COMPOUNDED INTERNAL LIQUID UNK

POWDER

99505004 BACKORDER INTERNAL POWDER UNK

99505000 MISCELLANEOUS COMPOUNDED INTERNAL POWDER UNK

92:00.00 UNCLASSIFIED THERAPEUTIC AGENTS

EXTEMPORANEOUS MIXTURE (GENDER AFFIRMING)

Limited use benefit (prior approval required).

For gender affirming hormone therapy.

INJECTION

00915312 GENDER AFFIRMING HORMONES UNK

LIQUID

00915311 GENDER AFFIRMING TOPICAL HORMONES UNK

EXTEMPORANEOUS MIXTURE (LU)

Limited use benefit (prior approval required).

INJECTION

99506021 MISCELLANEOUS COMPOUNDED INJECTION/INFUSION UNK

MISCELLANEOUS

99504001 MISC LIMITED USE EXTERNAL COMPOUND MIXTURE UNK

OPHTHALMIC AND OTIC SOLUTION

99507000 MISCELLANEOUS COMPOUNDED EYE/EAR DROP UNK

ORAL LIQUID

99503033 MISC LIMITED USE COMPOUND INTERNAL UNK

99503032 OPIOID COMPOUNDED UNK

POWDER

99504000 MISCELLANEOUS COMPOUNDED EXTERNAL POWDER UNK

SUPPOSITORY

99508000 MISCELLANEOUS COMPOUNDED SUPPOSITORY UNK

GOSERELIN ACETATE

10.8MG/DEPOT IMPLANT

02225905 ZOLADEX LA UNK

OCTREOTIDE ACETATE

10MG/VIAL POWDER FOR SUSPENSION (SUSTAINED-RELEASE)

02239323 SANDOSTATIN LAR NVR

20MG/VIAL POWDER FOR SUSPENSION (SUSTAINED-RELEASE)

02239324 SANDOSTATIN LAR NVR

30MG/VIAL POWDER FOR SUSPENSION (SUSTAINED-RELEASE)

02239325 SANDOSTATIN LAR NVR

50MCG/ML SOLUTION

02248639 OCTREOTIDE ACETATE OMEGA OMG

00839191 SANDOSTATIN NVR

100MCG/ML SOLUTION

02248640 OCTREOTIDE ACETATE OMEGA OMG

00839205 SANDOSTATIN NVR

200MCG/ML SOLUTION

02248642 OCTREOTIDE ACETATE OMEGA OMG

02049392 SANDOSTATIN NVR

500MCG/ML SOLUTION

02248641 OCTREOTIDE ACETATE OMEGA OMG

92:00.00 UNCLASSIFIED THERAPEUTIC AGENTS

PENTOSAN POLYSULFATE SODIUM

100MG CAPSULE
02029448 ELMIRON JSO

USTEKINUMAB

Limited use benefit (prior approval required).

- Psoriasis according to established criteria.

(Please refer to Appendix A).

45MG/0.5ML SOLUTION
02320673 STELARA JSO

90MG/ML SOLUTION
02320681 STELARA JSO

92:01.00 NATURAL HEALTH PRODUCTS

CANTHARIDIN

1%(W/V) LIQUID
80023975 CANTHARONE 07 DOR

NATURAL HEALTH PRODUCT

1% CREAM
80066699 CORTIVERA H VAN

PSYLLIUM MUCILLOID

ST **3G POWDER**

80013276 METAMUCIL FIBRE THERAPY SMOOTH TEXTURE PGI

80013287 METAMUCIL FIBRE THERAPY SMOOTH TEXTURE SUGAR FREE PGI

80015505 METAMUCIL SMOOTH TEXTURE UNFLAVOURED UNSWEETENED PGI

92:05.00 SERUMS

APIS MELLIFERA VENOM PROTEIN EXTRACT

1.1MG POWDER FOR SOLUTION
01948903 PHARMALGEN HONEY BEE VENOM ALK

120MCG POWDER FOR SOLUTION
01948911 PHARMALGEN HONEY BEE VENOM ALK

DOLICHOVESPULA ARENARIA VENOM PROTEIN

120MCG POWDER FOR SOLUTION
01948946 PHARMALGEN YELLOW HORNET VENOM PROTEIN ALK

DOLICHOVESPULA MACULATA VENOM PROTEIN EXTRACT

120MCG POWDER FOR SOLUTION
01949004 PHARMALGEN WHITE FACED HORNET VENOM ALK

HONEY BEE VENOM PROTEIN EXTRACT

120MCG POWDER FOR SOLUTION
02226197 VENOMIL HONEY BEE VENOM JUB

550MCG POWDER FOR SOLUTION
02220075 HYMENOPTERA VENOM PRODUCT HONEY BEE VENOM JUB

92:05.00 SERUMS

NON POLLEN

100,000U LIQUID
00299979 ALLERGENIC EXTRACT NON POLLENS ALK

POLISTES SPP VENOM PROTEIN EXTRACT

1.1MG POWDER FOR SOLUTION
01948970 PHARMALGEN WASP VENOM PROTEIN ALK

POLLEN

4,300U/ML LIQUID
00464988 POLLINEX R BEN

100,000U LIQUID
00299987 ALLERGENIC EXTRACT POLLENS ALK

POLLEN AND NON POLLEN

20,000U LIQUID
00648922 CENTER-AL ALK

VENOM PROTEIN EXTRACT

3,300MCG POWDER FOR SOLUTION
01948873 PHARMALGEN MIXED VESPID VENOM PROTEIN ALK

VESPUA SPP VENOM PROTEIN EXTRACT

1.1MG POWDER FOR SOLUTION
01948954 PHARMALGEN YELLOW JACKET VENOM PROTEIN ALK

120MCG POWDER FOR SOLUTION
01948962 PHARMALGEN YELLOW JACKET VENOM PROTEIN ALK

WASP VENOM PROTEIN

120MCG POWDER FOR SOLUTION
02226219 VENOMIL WASP VENOM PROTEIN JUB

550MCG POWDER FOR SOLUTION
02220091 HYMENOPTERA VENOM PRODUCT WASP VENOM PROTEIN JUB

WHITE FACED HORNET VENOM PROTEIN

120MCG POWDER FOR SOLUTION
02226235 VENOMIL WHITE-FACED HORNET VENOM PROTEIN JUB

WHITE FACED HORNET VENOM PROTEIN, YELLOW HORNET VENOM PROTEIN, YELLOW JACKET VENOM PROTEIN

120MCG POWDER FOR SOLUTION
01948881 PHARMALGEN MIXED VESPID VENOM PROTEIN ALK

02226294 VENOMIL MIXED VESPID VENOM PROTEIN JUB

550MCG POWDER FOR SOLUTION
02221314 HYMENOPTERA VENOM PRODUCT MIXED VESPID VENOM PROTEIN JUB

YELLOW HORNET VENOM PROTEIN

120MCG/ML POWDER FOR SOLUTION
02226251 VENOMIL YELLOW HORNET VENOM PROTEIN JUB

92:05.00 SERUMS

YELLOW HORNET VENOM PROTEIN

550MCG POWDER FOR SOLUTION

02220083	HYMENOPTERA VENOM PRODUCTS YELLOW HORNET VENOM PROTEIN	JUB
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YELLOW JACKET VENOM PROTEIN

120MCG POWDER FOR SOLUTION

02226286	VENOMIL YELLOW JACKET VENOM PROTEIN	JUB
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550MCG POWDER FOR SOLUTION

02220113	HYMENOPTERA VENOM PRODUCT YELLOW JACKET VENOM PROTEIN	JUB
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92:08.00 5 ALFA REDUCTASE INHIBITORS

DUTASTERIDE

ST **0.5MG CAPSULE**

02412691	ACT DUTASTERIDE	TEV
02404206	APO-DUTASTERIDE	APX
02469308	AURO-DUTASTERIDE	AUR
02247813	AVODART	GSK
02421712	DUTASTERIDE	PDL
02429012	DUTASTERIDE	SIV
02443058	DUTASTERIDE	SAN
02416298	MED-DUTASTERIDE	GMP
02428873	MINT-DUTASTERIDE	MIN
02393220	PMS-DUTASTERIDE	PMS
02427753	RIVA-DUTASTERIDE	RIV
02424444	SANDOZ DUTASTERIDE	SDZ
02408287	TEVA-DUTASTERIDE	TEV

FINASTERIDE

ST **5MG TABLET**

02355043	ACH-FINASTERIDE	ACC
02354462	ACT FINASTERIDE	ACG
02365383	APO-FINASTERIDE	APX
02405814	AURO-FINASTERIDE	AUR
02376709	DOM-FINASTERIDE	DPC
02350270	FINASTERIDE	PDL
02445077	FINASTERIDE	SAN
02447541	FINASTERIDE	SIV
02357224	JAMP-FINASTERIDE	JMP
02389878	MINT-FINASTERIDE	MIN
02310112	PMS-FINASTERIDE	PMS
02010909	PROSCAR	FRS
02371820	RAN-FINASTERIDE	RBY
02455013	RIVA-FINASTERIDE	RIV
02322579	SANDOZ FINASTERIDE	SDZ
02348500	TEVA-FINASTERIDE	TEV
02428741	VAN-FINASTERIDE	VAN

92:12.00 ANTIDOTES

LEUCOVORIN CALCIUM

5MG TABLET

02170493	LEDERLE LEUCOVORIN	PFI
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92:16.00 ANTIGOUT AGENTS

ALLOPURINOL

ST **100MG TABLET**

00555681	ALLOPURINOL	PDL
02402769	APO-ALLOPURINOL	APX
02421593	JAMP-ALLOPURINOL	JMP
02396327	MAR-ALLOPURINOL	MAR
00402818	ZYLOPRIM	AAP

ST **200MG TABLET**

02130157	ALLOPURINOL	PDL
02402777	APO-ALLOPURINOL	APX
02421607	JAMP-ALLOPURINOL	JMP
02396335	MAR-ALLOPURINOL	MAR
00479799	ZYLOPRIM	AAP

ST **300MG TABLET**

00294322	ALLOPURINOL	APX
00555703	ALLOPURINOL	PDL
02402785	APO-ALLOPURINOL	APX
02421615	JAMP-ALLOPURINOL	JMP
02396343	MAR-ALLOPURINOL	MAR
00402796	ZYLOPRIM	AAP

ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503018	ALLOPURINOL ORAL LIQUID	UNK
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COLCHICINE

ST **0.6MG TABLET**

00572349	COLCHICINE	ODN
02373823	JAMP-COLCHICINE	JMP
02402181	PMS-COLCHICINE	PMS
00287873	SANDOZ COLCHICINE	SDZ

FEBUXOSTAT

Limited use benefit (prior approval required).

For patients with symptomatic gout who have documented hypersensitivity to allopurinol.

ST **80MG TABLET**

02357380	ULORIC	TAK
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92:20.00 IMMUNOMODULATORY AGENTS

GLATIRAMER ACETATE

Limited use benefit (prior approval required).

• As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

20MG SOLUTION

02245619	COPAXONE	TEV
02460661	GLATECT	PMS

92:20.00 IMMUNOMODULATORY AGENTS

INTERFERON BETA-1A

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

30MCG INJECTION

09857395	AVONEX PEN	UNK
99100763	AVONEX PEN	UNK

60MCG POWDER FOR SOLUTION

02267594	AVONEX	UNK
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22MCG SOLUTION

02237319	REBIF	SRO
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30MCG SOLUTION

02269201	AVONEX	UNK
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44MCG SOLUTION

02237318	REBIF	SRO
02237320	REBIF	SRO

66MCG SOLUTION

02318253	REBIF	SRO
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132MCG SOLUTION

02318261	REBIF	SRO
02318288	REBIF	SRO

INTERFERON BETA-1B

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

0.3MG INJECTION

99100555	BETASERON INITIATION KIT	BAY
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0.3MG POWDER FOR SOLUTION

02169649	BETASERON	BAY
02337819	EXTAVIA	NVR

92:20.00 IMMUNOMODULATORY AGENTS

OCRELIZUMAB

Limited use benefit (prior approval required).

1. For the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence who meet all of the following criteria:

- Prescribed by a neurologist experienced in the management of RRMS; AND
- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids. Expanded Disability Status Scale score (EDSS) of 5.5 or less.
- Patient is 18 years of age or older.

a. A clinical relapse is defined as the appearance of new symptoms or worsening of old symptoms, lasting at least 24 hours in the absence of fever, preceded by stability for at least one month.

b. MRI activity is defined as any new multiple sclerosis lesion/s, expanding lesion/s, and/or enhancing lesion/s.

OR

2. For the treatment of primary progressive multiple sclerosis (PPMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence who meet all of the following criteria:

- Initial Coverage (one year)
- Prescribed by a neurologist experienced in the management of PPMS; AND
- Expanded Disability Status Scale (EDSS) between 3.0 and 6.5; AND
- Score of at least 2.0 on the Functional Systems scale (FSS) for the pyramidal system due to lower extremity findings; AND
- Disease duration of less than 15 years for those with an EDSS greater than 5.0 or less than 10 years for those with an EDSS of 5.0 or less; AND
- Patient is 18 years of age or older.

Renewal Coverage for PPMS (one year):

- EDSS of less than 7.0.

30MG SOLUTION

02467224	OCREVUS	HLR
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TERIFLUNOMIDE

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

14MG TABLET

02416328	AUBAGIO	GEE
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92:24.00 BONE RESORPTION INHIBITORS

ALENDRONATE SODIUM

ST **5MG TABLET**

02381478	ACH-ALENDRONATE	ACC
02248727	APO-ALENDRONATE	APX

92:24.00 BONE RESORPTION INHIBITORS

ALENDRONATE SODIUM

ST **5MG TABLET**

02384698	RAN-ALENDRONATE	RBY
02248251	TEVA-ALENDRONATE	TEV
02428717	VAN-ALENDRONATE	VAN

ST **10MG TABLET**

02381486	ACH-ALENDRONATE	ACC
02248728	APO-ALENDRONATE	APX
02388545	AURO-ALENDRONATE	AUR
02384701	RAN-ALENDRONATE	RBY
02288087	SANDOZ ALENDRONATE	SDZ
02247373	TEVA-ALENDRONATE	TEV
02428725	VAN-ALENDRONATE	VAN

ST **40MG TABLET**

02258102	ACT ALENDRONATE	ACG
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ST **70MG TABLET**

02381494	ACH-ALENDRONATE	ACC
02299712	ALENDRONATE	SIV
02352966	ALENDRONATE	SAN
02303078	ALENDRONATE-70	PDL
02248730	APO-ALENDRONATE	APX
02388553	AURO-ALENDRONATE	AUR
02282763	DOM-ALENDRONATE	DPC
02245329	FOSAMAX	FRS
02385031	JAMP-ALENDRONATE	JMP
02394871	MINT-ALENDRONATE	MIN
02273179	PMS-ALENDRONATE	PMS
02284006	PMS-ALENDRONATE	PMS
02384728	RAN-ALENDRONATE	RBY
02270889	RIVA-ALENDRONATE	RIV
02288109	SANDOZ ALENDRONATE	SDZ
02261715	TEVA-ALENDRONATE	TEV
02428733	VAN-ALENDRONATE	VAN

ALENDRONATE SODIUM, CHOLECALCIFEROL

ST **70MG & 2,800U TABLET**

02454467	APO-ALENDRONATE/VITAMIN D3	APX
02276429	FOSAVANCE	FRS
02403633	TEVA-ALENDRONATE/CHOLECALCIFEROL	TEV

ST **70MG & 5,600U TABLET**

02454475	APO-ALENDRONATE/VITAMIN D3	APX
02314940	FOSAVANCE	FRS
02429160	SANDOZ ALENDRONATE/CHOLECALCIFEROL	SDZ
02403641	TEVA-ALENDRONATE/CHOLECALCIFEROL	TEV

92:24.00 BONE RESORPTION INHIBITORS

DENOSUMAB (PROLIA)

Limited use benefit (prior approval required).

For the treatment of osteoporosis in patients who have a significant fracture risk defined as either:
 - moderate 10-year fracture risk (10% to 20%) with a prior fragility fracture; OR
 - high 10-year fracture risk ($\geq 20\%$);
AND
 - Have a contraindication to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment);
OR
 - Have failed or have an intolerance to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment).

60MG/ML SOLUTION

02343541	PROLIA	AMG
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DENOSUMAB (XGEVA)

Limited use benefit (prior approval required).

For the prevention of skeletal-related events (SREs) in patients with castrate-resistant prostate cancer (CRPC) with:
 • one or more documented bone metastases; **AND**
 • good performance status (ECOG performance status score of 0, 1, or 2).

120MG/1.7ML SOLUTION

02368153	XGEVA	AMG
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ETIDRONATE DISODIUM

ST **200MG TABLET**

02248686	ACT ETIDRONATE	ACG
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ETIDRONATE DISODIUM, CALCIUM CARBONATE

ST **400MG & 500MG TABLET**

02263866	ACT ETIDROCAL	ACG
02324199	NOVO-ETIDRONATECAL	TEV

PAMIDRONATE DISODIUM

6MG SOLUTION

02249677	PAMIDRONATE	OMG
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9MG SOLUTION

02246599	PAMIDRONATE	FKD
02249685	PAMIDRONATE DISODIUM OMEGA	OMG

30MG SOLUTION

02264951	PAMIDRONATE	SDZ
02244550	PAMIDRONATE DISODIUM	PFI

60MG SOLUTION

02264978	PAMIDRONATE	SDZ
02244551	PAMIDRONATE DISODIUM	PFI

90MG SOLUTION

02264986	PAMIDRONATE	SDZ
02244552	PAMIDRONATE DISODIUM	PFI
02245999	PMS-PAMIDRONATE	PMS

RISEDRONATE SODIUM

ST **5MG TABLET**

02298376	TEVA-RISEDRONATE	TEV
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ST **30MG TABLET**

02298384	TEVA-RISEDRONATE	TEV
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ST **35MG TABLET**

02246896	ACTONEL	ALL
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92:24.00 BONE RESORPTION INHIBITORS

RISEDRONATE SODIUM

ST 35MG TABLET

02353687	APO-RISEDRONATE	APX
02406306	AURO-RISEDRONATE	AUR
02309831	DOM-RISEDRONATE	DPC
02368552	JAMP-RISEDRONATE	JMP
02302209	PMS-RISEDRONATE	PMS
02347474	RISEDRONATE	PDL
02370255	RISEDRONATE	SAN
02411407	RISEDRONATE-35	SIV
02341077	RIVA-RISEDRONATE	RIV
02327295	SANDOZ RISEDRONATE	SDZ
02298392	TEVA-RISEDRONATE	TEV

ST 150MG TABLET

02316838	ACTONEL	ALL
02377721	APO-RISEDRONATE	APX
02424177	PMS-RISEDRONATE	PMS
02413809	TEVA-RISEDRONATE	TEV

ZOLEDRONIC ACID MONOHYDRATE

Limited use benefit (prior approval required).

Maximum dose covered is 5mg per 12-month period

For the treatment of Paget's disease;

OR

For the treatment of osteoporosis in patients who have a significant fracture risk defined as either:

- moderate 10-year fracture risk (10% to 20%) with a prior fragility fracture; OR

- high 10-year fracture risk (≥ 20%)

AND

- Have a contraindication to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment);OR

- Have failed or have an intolerance to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment).

5MG/100ML SOLUTION

02269198	ACLASTA	NVR
02415100	TARO-ZOLEDRONIC ACID	TAR
02408082	ZOLEDRONIC ACID	TEV
02422433	ZOLEDRONIC ACID	REC

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS

ABATACEPT

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Polyarticular juvenile idiopathic arthritis (pJIA) according to established criteria.

(Please refer to Appendix A).

250MG POWDER FOR SOLUTION

02282097	ORENCIA	BMS
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125MG SOLUTION

02402475	ORENCIA	BMS
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92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS

ADALIMUMAB

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Psoriatic Arthritis according to established criteria.
- Ankylosing Spondylitis according to established criteria.
- Psoriasis according to established criteria.
- Crohn's disease according to established criteria.
- Polyarticular juvenile idiopathic arthritis (pJIA) according to established criteria.
- Ulcerative colitis according to established criteria.
- Hidradenitis Suppurativa according to established criteria.

(Please refer to Appendix A).

40MG/VIAL SOLUTION

02258595	HUMIRA	ABV
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CERTOLIZUMAB PEGOL

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Psoriatic arthritis according to established criteria.
- Ankylosing spondylitis according to established criteria.

(Please refer to Appendix A).

200MG SOLUTION

02465574	CIMZIA	UCB
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200MG/ML SOLUTION

02331675	CIMZIA	UCB
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ETANERCEPT

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Psoriatic Arthritis according to established criteria.
- Ankylosing Spondylitis according to established criteria.
- Polyarticular juvenile idiopathic arthritis (pJIA) according to established criteria.

(Please refer to Appendix A).

25MG/VIAL INJECTION

02242903	ENBREL	PED
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50MG/ML INJECTION

02274728	ENBREL	PED
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99100373	ENBREL SURECLICK	AMG
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ETANERCEPT (BRENZYS)

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Ankylosing Spondylitis according to established criteria.

(Please refer to Appendix A).

50MG SOLUTION

02455323	BRENZYS	UNK
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02455331	BRENZYS	UNK
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**92:36.00 DISEASE-MODIFYING
ANTIRHEUMATIC AGENTS**

ETANERCEPT (ERELZI)

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Psoriatic Arthritis according to established criteria.
- Ankylosing Spondylitis according to established criteria.
- Polyarticular juvenile idiopathic arthritis (pJIA) according to established criteria.

(Please refer to Appendix A).

25MG SOLUTION

02462877 ERELZI SDZ

50MG SOLUTION

02462850 ERELZI SDZ

02462869 ERELZI SDZ

GOLIMUMAB

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Psoriatic Arthritis according to established criteria.
- Ankylosing Spondylitis according to established criteria.
- Ulcerative colitis according to established criteria.

(Please refer to Appendix A).

50MG/0.5ML SOLUTION

02324776 SIMPONI JSO

02324784 SIMPONI JSO

100MG/ML SOLUTION

02413175 SIMPONI JSO

02413183 SIMPONI JSO

INFLIXIMAB (INFLECTRA)

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Psoriatic Arthritis according to established criteria.
- Ankylosing Spondylitis according to established criteria.
- Psoriasis according to established criteria.
- Crohn's disease according to established criteria.
- Fistulizing Crohn's disease according to established criteria.
- Ulcerative colitis according to established criteria.

(Please refer to Appendix A).

100MG POWDER FOR SOLUTION

02419475 INFLECTRA HOS

02470373 RENFLEXIS UNK

INFLIXIMAB (REMICADE)

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Crohn's disease according to established criteria.
- Fistulizing Crohn's disease according to established criteria.

(Please refer to Appendix A).

100MG/VIAL POWDER FOR SOLUTION

02244016 REMICADE JSO

**92:36.00 DISEASE-MODIFYING
ANTIRHEUMATIC AGENTS**

LEFLUNOMIDE

ST **10MG TABLET**

02256495 APO-LEFLUNOMIDE APX

02241888 ARAVA SAC

02351668 LEFLUNOMIDE SAN

02415828 LEFLUNOMIDE PDL

02288265 PMS-LEFLUNOMIDE PMS

02283964 SANDOZ LEFLUNOMIDE SDZ

02261251 TEVA-LEFLUNOMIDE TEV

ST **20MG TABLET**

02256509 APO-LEFLUNOMIDE APX

02241889 ARAVA SAC

02351676 LEFLUNOMIDE SAN

02415836 LEFLUNOMIDE PDL

02288273 PMS-LEFLUNOMIDE PMS

02283972 SANDOZ LEFLUNOMIDE SDZ

02261278 TEVA-LEFLUNOMIDE TEV

TOCILIZUMAB (IV)

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Systemic juvenile idiopathic arthritis (sJIA) according to established criteria.
- Polyarticular juvenile idiopathic arthritis (pJIA) according to established criteria.

(Please refer to Appendix A).

80MG/4ML SOLUTION

02350092 ACTEMRA HLR

200MG/10ML SOLUTION

02350106 ACTEMRA HLR

400MG/20ML SOLUTION

02350114 ACTEMRA HLR

TOCILIZUMAB (SC)

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.
- Giant Cell Arteritis according to established criteria.

(Please refer to Appendix A).

162MG SOLUTION

02424770 ACTEMRA HLR

TOFACITINIB CITRATE

Limited use benefit (prior approval required).

For the treatment of:

- Rheumatoid Arthritis according to established criteria.

(Please refer to Appendix A).

5MG TABLET

02423898 XELJANZ PFI

92:44.00 IMMUNOSUPPRESSIVE AGENTS

ALEMTUZUMAB

Limited use benefit (prior approval required).

Coverage is provided for two years (i.e. two treatment courses/eight doses) for adult patients who meet ALL of the following criteria:

- For the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence; AND
- Prescribed by a specialist with experience in the treatment of multiple sclerosis; AND
- Active disease defined by clinical and imaging features (i.e. significant increase in T2 lesion load compared with that from a previous MRI scan OR at least one gadolinium-enhancing lesion); AND
- Failure to respond to full and adequate courses of at least ONE initial trial of at least 6 months of interferon, glatiramer, dimethyl fumarate or teriflunomide therapy OR documented intolerance to at least 2 therapies; AND
- At least one relapse while on at least six months of a disease modifying therapy (an interferon, glatiramer, dimethyl fumarate or teriflunomide) within the last 10 years; AND
- At least two attacks (first episode or relapse) in the previous two years, with at least one attack in the previous year; AND
- An Expanded Disability Status Scale (EDSS) score of five (5) or less.

12MG SOLUTION

02418320 LEMTRADA GEE

AZATHIOPRINE

ST **50MG TABLET**

02242907 APO-AZATHIOPRINE APX
 02243371 AZATHIOPRINE-50 PDL
 00004596 IMURAN ASP
 02236819 TEVA-AZATHIOPRINE TEV

ST **PDIN FOR EXTEMPORANEOUS MIXTURE**

99503019 AZATHIOPRINE ORAL LIQUID UNK

CYCLOSPORINE

Limited use benefit (prior approval required).

For transplant therapy.

ST **10MG CAPSULE**

02237671 NEORAL NVR

ST **25MG CAPSULE**

02150689 NEORAL NVR
 02247073 SANDOZ CYCLOSPORINE SDZ

ST **50MG CAPSULE**

02150662 NEORAL NVR
 02247074 SANDOZ CYCLOSPORINE SDZ

ST **100MG CAPSULE**

02150670 NEORAL NVR
 02242821 SANDOZ CYCLOSPORINE SDZ

ST **100MG/ML SOLUTION**

02244324 APO-CYCLOSPORINE APX
 02150697 NEORAL NVR

92:44.00 IMMUNOSUPPRESSIVE AGENTS

FINGOLIMOD (FINGOLIMOD HYDROCHLORIDE)

Limited use benefit (prior approval required).

Initial Coverage (one year):

For the treatment of patients with Relapsing Remitting Multiple Sclerosis (RRMS) who meet ALL of the following criteria:

- Failure to respond to full and adequate courses of at least ONE initial disease-modifying therapy (an interferon, glatiramer acetate, dimethyl fumarate or teriflunomide) OR documented intolerance to at least 2 therapies; AND
- One or more clinically disabling relapses in the previous year; AND
- Significant increase in T2 lesion load compared with that from a previous MRI scan or at least one gadolinium-enhancing lesion; AND
- Requested and followed by a neurologist experienced in the management of RRMS; AND
- Recent Expanded Disability Status Scale (EDSS) score.

Renewal Coverage (two years):

- EDSS scores must be provided (exam must have occurred within that last 90 days).
- Patients must be stable or have experienced no more than 1 disabling attack/relapse in the past year.

0.5MG CAPSULE

02365480 GILENYA NVR

MEPOLIZUMAB

Limited use benefit (prior approval required).

For initial 12-month coverage:

For the adjunctive treatment of severe eosinophilic asthma in adults who are inadequately controlled with high-dose inhaled corticosteroids plus one or more additional asthma controller(s) (e.g. long-acting beta-agonist); AND

- Have had a blood eosinophil count of $\geq 0.15 \times 10^9/L$ before initiation of Nucala (levels must have been drawn within 3 months of the start of treatment); OR
- Have had a blood eosinophil count of $\geq 0.3 \times 10^9/L$ within the 12-month period prior to starting Nucala; AND
- Show reversibility on spirometry (a rise in FEV1 of at least 12% AND at least 200 mL); AND
- Have experienced two or more clinically significant asthma exacerbations* in the past 12 months period prior to starting Nucala; or
- Have received maintenance therapy with daily oral corticosteroids for at least 3 months prior to starting Nucala.

For 12-month renewal coverage:

- Patient has experienced a decrease in clinically significant asthma exacerbations* with Nucala treatment; OR
- Patient's oral corticosteroid maintenance dose decreased by at least 25 % from the pre-treatment dose.

Coverage for Nucala is provided for a maximum dose of 100 mg every four weeks.

* A clinically significant asthma exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least three days or the patient visited an emergency department or was hospitalized.

100MG POWDER FOR SOLUTION

02449781 NUCALA GSK

92:44.00 IMMUNOSUPPRESSIVE AGENTS

MYCOPHENOLATE MOFETIL

Limited use benefit (prior approval required).

For transplant therapy.

ST **250MG CAPSULE**

02383780	ACH-MYCOPHENOLATE	ACC
02352559	APO-MYCOPHENOLATE	APX
02192748	CELLCEPT	HLR
02386399	JAMP-MYCOPHENOLATE	JMP
02457369	MYCOPHENOLATE MOFETIL	SAN
02371154	MYLAN-MYCOPHENOLATE	MYL
02320630	SANDOZ MYCOPHENOLATE	SDZ
02364883	TEVA-MYCOPHENOLATE	TEV
02433680	VAN-MYCOPHENOLATE	VAN

ST **200MG POWDER FOR SUSPENSION**

02242145	CELLCEPT	HLR
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ST **500MG TABLET**

02352567	APO-MYCOPHENOLATE	APX
02237484	CELLCEPT	HLR
02380382	JAMP-MYCOPHENOLATE	JMP
02378574	MYCOPHENOLATE	ACC
02457377	MYCOPHENOLATE MOFETIL	SAN
02370549	MYLAN-MYCOPHENOLATE	MYL
02313855	SANDOZ MYCOPHENOLATE	SDZ
02348675	TEVA-MYCOPHENOLATE	TEV
02432625	VAN-MYCOPHENOLATE	VAN

MYCOPHENOLATE SODIUM

Limited use benefit (prior approval required).

For transplant therapy.

ST **180MG TABLET (ENTERIC COATED)**

02372738	APO-MYCOPHENOLIC ACID	APX
02264560	MYFORTIC	NVR

ST **360MG TABLET (ENTERIC COATED)**

02372746	APO-MYCOPHENOLIC ACID	APX
02264579	MYFORTIC	NVR

SIROLIMUS

Limited use benefit (prior approval required).

Coverage will be provided as a second line therapy for patients failing mycophenolate mofetil.

ST **1MG/ML SOLUTION**

02243237	RAPAMUNE	PFI
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ST **1MG TABLET**

02247111	RAPAMUNE	PFI
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TACROLIMUS MONOHYDRATE

Limited use benefit (prior approval required).

For transplant therapy.

ST **0.5MG CAPSULE**

02243144	PROGRAF	AST
02416816	SANDOZ TACROLIMUS	SDZ

ST **1MG CAPSULE**

02175991	PROGRAF	AST
02416824	SANDOZ TACROLIMUS	SDZ

92:44.00 IMMUNOSUPPRESSIVE AGENTS

TACROLIMUS MONOHYDRATE

Limited use benefit (prior approval required).

For transplant therapy.

ST **5MG CAPSULE**

02175983	PROGRAF	AST
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ST **0.5MG CAPSULE (EXTENDED RELEASE)**

02296462	ADVAGRAF	AST
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ST **1MG CAPSULE (EXTENDED RELEASE)**

02296470	ADVAGRAF	AST
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ST **3MG CAPSULE (EXTENDED RELEASE)**

02331667	ADVAGRAF	AST
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ST **5MG CAPSULE (EXTENDED RELEASE)**

02296489	ADVAGRAF	AST
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ST **5MG CAPSULE (IMMEDIATE RELEASE)**

02416832	SANDOZ TACROLIMUS	SDZ
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5MG/ML SOLUTION

02176009	PROGRAF	AST
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VEDOLIZUMAB

Limited use benefit (prior approval required).

For the treatment of:

- Crohn's disease according to established criteria.
- Ulcerative colitis according to established criteria.

(Please refer to Appendix A).

300MG POWDER FOR SOLUTION

02436841	ENTYVIO	TAK
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92:92.00 OTHER MISCELLANEOUS THERAPEUTIC AGENTS

ABOBOTULINUMTOXINA

Limited use benefit (prior approval required).

Treatment of cervical dystonia (spasmodic torticollis) in adults; OR
Symptomatic treatment of focal spasticity affecting upper limbs in adults; OR
Lower limb spasticity in patients 2 years of age and older.

300U POWDER FOR SOLUTION

02460203	DYSPORT THERAPEUTIC	IPS
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500U POWDER FOR SOLUTION

02456117	DYSPORT THERAPEUTIC	IPS
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CYPROTERONE ACETATE

50MG TABLET

00704431	ANDROCUR	BAY
02245898	CYPROTERONE	AAP
02390760	MED-CYPROTERONE	GMP
02395797	RIVA-CYPROTERONE	RIV

CYPROTERONE ACETATE, ETHINYL ESTRADIOL

2MG & 35MCG TABLET

02290308	CYESTRA-35	PAL
02233542	DIANE-35	BAY
02425017	RAN-CYPROTERONE/ETHINYL ESTRADIOL	RBY
02309556	TEVA-CYPROTERONE / ETHINYL ESTRADIOL	TEV

**92:92.00 OTHER MISCELLANEOUS
THERAPEUTIC AGENTS**

INCOBOTULINUMTOXINA

Limited use benefit (prior approval required).

For the treatment of:

- strabismus and blepharospasm associated with dystonia, including benign essential blepharospasm or VII nerve disorder in patients 12 years of age or older; OR
- cervical dystonia (spasmodic torticollis).

50UNIT/VIAL POWDER FOR SOLUTION

02371081 XEOMIN MEZ

100U/VIAL POWDER FOR SOLUTION

02324032 XEOMIN MEZ

LANREOTIDE ACETATE

60MG/0.3ML SOLUTION (EXTENDED RELEASE)

02283395 SOMATULINE AUTOGEL IPS

90MG/0.3ML SOLUTION (EXTENDED RELEASE)

02283409 SOMATULINE AUTOGEL IPS

120MG/0.5ML SOLUTION (EXTENDED RELEASE)

02283417 SOMATULINE AUTOGEL IPS

ONABOTULINUMTOXINA

Limited use benefit (prior approval required).

For the treatment of:

- strabismus and blepharospasm associated with dystonia, including benign essential blepharospasm or VII nerve disorder in patients 12 years of age or older; OR
- cervical dystonia (spasmodic torticollis); OR
- urinary incontinence due to neurogenic detrusor over activity resulting from neurogenic bladder associated with MS or subcervical spinal cord injury; OR
- overactive bladder.

50IU INJECTION

09857386 BOTOX ALL

200IU INJECTION

09857387 BOTOX ALL

100IU POWDER FOR SOLUTION

01981501 BOTOX ALL

94:00 DEVICES

94:00.00 DEVICES

SPACER DEVICE

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 2 spacer devices every 12 months.

DEVICE

96899962	AEROCHAMBER AC BOYZ	TRU
96899963	AEROCHAMBER AC GIRLZ	TRU
96899969	AEROCHAMBER PLUS FLOWVU LARGE	TRU
96899970	AEROCHAMBER PLUS FLOWVU MEDIUM	TRU
96899968	AEROCHAMBER PLUS FLOWVU MOUTH	TRU
96899971	AEROCHAMBER PLUS FLOWVU SMALL	TRU
96899977	AEROTRACH PLUS	UNK
96899956	COMPACT SPACE PLUS LARGE MASK	MIN
96899955	COMPACT SPACE PLUS MEDIUM MASK	MIN
96899953	COMPACT SPACE PLUS NO MASK	MIN
96899954	COMPACT SPACE PLUS SMALL MASK	MIN
99400507	E-Z SPACER	WEP
99400511	E-Z SPACER (MASK ONLY)	WEP
99400508	E-Z SPACER WITH SMALL MASK	WEP
00901012	INSPIRA CHAMBER W LARGE MASK	LUP
00900003	INSPIRA CHAMBER W MEDIUM MASK	LUP
00900001	INSPIRA CHAMBER W MOUTHPIECE	LUP
00900002	INSPIRA CHAMBER W SMALL MASK	LUP
99400501	OPTICHAMBER	AUC
96899961	OPTICHAMBER DIAMOND (CHAMBER)	AUC
96899958	OPTICHAMBER DIAMOND LARGE MASK	AUC
96899959	OPTICHAMBER DIAMOND MEDIUM MASK	AUC
96899960	OPTICHAMBER DIAMOND SMALL MASK	AUC
99400504	OPTICHAMBER LARGE MASK	AUC
99400503	OPTICHAMBER MEDIUM MASK	AUC
99400502	OPTICHAMBER SMALL MASK	AUC
99400505	OPTIHALER	AUC
99400787	POCKET CHAMBER	MCA
99400791	POCKET CHAMBER WITH ADULT MASK	MCA
99400788	POCKET CHAMBER WITH INFANT MASK	MCA
99400790	POCKET CHAMBER WITH MEDIUM MASK	MCA
99400789	POCKET CHAMBER WITH SMALL MASK	MCA
96899974	RESPICHAMBER SILICONE MEDIUM MASK	TRU

94:00.00 DEVICES

SPACER DEVICE

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 2 spacer devices every 12 months.

DEVICE

96899973	RESPICHAMBER SILICONE SMALL MASK	TRU
96899972	RESPICHAMBER VHC W MOUTHPIECE	TRU

94:01.00 DEVICES (DIABETIC)

ADHESHIVE WIPES

MISCELLANEOUS

97799671	SKIN PREP ADHESHIVE WIPES	UNK
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DRESSING

DRESS

99401078	SN IV3000 1-HAND TRANS	SMW
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INSULIN PUMP SUPPLIES

Limited use benefit (prior approval required).

- Insulin pump supplies are approved for NIHB clients following the approval of an insulin pump by NIHB; OR
- Insulin pump supplies are approved for NIHB clients with Type 1 diabetes if an insulin pump was partially or totally covered by another insurance.

DEVICE

97799674	CARTRIDGE FOR IR200	UNK
97799342	INSET 30 INFUSION SETS	UNK
99401038	INSULIN PUMP BATTERY	AUC
09991458	IV3000	SMW

COMFORT ANGLED DEVICE

97799682	COMFORT ANGLED INFSET 17MM	UNK
97799683	COMFORT ANGLED INFSET 17MM	UNK

COMFORT SHORT ANGLED DEVICE

97799678	COMFORT SRT ANGLED INFSET 13	UNK
97799679	COMFORT SRT ANGLED INFSET 13	UNK

CONTACT DETACH DEVICE

97799672	CONTACT DETACH 90 DEGREE 6MMX60CM	UNK
97799610	CONTACT DETACH 90 DEGREE 8MMX60CM	UNK

INSET II DEVICE

97799685	INSET II 90 DEGREE 6MMX110CM	UNK
97799687	INSET II 90 DEGREE 6MMX60CM	UNK
97799684	INSET II 90 DEGREE 9MMX110CM	UNK
97799686	INSET II 90 DEGREE 9MMX60CM	UNK

MIO DEVICE

97799491	MIO BLUE 6MMX18	MDT
97799438	MIO BLUE 6MMX23	MDT
97799490	MIO CLEAR 6MMX32	MDT
97799489	MIO CLEAR 9MMX32	MDT
97799492	MIO PINK 6MMX18	MDT
97799437	MIO PINK 6MMX23	MDT

OMNIPOD DEVICE

09991327	PODS	UNK
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94:01.00 DEVICES (DIABETIC)

INSULIN PUMP SUPPLIES

Limited use benefit (prior approval required).

- Insulin pump supplies are approved for NIHB clients following the approval of an insulin pump by NIHB; OR
- Insulin pump supplies are approved for NIHB clients with Type 1 diabetes if an insulin pump was partially or totally covered by another insurance.

PARADIGM SILHOUETTE DEVICE

97799715	PARADIGM SILHOUETTE 13MMX 43	MDT
97799485	PARADIGM SILHOUETTE 13MMX18"	MDT
97799716	PARADIGM SILHOUETTE 13MMX23	MDT
97799484	PARADIGM SILHOUETTE 13MMX32"	MDT
97799718	PARADIGM SILHOUETTE 17MMX23	MDT
97799483	PARADIGM SILHOUETTE 17MMX32"	MDT
97799719	PARADIGM SILHOUETTE 17MMX43	MDT
97799529	PARADIGM SILHOUETTE CANNULA 13MM	MDT
97799528	PARADIGM SILHOUETTE CANNULA 17MM	MDT

QUICK-SET DEVICE

97799486	QUICK-SET 6MMX18	MDT
97799744	QUICK-SET 6MMX23 TUBING	MDT
97799487	QUICK-SET 6MMX32	MDT
97799743	QUICK-SET 6MMX43 TUBING	MDT
97799742	QUICK-SET 9MMX23 TUBING	MDT
97799488	QUICK-SET 9MMX32	MDT
97799741	QUICK-SET 9MMX43 TUBING	MDT

RAPID-D DEVICE

97799650	RAPID-D 10MM/110CM	ROD
97799652	RAPID-D 10MM/60CM	ROD
97799651	RAPID-D 10MM/80CM	ROD
97799656	RAPID-D 6MM/110CM	ROD
97799658	RAPID-D 6MM/60CM	ROD
97799657	RAPID-D 6MM/80CM	ROD
97799653	RAPID-D 8MM/110CM	ROD
97799655	RAPID-D 8MM/60CM	ROD
97799654	RAPID-D 8MM/80CM	ROD

SURE-T DEVICE

97799521	PARADIGM SURE-T 29G 6MMX18	MDT
97799520	PARADIGM SURE-T 29G 6MMX23	MDT
97799519	PARADIGM SURE-T 29G 8MMX23	MDT

TENDER DEVICE

97799644	TENDER-1 17MM/110CM	ROD
97799646	TENDER-1 17MM/60CM	ROD
97799645	TENDER-1 17MM/80CM	ROD
97799638	TENDER-2 17MM/110CM	ROD
97799640	TENDER-2 17MM/60CM	ROD
97799639	TENDER-2 17MM/80CM	ROD

TENDER "MINI" DEVICE

97799647	TENDER-1 MINI INF SET 13MM/110CM	ROD
97799649	TENDER-1 MINI INFSET 13MM/60CM	ROD
97799648	TENDER-1 MINI INFSET 13MM/80CM	ROD

94:01.00 DEVICES (DIABETIC)

INSULIN PUMP SUPPLIES

Limited use benefit (prior approval required).

- Insulin pump supplies are approved for NIHB clients following the approval of an insulin pump by NIHB; OR
- Insulin pump supplies are approved for NIHB clients with Type 1 diabetes if an insulin pump was partially or totally covered by another insurance.

TENDER "MINI" DEVICE

97799641	TENDER-2 MINI INF SET 13MM/110CM	ROD
97799643	TENDER-2 MINI INFSET 13MM/60CM	ROD
97799642	TENDER-2 MINI INFSET 13MM/80CM	ROD

ULTRAFLEX DEVICE

97799665	ULTRAFLEX 1 10MM/110CM	ROD
97799667	ULTRAFLEX 1 10MM/60CM	ROD
97799666	ULTRAFLEX 1 10MM/80CM	ROD
97799668	ULTRAFLEX 1 8MM/110CM	ROD
97799670	ULTRAFLEX 1 8MM/60CM	ROD
97799669	ULTRAFLEX 1 8MM/80CM	ROD

643MMX" DEVICE

09991616	INSET 6MMX43"	UNK
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DRESS

09991615	IV3000 STANDARD	SMW
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3ML NEEDLE

00951417	T : SLIM X2 CARTRIDGE (SK)	UNK
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PATCH

09991614	MMT-174 ADHESIVE	UNK
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SYRINGE

97799707	RESERVOIR PARADIGM 5X1.8ML	MDT
97799706	RESERVOIR PARADIGM 7X3.0ML	MDT

ISOPROPYL ALCOHOL

70% PAD

00480452	ALCOHOL PREP	PDI
00809357	ALCOHOL SWABS	BTD
00977187	ALCOHOL SWABS 6893 BUTTERFLY	BTD
00977195	ALCOHOL SWABS 6896 (150)	BTD
02247809	ALCOHOL SWABS ANTISEPTIC SKIN CLEANERS	TIP
99038349	ALCOHOL SWABS BD REGULAR	BTD
97799880	BD ALCOHOL SWABS	BTD
99438102	MONOJECT ALCOHOL WIPES	COV
00795232	WEBCOL ALCOHOL PREP	COV

94:01.00 DEVICES (DIABETIC)

LANCET

Limited use benefit (prior approval not required).

The number of lancets that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 600 lancets per 100 days.
- Clients managing diabetes with high risk of causing hypoglycemia will be allowed 400 lancets per 365 days.
- Clients managing diabetes medication with low risk of causing hypoglycemia will be allowed 200 lancets per 365 days.
- Clients managing diabetes through diet/lifestyle therapy only (no insulin or anti-diabetes medications) will be allowed 200 lancets per 365 days.

Please note that the test strip limit is 500/100 days. Due to lancet pack sizes, 600 per 100 days will be reimbursed.

LANCET

97799494	ACCU-CHEK FASTCLIK LANCET	ROD
97799495	ACCU-CHEK FASTCLIK LANCET	ROD
97799817	ACCU-CHEK MULTICLIX LANCET	ROD
97799946	ACCU-CHEK MULTICLIX LANCET	ROD
97799945	ACCU-CHEK SOFTCLIX LANCET	ROD
97799466	BG STAR LANCET	SAC
97799541	EZ HEALTH ORACLE LANCET	TRE
97799825	FINGERSTIX LANCET	BAY
97799292	FIRST CANADIAN HEALTH LANCETS	ARA
97799826	FREESTYLE LANCET	BAY
97799918	MICROLET LANCET	BAY
97799810	MPD THIN LANCET (NS)	MPD
97799811	MPD THIN LANCET (NS)	MPD
97799807	MPD ULTRA THIN LANCET (100)	MPD
97799808	MPD ULTRA THIN LANCET (200)	MPD
97799970	ONETOUCH ULTRASOFT LANCET	JAJ
97799348	ULTILET CLASSIC LANCET	UNK

21G LANCET

97799804	MONOLET 21G LANCET	TYC
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28G LANCET

97799232	DROPLET PERSONAL LANCET 28G	SFA
97799253	FIRST CANHEALTH 28G LANCET	ARA
97799766	ITEST SAFETY 28G LANCET	AUC
97799801	MONOLET THIN (MONOJECT) 28G	TYC

30G LANCET

97799254	FIRST CANHEALTH 30G LANCET	ARA
97799388	MEDI+SURE SOFT 30G TWIST	MEC
97799389	MEDI+SURE SOFT 33G TWIST	MEC
97799431	ONE TOUCH DELICA 30G LANCET	JAJ

33G LANCET

97799690	BD ULTRAFINE 33G LANCET	BTD
97799234	DROPLET PERSONAL LANCET 33G	SFA
97799255	FIRST CANHEALTH 33G LANCET	ARA
97799767	ITEST ULTRA-THIN 33G LANCET	AUC
97799501	ONETOUCH DELICA 33G LANCET	JAJ

MAGNIFIER

DEVICE

99400550	SYRINGE SCALE MAGNIFIER	UNK
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94:01.00 DEVICES (DIABETIC)

PEN NEEDLE

ST NEEDLE

97799433	BD AUTOSHIELD DUO SAFETY PEN NEEDLE	BTD
09991447	BD BLUNT 18GX1 1/2 FILTER	BTD
09991387	BD PRECISIONGLIDE 25GX1 NEEDLE	BTD
00909114	BD ULTRA-FINE III PEN NEEDLE	BTD
00897590	NOVOLIN-PEN NEEDLE	NOO
97799280	SURECOMFORT 29GX1/2 NEEDLE	UNK
97799269	SURECOMFORT 30GX5/16 NEEDLE	UNK
97799279	SURECOMFORT 31GX3/16 NEEDLE	UNK
97799268	SURECOMFORT 31GX5/16 NEEDLE	UNK
97799278	SURECOMFORT 32GX1/4 NEEDLE	UNK
97799267	SURECOMFORT 32GX5/32 NEEDLE	UNK

ST 29GX10MM NEEDLE

97799238	DROPLET PEN NEEDLE 10MM 29G	SFA
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ST 29GX12.7MM NEEDLE

97799561	SUPER-FINE STANDARD 29G-12.7MM	PMS
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ST 29GX12MM NEEDLE

97799235	DROPLET PEN NEEDLE 12MM 29G	SFA
97799566	INSUPEN 29GX12MM NEEDLE	DPI
97799543	ULTICARE 29GX12MM PEN NEEDLE	UMI
97799991	UNIFINE 29G 12MM NEEDLE	AUC

ST 29GX8MM NEEDLE

97799526	BD AUTOSHIELD PEN NEEDLES	BTD
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ST 30GX6MM NEEDLE

97799911	NOVOFINE 30GX 6MM NEEDLE	NVC
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ST 30GX8MM NEEDLE

97799567	INSUPEN 30GX8MM NEEDLE	DPI
97799910	NOVOFINE 30GX 8MM NEEDLE	NVC

ST 31GX4.5MM NEEDLE

97799404	CLICKFINE PEN NEEDLE 31G 4.5MM	AUC
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ST 31GX5MM NEEDLE

97799282	BD ULTRAFINE 31G 5MM PEN NEEDLE	BTD
97799239	DROPLET PEN NEEDLE 5MM 31G	SFA
97799563	SUPER-FINE MICRO 31G-5MM NEEDLE	PMS
97799426	UNIFINE PENTIPS 31GX5MM	AUC

ST 31GX6MM NEEDLE

97799405	CLICKFINE PEN NEEDLE 31G 6MM	AUC
97799237	DROPLET PEN NEEDLE 6MM 31G	SFA
97799364	INSULIN PEN NEEDLE 31GX6MM	MDT
97799569	INSUPEN 31GX6MM NEEDLE	DPI
97799545	ULTICARE 31GX6MM PEN NEEDLE	UMI
97799993	UNIFINE 31G.6MM NEEDLE	AUC

ST 31GX8MM NEEDLE

97799281	BD ULTRAFINE 31G 8MM PEN NEEDLE	BTD
97799406	CLICKFINE PEN NEEDLE 31G 8MM	AUC
97799236	DROPLET PEN NEEDLE 8MM 31G	SFA
97799366	INSULIN PEN NEEDLE 31GX8MM	MDT
97799568	INSUPEN 31GX8MM NEEDLE	DPI

94:01.00 DEVICES (DIABETIC)

PEN NEEDLE

ST 31GX8MM NEEDLE

97799441	LIFE BRAND PEN NEEDLE 31G 8MM	HOD
97799562	SUPER-FINE XTRA 31G-8MM NEEDLE	PMS
97799544	ULTICARE 31GX8MM PEN NEEDLE	UMI
00963976	ULTRAFINE III NEEDLE 31G 8MM	BTD
97799992	UNIFINE 31G.8MM NEEDLE	AUC

ST 32GX4MM NEEDLE

97799527	BD ULTRA-FINE NANO PEN NEEDLE	BTD
97799243	DROPLET PEN NEEDLE 4MM 32G	SFA
97799367	INSULIN PEN NEEDLE 32GX4MM	MDT
97799399	INSUPEN 32GX4MM NEEDLE	DPI
97799334	MONTKIDDY BLUE NEEDLE 32GX4MM	MDT
97799337	MONTKIDDY GREEN NEEDLE 32GX4MM	MDT
97799335	MONTKIDDY PINK NEEDLE 32GX4MM	MDT
97799336	MONTKIDDY YELLOW NEEDLE 32GX4MM	MDT
97799386	NOVOFINE PLUS 4MM NEEDLE	NOO
97799440	ULTICARE 32GX4MM PEN NEEDLE	DPI

ST 32GX5MM NEEDLE

97799242	DROPLET PEN NEEDLE 5MM 32G	SFA
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ST 32GX6MM NEEDLE

97799241	DROPLET PEN NEEDLE 6MM 32G	SFA
97799363	INSULIN PEN NEEDLE 32GX6MM	MDT
97799571	INSUPEN 32GX6MM NEEDLE	DPI

ST 32GX8MM NEEDLE

97799240	DROPLET PEN NEEDLE 8MM 32G	SFA
97799365	INSULIN PEN NEEDLE 32GX8MM	MDT
97799570	INSUPEN 32GX8MM NEEDLE	DPI

ST 33GX4MM NEEDLE

97799383	INSUPEN 33GX4MM NEEDLE	DPI
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324GXMM NEEDLE

97799160	BD NANO PRO 32GX4MM PEN NEEDLE	BTD
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21G NEEDLE

09991504	BD BUTTERFLY NEEDLE 21G	BTD
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ST 29G NEEDLE

97799897	BD ULTRA-FINE PEN NEEDLE 29G	BTD
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ST 30G NEEDLE

97799467	NOVOTWIST TIP 30G NEEDLE	NOO
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ST 32G NEEDLE

97799821	NOVOFINE 32G TIP PEN NEEDLE	NOO
97799468	NOVOTWIST TIP 32G NEEDLE	NOO

SHARPS CONTAINER

DEVICE

99401026	BC SHARPS CONTAINER 1.4L	BTD
99401027	BD SHARPS CONTAINER 3.1L	BTD
09991639	BD SHARPS CONTAINER 3L	BTD
99401033	SHARPS NESTABLE YELLOW LARGE 22.7L	UNK

94:01.00 DEVICES (DIABETIC)

SYRINGE & NEEDLE

ST 27GX1/2 NEEDLE

09991381	BD PRECISIONGLIDE 27GX1/2	BTD
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ST 18G NEEDLE

09991402	BD PRECISIONGLIDE 18GX1 1/2	BTD
09991401	BD PRECISIONGLIDE 18GX1 NEEDLE	BTD

ST 25G NEEDLE

09991385	BD PRECISIONGLIDE 25GX5/8	BTD
09991386	BD PRECISIONGLIDE 25GX7/8	BTD

ST 26G NEEDLE

09991384	BD PRECISIONGLIDE 26GX1/2	BTD
09991383	BD PRECISIONGLIDE 26GX3/8	BTD

ST 27G NEEDLE

09991382	BD PRECISIONGLIDE 27GX1 1/4	BTD
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SYRINGE

09991609	BD POSIFLUSH SP	BTD
09991659	BD POSIFLUSH SP	BTD
00977020	PLASTIPAK MICRO	BTD
97799510	ULTICARE LOW DEAD SPACE SYRINGE	UMI

ST 0.25CC SYRINGE

99002132	INSULIN SYR W/NEEDL 0.25CC	UNK
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0.3CC SYRINGE

00977961	BD MICRO-FINE 0.3CC SYRINGE	BTD
99002140	INSULIN SYR W/NEEDLE 0.3CC	UNK

ST 0.5CC SYRINGE

00920096	E-Z JE	RIV
99002159	INSULIN SYR W/NEEDLE 0.5CC	UNK
00977136	MONOJECT	BTD

ST 0.5CC/1CC SYRINGE

00977128	MONOJECT	MDT
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ST 1CC SYRINGE

00920061	E-Z JE	RIV
99002167	INSULIN SYR W/NEEDLE 1CC	UNK

ST 1ML SYRINGE

09991376	BD LUER-LOK TIP 1ML SYRINGE	BTD
09991375	BD SLIP TIP 1ML SYRINGE	BTD

ST 3ML SYRINGE

09991371	BD LUER-LOK TIP 3ML SYRINGE	BTD
09991372	BD SLIP TIP 3ML SYRINGE	BTD

ST 5ML SYRINGE

09991373	BD LUER-LOK TIP 5ML SYRINGE	BTD
09991374	BD SLIP TIP 5ML SYRINGE	BTD

ST 8MM SYRINGE

97799261	SURECOMFORT 5/16 IN 30GX0.3CC	UNK
97799272	SURECOMFORT 5/16 IN 30GX0.5CC	UNK
97799265	SURECOMFORT 5/16 IN 30GX1CC	UNK
97799273	SURECOMFORT 5/16 IN 31GX0.3CC	UNK
97799274	SURECOMFORT 5/16 IN 31GX0.3CC	UNK
97799263	SURECOMFORT 5/16 IN 31GX0.5CC	UNK
97799262	SURECOMFORT 5/16 IN 31GX1CC	UNK

ST 10ML SYRINGE

09991363	BD LUER-LOK TIP 10ML SYRINGE	BTD
09991364	BD SLIP TIP 10ML SYRINGE	BTD

94:01.00 DEVICES (DIABETIC)

SYRINGE & NEEDLE

ST 12MM SYRINGE		
97799275	SURECOMFORT 1/2 IN 28GX1CC SYRINGE	UNK
ST 12.7MM SYRINGE		
97799257	SURECOMFORT 1/2 IN 28GX0.5CC	UNK
97799260	SURECOMFORT 1/2 IN 29GX0.3CC	UNK
97799259	SURECOMFORT 1/2 IN 29GX0.5CC	UNK
97799258	SURECOMFORT 1/2 IN 29GX1CC	UNK
97799264	SURECOMFORT 1/2 IN 30GX0.3CC	UNK
97799270	SURECOMFORT 1/2 IN 30GX0.5CC	UNK
97799271	SURECOMFORT 1/2 IN 30GX1CC	UNK
ST 18GX1 1/2 SYRINGE		
09991349	BD LUER-LOK TIP 18GX1 1/2 SYRINGE	BTD
ST 20ML SYRINGE		
09991368	BD LUER-LOK TIP 20ML SYRINGE	BTD
09991369	BD SLIP TIP 20ML SYRINGE	BTD
ST 21GX1 SYRINGE		
09991360	BD TUBERCULIN 21GX1 SYRINGE	BTD
ST 22GX1 1/2 SYRINGE		
09991341	BD LUER-LOK TIP 22GX1 1/2 SYRINGE	BTD
ST 23GX5/8 SYRINGE		
09991339	BD LUER-LOK TIP 25GX5/8 SYRINGE	BTD
ST 25GX1 SYRINGE		
09991338	BD LUER-LOK TIP 25GX1 SYRINGE	BTD
ST 25GX1 1/2 SYRINGE		
09991337	BD LUER-LOK TIP 25GX1 1/2 SYRINGE	BTD
ST 25GX5/8 SYRINGE		
09991359	BD TUBERCULIN 25GX5/8 SYRINGE	BTD
ST 26GX3/8 SYRINGE		
09991358	BD TUBERCULIN 26GX3/8 SYRINGE	BTD
ST 26GX5/8 SYRINGE		
09991361	BD SLIP TIP SUB Q 26G SYRINGE	BTD
ST 27GX1/2 SYRINGE		
09991356	BD TUBERCULIN 27GX1/2 SYRINGE	BTD
09991357	BD TUBERCULIN 27GX1/2 SYRINGE	BTD
28GX0.5CC SYRINGE		
00920177	BD MICRO-FINE 28GX0.5CC SYRINGE	BTD
97799518	ULTICARE 1/2 IN 28GX0.5CC SYRINGE	UMI
28GX1CC SYRINGE		
00920185	BD MICRO-FINE 28GX1CC SYRINGE	BTD
97799517	ULTICARE 1/2 IN 28GX1CC SYRINGE	UMI
ST 29GX0.3CC SYRINGE		
97799509	ULTI SYG 1/2 IN 29GX0.3CC	UMI
97799999	ULTICARE 29GX0.3CC	AUC
97799887	ULTRA 29G3/10CC	BTD

94:01.00 DEVICES (DIABETIC)

SYRINGE & NEEDLE

ST 29GX0.5CC SYRINGE		
97799888	BD ULTRA 29G.1/2CC SYRINGE	BTD
97799508	ULTI SYG 1/2 IN 29GX0.5CC	UMI
97799998	ULTICARE 29GX0.5CC	AUC
ST 29GX1CC SYRINGE		
97799889	BD ULTRA 29G.1CC SYRINGE	BTD
97799507	ULTI SYG 1/2 IN 29GX1CC SYRINGE	UMI
97799997	ULTICARE 29GX0.1CC	AUC
ST 30GX0.3CC SYRINGE		
97799551	ULTI SYG 1/2 IN 30GX0.3CC	UMI
97799506	ULTI SYG 5/16 IN 30GX0.3CC	UMI
97799996	ULTICARE 30GX0.3CC	AUC
97799886	ULTRA-FINE II 30GX0.3 CC SYRINGE	BTD
ST 30GX0.5CC SYRINGE		
97799885	BD ULTRA-FINE II 30GX0.5CC SYRINGE	BTD
97799550	ULTI SYG 1/2 IN 30GX0.5CC	UMI
97799505	ULTI SYG 5/16 IN 30GX0.5CC	UMI
97799995	ULTICARE 30GX0.5CC	AUC
ST 30GX1CC SYRINGE		
97799549	ULTI SYG 1/2 IN 30GX1CC SYRINGE	UMI
97799504	ULTI SYG 5/16 IN 30GX1CC SYRINGE	UMI
97799994	ULTICARE 30GX0.1CC	AUC
97799890	ULTRA-FINE II 30G.1CC	BTD
ST 30ML SYRINGE		
09991377	BD LUER-LOK TIP 30ML SYRINGE	BTD
09991378	BD SLIP TIP 30ML SYRINGE	BTD
ST 31GX0.3CC SYRINGE		
97799369	INSULIN 31GX0.3CC	MDT
97799548	ULTI SYG 5/16 IN 31GX0.3CC	UMI
97799513	ULTICARE 5/16 IN 31GX0.3CC SYRINGE	UMI
ST 31GX0.5CC SYRINGE		
97799370	INSULIN 31GX0.5CC	MDT
97799547	ULTI SYG 5/16 IN 31GX0.5CC	UMI
97799512	ULTICARE 5/16 IN 31GX0.5CC SYRINGE	UMI
ST 31GX1CC SYRINGE		
97799371	INSULIN 31GX1CC	MDT
97799546	ULTI SYG 5/16 IN 31GX1CC SYRINGE	UMI
97799511	ULTICARE 5/16 IN 31GX1CC SYRINGE	UMI
ST 31GX6MMX0.3CC SYRINGE		
97799425	BD SYRINGE WITH ULTRA-FINE NEEDLE	BTD
ST 31X6MMX0.5CC SYRINGE		
97799385	BD SYRINGE + NEEDLE	BTD
ST 31X6MMX1CC SYRINGE		
97799384	BD SYRINGE + NEEDLE	BTD
ST 60ML SYRINGE		
09991455	BD LUER-LOK TIP 60ML SYRINGE	BTD
09991454	BD SLIP TIP 60ML SYRINGE	BTD

94:01.00 DEVICES (DIABETIC)

SYRINGE CASE

DEVICE

99400552	MYHEALTH SYRINGE CASE-7	AUC
99400551	MYHEALTH SYRINGE CASE- SINGLE	AUC

96:00 PHARMACEUTICAL AIDS**96:00.00 PHARMACEUTICAL AIDS****ADMINISTRATION DIN****MISCELLANEOUS**

00903725 REFUSAL TO FILL UNK

DEVICE (METHADONE)

Limited use benefit (prior approval is not required).

Coverage is granted for 1 device.

MISCELLANEOUS

91500016 METHADONE LOCK BOX UNK

INFANT FORMULA

Limited use benefit (prior approval required).

Infant formula coverage for children < 1 YEAR OF AGE
(Corrected Gestational Age for Prematurity).**ORAL LIQUID**

95900000	ALIMENTUM	ABB
95900001	ALIMENTUM	UNK
95900003	ENFAMIL A+	MJO
95900007	ENFAMIL A+ READY TO FEED	MJO
95900152	ENFAMIL ENFACARE A+	MJO
95900026	NUTRAMIGEN A+	MJO

POWDER

95900047	ALIMENTUM PDR (400G)	ABB
95900164	ENFAMIL A+ 663G PDR	MJO
95900009	ENFAMIL ENFACARE A+ 363G PDR	MJO
95900025	NEOCATE W/ DHA & ARA 400G PDR	UNK
95900027	NUTRAMIGEN A+ LGG 566G PDR	MJO
95900035	PURAMINO A+ PDR	MJO
95900036	SIMILAC ADVANCE NEOSURE 363G	UNK

NUTRITIONAL SUPPLEMENT**POWDER**

09991319 SOURCE THICKEN UP 227G NVC

THICKENING AGENT**KIT**09991194 SIMPLY THICK 64OZ BOTTLE
PUMP UNK**POWDER**

12137029	RESOURCE THICKEN CLEAR	NVC
09991163	RESOURCE THICKEN UP 6.4G	NVC

THICKENING GEL**ORAL LIQUID**

09991164	SIMPLY THICK HONEY	UNK
09991035	SIMPLY THICK NECTAR	UNK

WATER**SOLUTION**

00905178	STERILE WATER	UNK
99002264	STERILE WATER	UNK

SYRINGE

09991563 STERILE WATER PF UNK

APPENDIX A
LIMITED USE BENEFITS AND CRITERIA

08:00 ANTI-INFECTIVE AGENTS**08:12.02 AMINOGLYCOSIDES****AMIKACIN SULFATE**

Limited use benefit (prior approval required).

250MG LIQUID

02242971 AMIKACIN SULFATE SDZ

TOBRAMYCIN

Limited use benefit (prior approval required).

28MG CAPSULE

02365154 TOBI PODHALER NVR

1.2G POWDER FOR SOLUTION

00533688 TOBRAMYCIN FKD

02285150 TOBRAMYCIN RAX

10MG/ML SOLUTION

02230639 TOBRAMYCIN FKD

02241209 TOBRAMYCIN SDZ

40MG/ML SOLUTION

02420287 JAMP-TOBRAMYCIN JMP

02230640 TOBRAMYCIN FKD

02241210 TOBRAMYCIN SDZ

02382814 TOBRAMYCIN MYL

99005069 TOBRAMYCINE UNK

60MG SOLUTION

02389622 TEVA-TOBRAMYCIN TEV

300MG SOLUTION

02443368 TOBRAMYCIN INHALATION SDZ

08:12.06 CEPHALOSPORINS**CEFTAZIDIME**

Limited use benefit (prior approval required).

1G POWDER FOR SOLUTION

00886971 CEFTAZIDIME FKD

02437848 CEFTAZIDIME RAX

02212218 FORTAZ 1G GSK

2G POWDER FOR SOLUTION

00886955 CEFTAZIDIME FKD

02437856 CEFTAZIDIME RAX

02212226 FORTAZ 2G GSK

3G POWDER FOR SOLUTION

02439522 CEFTAZIDIME RAX

6G POWDER FOR SOLUTION

00886963 CEFTAZIDIME FKD

02437864 CEFTAZIDIME RAX

02212234 FORTAZ 6G GSK

08:12.07 MISCELLANEOUS B-LACTAM ANTIBIOTICS**AZTREONAM**

Limited use benefit (prior approval required).

For the management of cystic fibrosis (CF) in patients if the following criteria are met:

- Patient has CF with chronic pulmonary Pseudomonas aeruginosa infections; AND
- Prescribed by a clinician with experience in the diagnosis and treatment of CF.

75MG POWDER FOR SOLUTION

02329840 CAYSTON

GIL

ERTAPENEM

Limited use benefit (prior approval required).

1G POWDER FOR SOLUTION

02247437 INVANZ

FRS

MEROPENEM

Limited use benefit (prior approval required).

500MG POWDER FOR SOLUTION

02378787 MEROPENEM

SDZ

1G POWDER FOR SOLUTION

02378795 MEROPENEM

SDZ

02436507 MEROPENEM

RAX

08:12.12 MACROLIDES**FIDAXOMICIN**

Limited use benefit (prior approval required).

For the treatment of confirmed severe Clostridium Difficile Infection (CDI); AND

Fidaxomicin has been prescribed or recommended by an infectious disease specialist or gastroenterologist; AND

There is a documented allergy (immune-mediated reaction) or severe intolerance to oral vancomycin resulting in discontinuation of vancomycin.

OR

• After an unsuccessful but adequate trial of oral vancomycin; AND

- Retreatment with vancomycin is not an option; AND

- The patient is at a high risk of hospitalization due to severe complications; AND

- Fidaxomicin is being used as monotherapy.

Notes:

a. Severe infection is defined as having any of the following symptoms: white blood cell count > 15,000 mm³ and fever; acute kidney injury with rising serum creatinine ≥ 1.5 times premorbid level or ≥ 175 micromoles/L; pseudomembranous colitis, hypotension, shock or megacolon.

b. An adequate trial of oral vancomycin is considered to be at least 10 days of therapy with a dose of at least 125mg four times daily.

c. Retreatment with fidaxomicin in recurrent CDI will be considered in symptomatic patients who require treatment of a previously resolved CDI episode. This is defined as a subsequent CDI episode occurring within 2 to 8 weeks of a previous episode from the date of diagnosis.

200MG TABLET

02387174 DIFICID

FRS

08:12.16 PENICILLINS**PIPERACILLIN, TAZOBACTAM**

Limited use benefit (prior approval required).

2G & 0.25G POWDER FOR SOLUTION

02299623 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

SDZ

02370158 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

TEV

3G & 0.375G POWDER FOR SOLUTION

02299631 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

SDZ

02308452 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

APX

02362627 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

RAX

02370166 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

TEV

4G & 0.5G POWDER FOR SOLUTION

02299658 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

SDZ

02308460 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

APX

02362635 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

RAX

02370174 PIPERACILLIN SODIUM/TAZOBACTAM SODIUM

TEV

08:12.16 PENICILLINS**PIPERACILLIN, TAZOBACTAM**

Limited use benefit (prior approval required).

12G & 1.5G POWDER FOR SOLUTION

02330547	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	SDZ
02377748	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	RAX

36G & 4.5G POWDER FOR SOLUTION

02439131	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	RAX
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08:12.18 QUINOLONES**LEVOFLOXACIN HEMIHYDRATE**

Limited use benefit (prior approval not required).

Coverage will be limited to 14 tablets every 14 days, followed by a 14 day lockout.

240MG SOLUTION

02442302	QUINSAIR	UNK
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250MG TABLET

02315424	ACT LEVOFLOXACIN	TEV
02284707	APO-LEVOFLOXACIN	APX
02284677	PMS-LEVOFLOXACIN	PMS
02298635	SANDOZ LEVOFLOXACIN	SDZ
02248262	TEVA-LEVOFLOXACIN	TEV

500MG TABLET

02315432	ACT LEVOFLOXACIN	TEV
02284715	APO-LEVOFLOXACIN	APX
02415879	LEVOFLOXACIN	PDL
02284685	PMS-LEVOFLOXACIN	PMS
02298643	SANDOZ LEVOFLOXACIN	SDZ
02248263	TEVA-LEVOFLOXACIN	TEV

750MG TABLET

02315440	ACT LEVOFLOXACIN	TEV
02325942	APO-LEVOFLOXACIN	APX
02305585	PMS-LEVOFLOXACIN	PMS
02298651	SANDOZ LEVOFLOXACIN	SDZ
02285649	TEVA-LEVOFLOXACIN	TEV

MOXIFLOXACIN HYDROCHLORIDE

Limited use benefit (prior approval not required).

Coverage will be limited to 14 tablets every 14 days, followed by a 14 day lockout.

400MG TABLET

02404923	APO-MOXIFLOXACIN	APX
02432242	AURO-MOXIFLOXACIN	AUR
02242965	AVELOX	BAY
02447266	BIO-MOXIFLOXACIN	BMI
02443929	JAMP-MOXIFLOXACIN	JMP
02447061	JAMP-MOXIFLOXACIN	JMP
02447053	MAR-MOXIFLOXACIN	MAR
02457814	MED-MOXIFLOXACIN	GMP
02472791	M-MOXIFLOXACIN	MAN
02462974	MOXIFLOXACIN	PDL
02450976	RIVA-MOXIFLOXACIN	RIV
02383381	SANDOZ MOXIFLOXACIN	SDZ
02375702	TEVA-MOXIFLOXACIN	TEV

08:12.28 MISCELLANEOUS ANTIBIOTICS**COLISTIN**

Limited use benefit (prior approval required).

For the management of cystic fibrosis (CF) in patients if the following criteria are met:

- Patient has CF with chronic pulmonary *Pseudomonas aeruginosa* infections; AND
- Prescribed by a clinician with experience in the diagnosis and treatment of CF.

150MG POWDER FOR SOLUTION

02244849 COLISTIMETHATE FOR U.S.P

RAX

00476420 COLY-MYCIN M PARENTERAL

ERF

LINEZOLID

Limited use benefit (prior approval required).

Tablets:

For treatment of proven vancomycin-resistant enterococci (VRE) infections.

For the treatment of proven methicillin-resistant staphylococcus aureus (MRSA) infections in patients who cannot tolerate vancomycin.

I.V. Solution:

When linezolid cannot be administered orally in the above mentioned situations.

Oral Liquid:

When linezolid cannot be administered orally in the above mentioned situations;

Plus at least one of the following:

- For treatment of proven vancomycin-resistant enterococci (VRE) infections
- For the treatment of proven methicillin-resistant staphylococcus aureus (MRSA) infections in patients who cannot tolerate vancomycin.

100MG POWDER FOR SUSPENSION

02243686 ZYVOXAM

PFI

2MG/ML SOLUTION

02243685 ZYVOXAM

PFI

600MG TABLET

02426552 APO-LINEZOLID

APX

02422689 SANDOZ LINEZOLID

SDZ

02243684 ZYVOXAM

PFI

RIFAXIMIN

Limited use benefit (prior approval required).

For reducing the risk of overt hepatic encephalopathy (HE) recurrence in patients:

- Who are unable to achieve adequate control of HE recurrence with a maximal tolerated dose of lactulose alone; AND
- When used in combination with a maximal tolerated dose of lactulose.

ST **550MG TABLET**

02410702 ZAXINE

SLX

VANCOMYCIN HYDROCHLORIDE (INJECTION)

Limited use benefit (prior approval required).

POWDER

99100176 VANCOMYCIN

MDS

500MG POWDER FOR SOLUTION

02420295 JAMP-VANCOMYCIN

JMP

02406535 MYLAN-VANCOMYCIN

MYL

02139375 VANCOMYCIN

FKD

02230191 VANCOMYCIN

PFI

02394626 VANCOMYCIN

SDZ

02411032 VANCOMYCIN

RAX

02435713 VANCOMYCIN

GMP

02342855 VANCOMYCIN HYDROCHLORIDE

RAX

1,000MG POWDER FOR SOLUTION

02230192 VANCOMYCIN

PFI

02396386 VANCOMYCIN

RAX

02435721 VANCOMYCIN

GMP

08:12.28 MISCELLANEOUS ANTIBIOTICS**VANCOMYCIN HYDROCHLORIDE (INJECTION)**

Limited use benefit (prior approval required).

1G POWDER FOR SOLUTION

02420309	JAMP-VANCOMYCIN	JMP
02406543	MYLAN-VANCOMYCIN	MYL
02241821	PMS-VANCOMYCIN 1 G	PMS
02139383	VANCOMYCIN	FKD
02394634	VANCOMYCIN	SDZ
02342863	VANCOMYCIN HYDROCHLORIDE	RAX

5G POWDER FOR SOLUTION

02420317	JAMP-VANCOMYCIN	JMP
02406551	MYLAN-VANCOMYCIN	MYL
02139243	VANCOMYCIN	FKD
02378337	VANCOMYCIN	PFI
02394642	VANCOMYCIN	SDZ

10G POWDER FOR SOLUTION

02420325	JAMP-VANCOMYCIN	JMP
02406578	MYLAN-VANCOMYCIN	MYL
02241807	VANCOMYCIN	FKD
02378345	VANCOMYCIN	PFI
02394650	VANCOMYCIN	SDZ
02411040	VANCOMYCIN	RAX
02405830	VANCOMYCIN HYDROCHLORIDE	RAX

08:14.08 AZOLES**VORICONAZOLE**

Limited use benefit (prior approval required).

For the treatment of patients with invasive aspergillosis; OR
For the treatment of culture proven invasive candidiasis with documented resistance to fluconazole.**50MG TABLET**

02409674	APO-VORICONAZOLE	APX
02399245	SANDOZ VORICONAZOLE	SDZ
02396866	TEVA-VORICONAZOLE	TEV
02256460	VFEND	PFI

200MG TABLET

02409682	APO-VORICONAZOLE	APX
02399253	SANDOZ VORICONAZOLE	SDZ
02396874	TEVA-VORICONAZOLE	TEV
02256479	VFEND	PFI

08:18.08 ANTIRETROVIRALS**TENOFOVIR DISOPROXIL FUMARATE**

Limited use benefit (prior approval required).

For the treatment of patients with HIV-1 infection who have failed or have experienced adverse events to an alternative agent.
For the treatment of patients with chronic hepatitis B infection who have cirrhosis documented on radiologic or histologic grounds and a HBV concentration above 2,000 IU/mL.**245MG TABLET**

02247128	VIREAD	GIL
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300MG TABLET

02451980	APO-TENOFOVIR	APX
02460173	AURO-TENOFOVIR	AUR
02479087	JAMP-TENOFOVIR	JMP

08:18.08 ANTIRETROVIRALS**TENOFOVIR DISOPROXIL FUMARATE**

Limited use benefit (prior approval required).

For the treatment of patients with HIV-1 infection who have failed or have experienced adverse events to an alternative agent.

For the treatment of patients with chronic hepatitis B infection who have cirrhosis documented on radiologic or histologic grounds and a HBV concentration above 2,000 IU/mL.

300MG TABLET

02452634 MYLAN-TENOFOVIR DISOPROXIL

MYL

02472511 NAT-TENOFOVIR

NPH

02453940 PMS-TENOFOVIR

PMS

02403889 TEVA-TENOFOVIR

TEV

08:18.20 INTERFERONS**PEGINTERFERON ALFA-2A**

Limited use benefit (prior approval required).

For the treatment of patients with chronic hepatitis B infection who have a HBV DNA concentration above 2,000 IU/mL without decompensated cirrhosis, upon the written request of a hepatologist or other specialist in this area.

180MCG/0.5ML SOLUTION

02248077 PEGASYS

HLR

PEGINTERFERON ALFA-2B, RIBAVIRIN

Limited use benefit (prior approval required).

For the treatment of chronic hepatitis C in patients who are treatment naïve, upon the written request of a hepatologist or other specialist in this area.

- For genotypes 1, 4, 5 and 6, an initial 24 week supply will be approved. A further 24 week supply may be approved if patient has a viral reduction of at least 2 logs or HCV is undetectable at 12 weeks (48 weeks total).

- For genotypes 2 or 3, initial coverage for a maximum of 24 weeks will be approved. Renewals will not be covered.

50MCG/0.5ML & 200MG KIT

02254573 PEGETRON KIT

FRS

PEGINTERFERON BETA-1A

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND

- Patient is fully ambulatory for 100 meters without aids; AND

- Patient is 18 years of age or older.

94MCG INJECTION

02444402 PLEGRIDY

UNK

125MCG LIQUID

02444399 PLEGRIDY

UNK

08:18.32 NUCLEOSIDES AND NUCLEOTIDES**ADEFOVIR DIPIVOXIL**

Limited use benefit (prior approval required).

For the treatment of chronic hepatitis B infection when used in combination with lamivudine in patients who have developed failure to lamivudine, as defined by an increase in HBV DNA of $\geq 1 \log_{10}$ IU/mL above the nadir, measured on two separate occasions within an interval of at least one month, after the first three months of lamivudine therapy, and when failure to lamivudine is not due to poor adherence to therapy.

10MG TABLET

02420333 APO-ADEFOVIR

APX

02247823 HEPSERA

GIL

08:18.32 NUCLEOSIDES AND NUCLEOTIDES**ENTECAVIR MONOHYDRATE**

Limited use benefit (prior approval required).

For the treatment of chronic hepatitis B infection in patients with cirrhosis documented on radiologic or histologic grounds and a HBV DNA concentration above 2000IU/mL.

0.5MG TABLET

02396955 APO-ENTECAVIR	APX
02448777 AURO-ENTECAVIR	AUR
02282224 BARACLUDE	BMS
02467232 JAMP ENTECAVIR	JMP
02430576 PMS-ENTECAVIR	PMS

08:18.40 HCV ANTIVIRALS**DACLATASVIR**

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria:

Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND

Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

30MG TABLET

02444747 DAKLINZA	BMS
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60MG TABLET

02444755 DAKLINZA	BMS
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ELBASVIR, GRAZOPREVR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria:

Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND

Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

50MG & 100MG TABLET

02451131 ZEPATIER	FRS
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GLECAPREVR, PIBRENTASVIR

Limited use benefit (prior approval required).

For treatment-naïve or treatment-experienced adult patients with genotypes 1, 2, 3, 4, 5, 6 with; OR

For the treatment of direct acting antivirals (DAA)-experienced2 adult patients with genotype 1 with:

- Chronic hepatitis C at any fibrosis stage (F0-F4); AND
- Detectable levels of HCV RNA in the last 12 months;

For genotypes 1, 2, 3, 4, 5 or 6, treatment-experienced is defined as a patient who has been previously treated with interferon, peginterferon (P), ribavirin (R) and/or sofosbuvir (SOF) (PR, SOF + PR, SOF + RBV), but no prior treatment experience with an NS3/4A protease inhibitor or NS5A inhibitor.

For genotype 1, DAA treatment-experienced is defined as a patient who has been previously treated with DAA regimens containing NS5A inhibitor [daclatasvir (DCV) + SOF or DCV + PR or ledipasvir/sofosbuvir, but no prior treatment experience with NS3/4A protease inhibitors] or containing NS3/4A protease inhibitors [simeprevir+SOF or simeprevir+PR or boceprevir+PR or telaprevir+PR, but no prior treatment experience with an NS5Ainhibitor].

100MG & 40MG TABLET

02467550 MAVIRET	ABV
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08:18.40 HCV ANTIVIRALS**RIBAVIRIN**

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria:
Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND
Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

200MG TABLET

02439212 IBAVYR

PED

400MG TABLET

02425890 IBAVYR

PED

600MG TABLET

02425904 IBAVYR

PED

SOFOSBUVIR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria:
Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND
Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

400MG TABLET

02418355 SOVALDI

GIL

SOFOSBUVIR, LEDIPASVIR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria:
Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND
Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

400MG & 90MG TABLET

02432226 HARVONI

GIL

SOFOSBUVIR, VELPATASVIR

Limited use benefit (prior approval required).

For adult patients with chronic hepatitis C infection at any fibrosis stage (F0-F4) who meet ALL of the following criteria:
Treatment is prescribed by a hepatologist, gastroenterologist, or infectious disease specialist (or other prescriber experienced in treating patients with chronic hepatitis C); AND
Laboratory confirmed quantitative HCV RNA level taken in the last 12 months;

Retreatment for failure or re-infection in patients who have received an adequate prior course of direct-acting antivirals will be considered on a case-by-case basis.

400MG & 100MG TABLET

02456370 EPCLUSA

GIL

08:18.40 HCV ANTIVIRALS**SOFOSBUVIR, VELPATASVIR, VOXILAPREVIR**

Limited use benefit (prior approval required).

For treatment-experienced adult patients with:

- Chronic hepatitis C at any fibrosis stage (F0-F4); AND
- Detectable levels of HCV RNA in the last 12 months;

AND

Treatment-experienced having failed a prior therapy with an HCV regimen containing:

- NS5A inhibitor: daclatasvir (Daklinza), elbasvir (part of Zepatier), ledipasvir (part of Harvoni), ombitasvir (part of Holkira Pak), velpatasvir (part of Eplusa) for genotype 1, 2, 3, 4, 5 or 6; OR
- sofosbuvir (Sovaldi) without an NS5A inhibitor for genotype 1, 2, 3 or 4.

400MG & 100MG & 100MG TABLET

02467542 VOSEVI

GIL

08:36.00 URINARY ANTI-INFECTIVES**FOSFOMYCIN TROMETHAMINE**

Limited use benefit (prior approval required).

For the treatment of women (>12 years old) with:

- Urinary tract infections with organisms resistant to first line therapy; OR
- Urinary tract infections in pregnancy when first-line agents are contraindicated.

3G/PK POWDER FOR SOLUTION

02240335 MONUROL

PAL

3G POWDER FOR SOLUTION

02473801 JAMP-FOSFOMYCIN

JMP

10:00 ANTINEOPLASTIC AGENTS**10:00.00 ANTINEOPLASTIC AGENTS****ABIRATERONE ACETATE**

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of metastatic castration resistant prostate cancer patients (mCRPC) who are asymptomatic or mildly symptomatic after failure of androgen deprivation therapy (ADT) and who have not received prior chemotherapy if they meet the following criteria:

- Used in combination with prednisone; AND
- Patient has an ECOG performance status of 0 or 1.

For the treatment of metastatic castration resistant prostate cancer patients (mCRPC) who progressed on docetaxel-based chemotherapy if they meet the following criteria:

- Used in combination with prednisone; AND
- Patient has an ECOG performance status \leq 2; AND
- Abiraterone is not used as an add-on therapy to enzalutamide (Xtandi); AND
- Abiraterone has not been used in the pre-docetaxel setting.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression

250MG TABLET

02371065 ZYTIGA

JSO

500MG TABLET

02457113 ZYTIGA

JSO

10:00.00 ANTINEOPLASTIC AGENTS**AFATINIB DIMALEATE**

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the treatment of patients with advanced Non-Small Cell Lung Cancer (NSCLC) who meet ALL of the following criteria:

- First line treatment of patients; AND
- EGFR mutation positive; AND
- Advanced or metastatic adenocarcinoma of the lung; AND
- An ECOG performance status of 0 or 1.

Criteria for renewal every 6 months:

- There is no objective evidence of disease progression.

Use of afatinib precludes the use of any other EGFR inhibitor as a subsequent line of therapy.

20MG TABLET

02415666 GIOTRIF

BOE

30MG TABLET

02415674 GIOTRIF

BOE

40MG TABLET

02415682 GIOTRIF

BOE

ALECTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

First-line treatment of patients with locally advanced or metastatic non-small cell lung cancer (NSCLC); OR

Second-line treatment of patients with locally advanced not amenable to curative therapy or metastatic NSCLC who have disease progression on or intolerance to crizotinib.

AND

To be used as monotherapy; AND

Disease is anaplastic lymphoma kinase (ALK)-positive; AND

Patient has a good performance status.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

150MG CAPSULE

02458136 ALECENSARO

HLR

AXITINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the second-line treatment of patients with advanced or metastatic clear cell renal carcinoma after failure of prior therapy with a first-line agent.

Patients are only eligible for either everolimus or axitinib in the second-line setting, but not sequential use of both agents except in cases of intolerance.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

1MG TABLET

02389630 INLYTA

PFI

5MG TABLET

02389649 INLYTA

PFI

10:00.00 ANTINEOPLASTIC AGENTS**BOSUTINIB**

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

Patients has Philadelphia chromosome positive (Ph+) chronic myelogenous leukemia (CML); AND

Patient has an ECOG performance status of 0 to 2;

AND

- Documented resistance/disease progression to at least one prior oral tyrosine kinase inhibitor [TKI] (imatinib, dasatinib or nilotinib); OR
- Documented intolerance to one prior oral TKI (imatinib, dasatinib or nilotinib) where subsequent treatment with an alternative oral TKI is not clinically appropriate.

Criteria for renewal every 12 months:

Confirmation from the clinician that the patient has experienced hematologic and/or cytogenetic response and is expected to continue to do so AND has not developed unacceptable toxicities.

100MG TABLET

02419149 BOSULIF

PFI

500MG TABLET

02419157 BOSULIF

PFI

CERITINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- Second-line treatment of patients with locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer (NSCLC) who have disease progression on or intolerance to crizotinib; AND
- To be used as monotherapy; AND
- Disease is anaplastic lymphoma kinase (ALK)-positive; AND
- Patient has an ECOG performance status of 0 to 2.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

150MG CAPSULE

02436779 ZYKADIA

NVR

COBIMETINIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the first-line treatment of patients with metastatic or unresectable melanoma in combination with vemurafenib (Zelboraf).

AND for patients who meet the following criteria:

- Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND
- Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND
- Patient has an ECOG performance status of 0 to 1.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

20MG TABLET

02452340 COTELLIC

HLR

CRIZOTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

First-line treatment of patients with advanced non-small cell lung cancer (NSCLC); OR

Second-line treatment of patients with advanced NSCLC who have received one prior chemotherapy regimen.*

AND

- Patient is anaplastic lymphoma kinase (ALK)-positive; AND
- Patient has an ECOG performance status of 0 to 2.

*Patients who have progressed during or following first-line therapy with crizotinib are not eligible to receive crizotinib as a second-line therapy.

Criteria for renewal every 12 months:

The patient has experienced a hematologic and/or cytogenetic response to crizotinib and is expected to continue to do so.

200MG CAPSULE

02384256 XALKORI

PFI

10:00.00 ANTINEOPLASTIC AGENTS**DABRAFENIB**

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

- For the first-line treatment of patients with metastatic or unresectable melanoma as monotherapy; OR
- For the first-line treatment of patients with metastatic or unresectable melanoma in combination with trametinib (Mekinist)

AND for patients who meet the following criteria:

- Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND
 - Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND
 - Patient has an ECOG performance status of 0 to 1;
- AND
- Patient is previously untreated.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

50MG CAPSULE

02409607 TAFINLAR

NVR

75MG CAPSULE

02409615 TAFINLAR

NVR

ENZALUTAMIDE

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of patients with metastatic castration-resistant prostate cancer (mCRPC) who are/have:

- Asymptomatic or mildly symptomatic after failure of androgen deprivation therapy (ADT) who have not received prior chemotherapy; AND
- Have an ECOG performance status of 0 or 1 with no risk factors for seizures; OR
- Progressed on docetaxel-based chemotherapy with an ECOG performance status ≤ 2 and no risk factors for seizures; AND
- Would be an alternative to abiraterone for patients in the post-docetaxel setting but would not be an add-on therapy to abiraterone treatment.

Patients previously treated with abiraterone would not be eligible for enzalutamide unless unable to tolerate abiraterone.

Use of enzalutamide in the post-docetaxel setting is not permitted if previously used in the pre-chemotherapy setting.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression

40MG CAPSULE

02407329 XTANDI

AST

ERLOTINIB HYDROCHLORIDE

Limited use benefit (prior approval required).

Treatment of non-small cell lung cancer (NSCLC) after failure of at least one prior chemotherapy regimen, and whose EGFR expression status is positive or unknown.

25MG TABLET

02461862 APO-ERLOTINIB

APX

02483912 NAT-ERLOTINIB

NPH

02269007 TARCEVA

HLR

02377691 TEVA-ERLOTINIB

TEV

100MG TABLET

02461870 APO-ERLOTINIB

APX

02454386 PMS-ERLOTINIB

PMS

02269015 TARCEVA

HLR

02377705 TEVA-ERLOTINIB

TEV

150MG TABLET

02461889 APO-ERLOTINIB

APX

02454394 PMS-ERLOTINIB

PMS

02269023 TARCEVA

HLR

02377713 TEVA-ERLOTINIB

TEV

10:00.00 ANTINEOPLASTIC AGENTS**EVEROLIMUS**

Limited use benefit (prior approval required).

1. Advanced Breast Cancer

Criteria for initial 12-month coverage:

For documented hormone receptor positive, HER2 negative advanced breast cancer; AND

- Used in combination with exemestane; AND
- Patient has an ECOG performance status of 0 to 2; AND
- Patient's condition recurred or progressed on a non-steroidal aromatase inhibitor.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

2. Advanced or Metastatic Renal Cell Carcinoma (mRCC)

Criteria for initial 12-month coverage:

For documented advanced or metastatic clear cell renal carcinoma; AND

For use as second- or third-line treatment of mRCC.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

3. Pancreatic Neuroendocrine Tumors (pNET)

Criteria for initial 12-month coverage:

For documented, progressive, unresectable, well or moderately differentiated, locally advanced or metastatic pancreatic neuroendocrine tumors; AND

- Patient has an ECOG performance status of 0 to 2; AND
- For patients previously treated with other agents, please specify reason for discontinuation.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

4. Non-functional Neuroendocrine Tumors (NETs) of Gastrointestinal or Lung Origin (GIL)

Criteria for initial 12-month coverage:

For documented unresectable, locally advanced or metastatic, progressive, well-differentiated non-functional NET-GIL in adults ≥ 18 years of age; AND

- Patient has documented radiological disease progression within the previous six months; AND
- Patient has an ECOG performance status of 0 to 2.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

2.5MG TABLET

02369257 AFINITOR

NVR

5MG TABLET

02339501 AFINITOR

NVR

10MG TABLET

02339528 AFINITOR

NVR

2MG TABLET FOR SUSPENSION

02425645 AFINITOR DISPERZ

NVR

3MG TABLET FOR SUSPENSION

02425653 AFINITOR DISPERZ

NVR

5MG TABLET FOR SUSPENSION

02425661 AFINITOR DISPERZ

NVR

GEFITINIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the treatment of patients with locally advanced or metastatic non-small cell lung cancer (NSCLC) who meet ALL of the following criteria:

- First-line treatment; AND
- EGFR mutation positive; AND
- Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

250MG TABLET

02468050 APO-GEFITINIB

APX

10:00.00 ANTINEOPLASTIC AGENTS**GEFITINIB**

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the treatment of patients with locally advanced or metastatic non-small cell lung cancer (NSCLC) who meet ALL of the following criteria:

- First-line treatment; AND
- EGFR mutation positive; AND
- Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

250MG TABLET

02248676 IRESSA

AZC

IBRUTINIB

Limited use benefit (prior approval required).

1. For the treatment of previously untreated chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL) (first-line)

Criteria for initial 12-month coverage:

As a first-line treatment option for newly diagnosed treatment naive chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL); AND

- Patient's prescriber has deemed that it would be inappropriate for the patient to receive treatment with a fludarabine-based regimen; AND
- Patient has high risk CLL, such that ibrutinib is preferred over anti-CD20 therapy, with one of the following cytogenetic markers:
 - Chromosome 17p deletion [del(17p)]
 - TP 53 mutation
 - Unmutated immunoglobulin heavy chain variable region (IgHV)
- Other reason.

Note: Anti-CD20 therapy is not funded as a sequential treatment option after ibrutinib. Choice of ibrutinib as first-line therapy must take this into account. Ibrutinib is not funded as a sequential treatment option for patients who have progressed on idelalisib treatment in the relapsed setting.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

2. For the treatment of chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL) (second-line)

Criteria for initial 12-month coverage:

Demonstrated diagnosis of chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL); AND

- Patient has received at least one prior therapy to treat CLL/SLL; AND
- Patient's prescriber has deemed that it would be inappropriate for the patient to receive treatment or retreatment with a fludarabine-based regimen.

Note: Ibrutinib is not funded as a sequential treatment option for patients who have progressed on idelalisib treatment in the relapsed setting.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

3. For the treatment of relapsed/refractory mantle cell lymphoma (MCL)

Criteria for initial 12-month coverage:

Demonstrated diagnosis of relapsed/refractory mantle cell lymphoma (MCL); AND

- Patient has received at least one prior therapy to treat MCL.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

140MG CAPSULE

02434407 IMBRUVICA

JSO

IDELALISIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

- For the treatment of patients with relapsed chronic lymphocytic leukemia (CLL) in combination with rituximab. Treatment should continue until unacceptable toxicity or disease progression.

Criteria for renewal every 6 months:

- There is no objective evidence of disease progression.

100MG TABLET

02438798 ZYDELIG

GIL

150MG TABLET

02438801 ZYDELIG

GIL

10:00.00 ANTINEOPLASTIC AGENTS**IMATINIB MESYLATE**

Limited use benefit (prior approval required).

- For the treatment of patients with chronic myeloid leukemia (CML) in blast crisis, accelerated phase, or in chronic phase.
- For the treatment of patients with gastrointestinal stromal tumour.
- For newly diagnosed adult patients with Philadelphia chromosome-positive (CML).
- For the treatment of adult patients with Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ ALL).

100MG TABLET

02355337 APO-IMATINIB	APX
02253275 GLEEVEC	NVR
02397285 NAT-IMATINIB	NPH
02431114 PMS-IMATINIB	PMS
02399806 TEVA-IMATINIB	TEV

400MG TABLET

02355345 APO-IMATINIB	APX
02253283 GLEEVEC	NVR
02397293 NAT-IMATINIB	NPH
02431122 PMS-IMATINIB	PMS
02399814 TEVA-IMATINIB	TEV

LENALIDOMIDE

Limited use benefit (prior approval required).

1. For the treatment of Myelodysplastic syndrome (MDS)

Criteria for initial 6-month coverage:

- Demonstrated diagnosis of Myelodysplastic syndrome (MDS) on bone marrow aspiration; AND
- Documented presence of del(5q) abnormality by standard cytogenetic or fluorescence in situ hybridization; AND
- International prognostic scoring system (IPSS) risk category low or intermediate-1; AND
- Transfusion-dependent symptomatic anemia.

Criteria for renewal every 12 months:

- Patient has demonstrated a reduction in transfusion requirements of at least 50%.

2. For the treatment of Refractory/relapsed Multiple Myeloma after one prior therapy (MM-AOPT)

Criteria for initial 12-month coverage:

- Progressive Multiple Myeloma; AND
- For use in combination with dexamethasone; AND
- Patient is refractory to initial or subsequent treatments or has relapsed after the conclusion of prior treatments and is suitable for further chemotherapy; OR
- Patient has completed at least one full treatment regimen as initial therapy and has demonstrated an intolerance to their current chemotherapy.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression or development of unacceptable toxicity to lenalidomide requiring discontinuation of therapy.

3. For the treatment of Newly diagnosed Multiple Myeloma for patients who are not eligible for autologous stem cell transplant - (MM-TNE)

Criteria for initial 12-month coverage:

- As a first-line treatment option for newly diagnosed patients with multiple myeloma who are not candidates for autologous stem-cell transplant; AND
- For use in combination with dexamethasone; AND
- Who have an ECOG performance status of 0 to 2.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression or development of unacceptable toxicity to lenalidomide requiring discontinuation of therapy.

4. For the maintenance treatment for newly diagnosed Multiple Myeloma post-autologous stem cell transplant

Criteria for initial 12-month coverage:

- Newly diagnosed Multiple Myeloma; AND
- The disease is stable or improved, with no evidence of progression after autologous stem-cell transplant.

Coverage is provided for lenalidomide at an initial dose of 10 mg daily. Doses adjustments of up to 15 mg daily may be required based on individual patient characteristics/response.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression or development of unacceptable toxicity to lenalidomide requiring discontinuation of therapy.

2.5MG CAPSULE

02459418 REVLIMID	UNK
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5MG CAPSULE

02304899 REVLIMID	UNK
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10:00.00 ANTINEOPLASTIC AGENTS**LENALIDOMIDE**

Limited use benefit (prior approval required).

1. For the treatment of Myelodysplastic syndrome (MDS)

Criteria for initial 6-month coverage:

- Demonstrated diagnosis of Myelodysplastic syndrome (MDS) on bone marrow aspiration; AND
- Documented presence of del(5q) abnormality by standard cytogenetic or fluorescence in situ hybridization; AND
- International prognostic scoring system (IPSS) risk category low or intermediate-1; AND
- Transfusion-dependent symptomatic anemia.

Criteria for renewal every 12 months:

- Patient has demonstrated a reduction in transfusion requirements of at least 50%.

2. For the treatment of Refractory/relapsed Multiple Myeloma after one prior therapy (MM-AOPT)

Criteria for initial 12-month coverage:

- Progressive Multiple Myeloma; AND
- For use in combination with dexamethasone; AND
- Patient is refractory to initial or subsequent treatments or has relapsed after the conclusion of prior treatments and is suitable for further chemotherapy; OR
- Patient has completed at least one full treatment regimen as initial therapy and has demonstrated an intolerance to their current chemotherapy.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression or development of unacceptable toxicity to lenalidomide requiring discontinuation of therapy.

3. For the treatment of Newly diagnosed Multiple Myeloma for patients who are not eligible for autologous stem cell transplant - (MM-TNE)

Criteria for initial 12-month coverage:

- As a first-line treatment option for newly diagnosed patients with multiple myeloma who are not candidates for autologous stem-cell transplant; AND
- For use in combination with dexamethasone; AND
- Who have an ECOG performance status of 0 to 2.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression or development of unacceptable toxicity to lenalidomide requiring discontinuation of therapy.

4. For the maintenance treatment for newly diagnosed Multiple Myeloma post-autologous stem cell transplant

Criteria for initial 12-month coverage:

- Newly diagnosed Multiple Myeloma; AND
- The disease is stable or improved, with no evidence of progression after autologous stem-cell transplant.

Coverage is provided for lenalidomide at an initial dose of 10 mg daily. Doses adjustments of up to 15 mg daily may be required based on individual patient characteristics/response.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression or development of unacceptable toxicity to lenalidomide requiring discontinuation of therapy.

10MG CAPSULE

02304902 REVLIMID

UNK

15MG CAPSULE

02317699 REVLIMID

UNK

20MG CAPSULE

02440601 REVLIMID

UNK

25MG CAPSULE

02317710 REVLIMID

UNK

10:00.00 ANTINEOPLASTIC AGENTS**LENVATINIB**

Limited use benefit (prior approval required).

Criteria for initial 4-month coverage:

- Used as monotherapy for treatment of patients with locally recurrent or metastatic, progressive differentiated thyroid cancer (DTC); AND
 - DTC is refractory to radioactive iodine treatment; AND
 - Have an ECOG performance status of ≤ 2 ;
- AND

Patient meets the eligibility criteria of the SELECT trial as follows:

- Pathologically confirmed differentiated thyroid cancer (patients with anaplastic or medullary thyroid cancer are not eligible)
- Evidence of iodine-131 refractory disease according to at least one of the following criteria:
 - At least one measurable lesion without iodine uptake on any iodine-131 scan
 - At least one measurable lesion that had progressed according to RECIST criteria within 12 months after iodine-131 therapy despite iodine-131 avidity at the time of treatment
 - Total lifetime radioactive iodine dose greater than 600 mCi (millicurie)
- Radiologic evidence of progression within the previous 13 months
- No prior therapy with a tyrosine kinase inhibitor or have received one prior treatment regimen with a tyrosine kinase inhibitor

Criteria for renewal every 4 months:

There is no objective evidence of disease progression.

10MG CAPSULE

02450321 LENVIMA

EIS

14MG CAPSULE

02450313 LENVIMA

EIS

20MG CAPSULE

02450305 LENVIMA

EIS

24MG CAPSULE

02450291 LENVIMA

EIS

MIDOSTAURIN

Limited use benefit (prior approval required).

Criteria for 12-month coverage:

- Patient has newly diagnosed FMS-like tyrosine kinase 3 (FLT3)-mutated acute myeloid leukemia (AML); AND
- Patient's FLT3-mutation status has been confirmed; AND
- Midostaurin is being used in combination with standard cytarabine and daunorubicin (or idarubicin) induction and cytarabine consolidation chemotherapy; AND
- Patient has an ECOG performance status of 0 to 2.

25MG CAPSULE

02466236 RYDAPT

NVR

NILOTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

Patients has newly diagnosed Philadelphia chromosome positive (Ph+) chronic myelogenous leukemia in chronic phase; OR
Patient has chronic phase or accelerated phase Philadelphia chromosome positive (Ph+) chronic myelogenous leukemia;

AND

- Patient has disease progression/resistance to imatinib; OR
- Documented intolerance to a prior oral TKI (imatinib, dasatinib or bosutinib).

Criteria for renewal every 12 months:

- Confirmation from the clinician that the patient has experienced hematologic and/or cytogenetic response and is expected to continue to do so AND has not developed unacceptable toxicities.

150MG CAPSULE

02368250 TASIGNA

NVR

200MG CAPSULE

02315874 TASIGNA

NVR

10:00.00 ANTINEOPLASTIC AGENTS**OLAPARIB**

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

- Maintenance treatment of adult patients with high grade serous epithelial ovarian fallopian tube cancer; OR
- Primary peritoneal cancer;

AND

- Platinum-sensitive disease; AND
- Relapsed BRCA-mutated disease (germline or somatic as detected by approved testing)
- Have completed at least two previous lines of platinum-based chemotherapy; AND
- Radiologic response (complete or partial response) to their most recent platinum-based chemotherapy regimen as per the SOLO-2 trial; AND
- Patient has an ECOG performance status of 0 to 2;

AND

- Olaparib is used as monotherapy

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

50MG CAPSULE

02454408 LYNPARZA

AZC

100MG TABLET

02475200 LYNPARZA

AZC

150MG TABLET

02475219 LYNPARZA

AZC

OSIMERTINIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

Patient with locally advanced or metastatic non-small cell lung cancer (NSCLC) who has progressed on epidermal growth factor receptor (EGFR) tyrosine kinase inhibitor therapy;

AND

Patient is EGFR T790M mutation- positive; AND

Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

40MG TABLET

02456214 TAGRISSO

AZC

80MG TABLET

02456222 TAGRISSO

AZC

PALBOCICLIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of post-menopausal women with estrogen receptor (ER)-positive, human epidermal growth factor receptor 2 (HER2)-negative advanced breast cancer; AND

- The patient has not received any prior treatment for metastatic disease (first-line treatment); AND
- Palbociclib will be used in combination with an aromatase inhibitor; AND
- Patient has an ECOG performance status of 0 to 2; AND
- Patient is not resistant to prior (neo)adjuvant aromatase inhibitor therapy; AND
- Patient does not have active or uncontrolled metastases to the central nervous system.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

75MG CAPSULE

02453150 IBRANCE

PFI

100MG CAPSULE

02453169 IBRANCE

PFI

125MG CAPSULE

02453177 IBRANCE

PFI

10:00.00 ANTINEOPLASTIC AGENTS**PAZOPANIB**

Limited use benefit (prior approval required).

Initial coverage criteria (12 months)

For the first-line treatment of patients with advanced or metastatic clear cell renal carcinoma; AND
Patient has an ECOG performance status of 0 to 2.

Renewal coverage criteria (12 months)

There is no objective evidence of disease progression.

200MG TABLET

02352303 VOTRIENT

NVR

POMALIDOMIDE

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of relapsed or refractory multiple myeloma who meet all of the following criteria:

- Used in combination with dexamethasone; AND
- Patient has relapsed or is refractory to at least two treatment regimens, including both bortezomib and lenalidomide; AND
- Patient has demonstrated disease progression on the last regimen.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression or development of unacceptable toxicity to pomalidomide requiring discontinuation of therapy.

1MG CAPSULE

02419580 POMALYST

UNK

2MG CAPSULE

02419599 POMALYST

UNK

3MG CAPSULE

02419602 POMALYST

UNK

4MG CAPSULE

02419610 POMALYST

UNK

PONATINIB HYDROCHLORIDE

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

- For the treatment of patients who have confirmed T315i mutation positive disease, independent of previous TKI therapy; OR
- Treatment of last resort for patients with intolerances or contraindications to imatinib and all other second generation TKI's (dasatinib, nilotinib, bosutinib); OR
- For the treatment of patients with chronic phase chronic myeloid leukemia (CML) who have resistance/disease progression after at least two prior lines of TKI therapy where Iclusig would be available as third-line TKI option; OR
- For the treatment of patients with accelerated phase or blast phase CML or Ph+ ALL who have resistance or disease progression after at least one second generation TKI therapy;

AND

- An ECOG performance status of 0 to 2.

Note: Second generation TKI's (dasatinib, nilotinib, bosutinib) are not covered as options after ponatinib.

Criteria for renewal every 6 months:

- There is no objective evidence of disease progression.

15MG TABLET

02437333 ICLUSIG

ARI

45MG TABLET

02437341 ICLUSIG

ARI

10:00.00 ANTINEOPLASTIC AGENTS**REGORAFENIB**

Limited use benefit (prior approval required).

1. For the treatment of Gastrointestinal Stromal Tumors (GIST)

Criteria for initial six-month coverage:

- For patients with gastrointestinal stromal tumors (GIST) who have failed or are unable to tolerate imatinib and sunitinib therapy; AND
- Patient has an ECOG performance status of 0 or 1;

Note: Regorafenib will not be funded concomitantly with imatinib or sunitinib.

Criteria for assessment every 12 months:

- There is no objective evidence of disease progression.

2. For the treatment of Hepatocellular Carcinoma (HCC)

Criteria for initial six-month coverage:

- Patient diagnosed with unresectable HCC; AND
- Patient has been previously treated with sorafenib; AND
- Patient was able to tolerate sorafenib as defined in the RESORCE trial criteria ($\geq 400\text{mg/day}$ for ≥ 20 days of the last 28 days of treatment); AND
- Patient has a Child-Pugh class status of A; AND
- Patient has an ECOG* performance status of 0 to 1

Criteria for assessment every 12 months:

- There is no objective evidence of disease progression.

40MG TABLET

02403390 STIVARGA

BAY

RITUXIMAB

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Initial coverage is provided for 24 weeks at a dose of 1000 mg x 2 doses at 0 & 2 weeks.

- Prescribed by a rheumatologist

For the treatment of adult patients with severely active rheumatoid arthritis who have failed to respond to a trial of an anti-TNF agent. Treatment should be combined with methotrexate. Rituximab should not be used in combination with anti-TNF agents.

For continued coverage for rituximab beyond twenty-four weeks, patient must meet all the following criteria:

- Initially prescribed by a rheumatologist;

AND

Patient has been assessed after the twentieth to twenty-fourth week of rituximab therapy and meets the response criteria of:

- $>20\%$ reduction in number of tender and swollen joints; PLUS
- $>20\%$ improvement in Physician Global Assessment scale; PLUS either
- $>20\%$ improvement in Patient Global Assessment scale; OR
- $>20\%$ reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of GRANULOMATOSIS POLYANGIITIS OR MICROSCOPIC POLYANGIITIS

Coverage is provided at a dose of 375 mg/m²body surface area, administered as an IV infusion once weekly for 4 weeks.

For the induction of remission in patients with severely active granulomatosis with polyangiitis or microscopic polyangiitis; AND

- Who have failed an adequate trial of cyclophosphamide; OR
- Who have a contraindication to cyclophosphamide.

10MG/ML SOLUTION

02241927 RITUXAN

HLR

10:00.00 ANTINEOPLASTIC AGENTS**RUXOLITINIB**

Limited use benefit (prior approval required).

1. For the treatment of Myelofibrosis:

Criteria for initial 6-month coverage:

- Intermediate to high risk symptomatic myelofibrosis as assessed using the Dynamic International Prognostic Scoring System (DIPSS) Plus; OR
 - Patient has symptomatic splenomegaly;
- AND
- Patient has an ECOG performance status of 0 to 3; AND
 - Patient previously untreated OR refractory to other treatment.

Criteria for renewal every 12 months:

- Reduction in spleen size; OR
- Improvement in disease symptoms.

2. For the treatment of patients with polycythemia vera:

Criteria for initial 6-month coverage:

Disease is resistant to hydroxyurea (HU) according to the modified European LeukemiaNet Criteria defined as below:

After 3 months of at least 2g/day of HU or at the maximally tolerated HU dose, patient showed:

- Need for phlebotomy to keep hematocrit < 45%; OR
- Uncontrolled myeloproliferation (platelet > 400x10⁹/L and WBC > 10x10⁹/L); OR
- Failure to reduce massive splenomegaly > 50% as measured by palpation.

OR

Patient is intolerant to HU according to the modified European LeukemiaNet Criteria defined below:

After any dose of HU, patient showed:

- Absolute neutrophil count < 1.0 x 10⁹/L , or platelet < 100x10⁹/L or hemoglobin < 100 g/L at the lowest dose of HU required to achieve a response (response defined as hematocrit < 45% without phlebotomy, and/or all of the following : platelet ≤ 400x10⁹/L , WBC ≤ 10 x 10⁹/L , and non-palpable spleen); OR
- Presence of leg ulcers or other unacceptable HU-related non-hematological toxicities (such as mucocutaneous manifestations, gastrointestinal symptoms, pneumonitis or fever, defined as Common Terminology Criteria for Adverse Events (CTCAE) version 3.0 grade 3 or 4, or more than one week of CTCAE version 3.0 grade 2, or permanent discontinuation of HU, or interruption of HU until toxicity resolved, or hospitalization due to HU toxicity).

AND

- Patient has an ECOG performance status of 0 to 3.

Criteria for renewal every 12 months:

- Reduction in spleen size; OR
- Improvement in disease symptoms.

5MG TABLET

02388006 JAKAVI

NVR

10MG TABLET

02434814 JAKAVI

NVR

15MG TABLET

02388014 JAKAVI

NVR

20MG TABLET

02388022 JAKAVI

NVR

SUNITINIB MALATE

Limited use benefit (Prior approval required).

Criteria for initial 6-month coverage:

- For patients with histologically proven unresectable or recurrent/metastatic GIST who have failed or are unable to tolerate imatinib therapy.
- Sunitinib will not be funded concomitantly with imatinib.

OR

Criteria for initial 12-month coverage:

- Documented, progressive, unresectable, well or moderately differentiated, locally advanced or metastatic pancreatic neuroendocrine tumors; AND
- Patient has an ECOG performance status of 0 to 2.

Criteria for renewal every 12 months:

- There is no objective evidence of disease progression.

12.5MG CAPSULE

02280795 SUTENT

PFI

25MG CAPSULE

02280809 SUTENT

PFI

50MG CAPSULE

02280817 SUTENT

PFI

10:00.00 ANTINEOPLASTIC AGENTS**TRAMETINIB**

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the first-line treatment of patients with metastatic or unresectable melanoma as monotherapy; OR
For the first-line treatment of patients with metastatic or unresectable melanoma in combination with dabrafenib(Tafinlar)

AND for patients who meet the following criteria:

- Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND
 - Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND
 - Patient has an ECOG performance status of 0 to 1;
- AND
- Patient is previously untreated.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

0.5MG TABLET

02409623 MEKINIST

NVR

2MG TABLET

02409658 MEKINIST

NVR

VANDETANIB

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For patients with symptomatic and/or progressive medullary thyroid cancer with unresectable locally advanced or metastatic disease; AND
An ECOG* performance status of 0 to 2.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression.

100MG TABLET

02378582 CAPRELSA

SAC

300MG TABLET

02378590 CAPRELSA

SAC

VEMURAFENIB

Limited use benefit (prior approval required).

Criteria for initial 6-month coverage:

For the first-line treatment of patients with metastatic or unresectable melanoma as monotherapy; OR
For the first-line treatment of patients with metastatic or unresectable melanoma in combination with cobimetinib (Cotellic).

AND for patients who meet the following criteria:

- Patient has documented BRAF V600 mutation-positive unresectable or metastatic melanoma; AND
- Patient does not have brain metastases or brain metastases are asymptomatic or stable; AND
- Patient has an ECOG performance status of 0 to 1.

Criteria for renewal every 6 months:

There is no objective evidence of disease progression.

ST 240MG TABLET

02380242 ZELBORAF

HLR

VENETOCLAX

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of chronic lymphocytic leukemia (CLL) who meet all of the following criteria:

- Venclexta will be used as monotherapy; AND
- Patient has received at least one prior therapy; AND
- Patient has failed a B-cell receptor inhibitor (BCRi) or is intolerant to prior ibrutinib therapy; AND
- Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression or unacceptable toxicity.

Coverage is for a maximum duration of two years.

10MG TABLET

02458039 VENCLEXTA

ABV

10:00.00 ANTINEOPLASTIC AGENTS**VENETOCLAX**

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

For the treatment of chronic lymphocytic leukemia (CLL) who meet all of the following criteria:

- Venclexta will be used as monotherapy; AND
- Patient has received at least one prior therapy; AND
- Patient has failed a B-cell receptor inhibitor (BCRI) or is intolerant to prior ibrutinib therapy; AND
- Patient has an ECOG* performance status of 0 to 2.

Criteria for renewal every 12 months:

There is no objective evidence of disease progression or unacceptable toxicity.

Coverage is for a maximum duration of two years.

50MG TABLET

02458047 VENCLEXTA

ABV

100MG TABLET

02458055 VENCLEXTA

ABV

02458063 VENCLEXTA

ABV

12:00 AUTONOMIC DRUGS**12:04.00 PARASYMPATHOMIMETIC AGENTS****DONEPEZIL HYDROCHLORIDE**

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:

- Diagnosis of mild to moderate Alzheimer's disease; AND
 - Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 - Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 - Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
- Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:

- Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
- Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 5MG TABLET

02419866 ACCEL-DONEPEZIL

ACP

02362260 APO-DONEPEZIL

APX

02232043 ARICEPT

PFI

02400561 AURO-DONEPEZIL

AUR

02412853 BIO-DONEPEZIL

BMI

02402645 DONEPEZIL

ACC

02416417 DONEPEZIL

PDL

02420597 DONEPEZIL

SIV

02426846 DONEPEZIL

SAN

02475278 DONEPEZIL

RIV

02416948 JAMP-DONEPEZIL

JMP

02402092 MAR-DONEPEZIL

MAR

02467453 M-DONEPEZIL

MAN

02439557 NAT-DONEPEZIL

NPH

02322331 PMS-DONEPEZIL

PMS

02381508 RAN-DONEPEZIL

RBV

02412918 RIVA-DONEPEZIL

RIV

02328666 SANDOZ DONEPEZIL

SDZ

02428482 SEPTA DONEPEZIL

SPT

02340607 TEVA-DONEPEZIL

TEV

ST 10MG TABLET

02419874 ACCEL-DONEPEZIL

ACP

02362279 APO-DONEPEZIL

APX

02232044 ARICEPT

PFI

12:04.00 PARASYMPATHOMIMETIC AGENTS**DONEPEZIL HYDROCHLORIDE**

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:

- Diagnosis of mild to moderate Alzheimer's disease; AND
 - Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 - Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 - Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
- Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:

- Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
- Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 10MG TABLET

02400588 AURO-DONEPEZIL	AUR
02412861 BIO-DONEPEZIL	BMI
02402653 DONEPEZIL	ACC
02416425 DONEPEZIL	PDL
02420600 DONEPEZIL	SIV
02426854 DONEPEZIL	SAN
02416956 JAMP-DONEPEZIL	JMP
02402106 MAR-DONEPEZIL	MAR
02467461 M-DONEPEZIL	MAN
02439565 NAT-DONEPEZIL	NPH
02322358 PMS-DONEPEZIL	PMS
02381516 RAN-DONEPEZIL	RBV
02412934 RIVA-DONEPEZIL	RIV
02328682 SANDOZ DONEPEZIL	SDZ
02428490 SEPTA DONEPEZIL	SPT
02340615 TEVA-DONEPEZIL	TEV

GALANTAMINE HYDROBROMIDE

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:

- Diagnosis of mild to moderate Alzheimer's disease; AND
 - Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 - Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 - Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
- Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:

- Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
- Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 8MG CAPSULE (EXTENDED RELEASE)

02425157 AURO-GALANTAMINE ER	AUR
02443015 GALANTAMINE	SAN
02416573 GALANTAMINE ER	PDL
02420821 MAR-GALANTAMINE ER	MAR
02339439 MYLAN-GALANTAMINE ER	MYL
02316943 PAT-GALANTAMINE ER	KLA
02398370 PMS-GALANTAMINE ER	PMS

ST 16MG CAPSULE (EXTENDED RELEASE)

02425165 AURO-GALANTAMINE ER	AUR
02443023 GALANTAMINE	SAN
02416581 GALANTAMINE ER	PDL
02420848 MAR-GALANTAMINE ER	MAR
02339447 MYLAN-GALANTAMINE ER	MYL
02316951 PAT-GALANTAMINE ER	KLA
02398389 PMS-GALANTAMINE ER	PMS

12:04.00 PARASYMPATHOMIMETIC AGENTS**GALANTAMINE HYDROBROMIDE**

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:

- Diagnosis of mild to moderate Alzheimer's disease; AND
 - Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 - Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 - Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
- Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:

- Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
- Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 24MG CAPSULE (EXTENDED RELEASE)

02425173 AURO-GALANTAMINE ER	AUR
02443031 GALANTAMINE	SAN
02416603 GALANTAMINE ER	PDL
02420856 MAR-GALANTAMINE ER	MAR
02339455 MYLAN-GALANTAMINE ER	MYL
02316978 PAT-GALANTAMINE ER	KLA
02398397 PMS-GALANTAMINE ER	PMS

RIVASTIGMINE HYDROGEN TARTRATE

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:

- Diagnosis of mild to moderate Alzheimer's disease; AND
 - Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 - Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 - Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
- Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:

- Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
- Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 1.5MG CAPSULE

02336715 APO-RIVASTIGMINE	APX
02242115 EXELON	NVR
02401614 MED-RIVASTIGMINE	GMP
02306034 PMS-RIVASTIGMINE	PMS
02416999 RIVASTIGMINE	PDL
02324563 SANDOZ RIVASTIGMINE	SDZ

ST 3MG CAPSULE

02336723 APO-RIVASTIGMINE	APX
02242116 EXELON	NVR
02401622 MED-RIVASTIGMINE	GMP
02306042 PMS-RIVASTIGMINE	PMS
02417006 RIVASTIGMINE	PDL
02324571 SANDOZ RIVASTIGMINE	SDZ

ST 4.5MG CAPSULE

02336731 APO-RIVASTIGMINE	APX
02242117 EXELON	NVR
02401630 MED-RIVASTIGMINE	GMP
02306050 PMS-RIVASTIGMINE	PMS
02417014 RIVASTIGMINE	PDL
02324598 SANDOZ RIVASTIGMINE	SDZ

ST 6MG CAPSULE

02336758 APO-RIVASTIGMINE	APX
02242118 EXELON	NVR
02401649 MED-RIVASTIGMINE	GMP

12:04.00 PARASYMPATHOMIMETIC AGENTS**RIVASTIGMINE HYDROGEN TARTRATE**

Limited use benefit (prior approval required).

Initial 12 month coverage for cholinesterase inhibitors:

- Diagnosis of mild to moderate Alzheimer's disease; AND
 - Mini Mental State Exam (MMSE) score of 10-26, established within the last 60 days; OR
 - Montreal Cognitive Assessment (MoCA) score of 10-26, established within the last 60 days; OR
 - Global Deterioration Scale (GDS) score between 4 to 6, established within the last 60 days.
- Continued coverage beyond 12 months will be based on improvement or stabilization of cognition, function or behaviour.

Criteria for coverage at every 12 month interval:

- Clinically meaningful response as determined by stabilization or improvement while on therapy; AND
- Alzheimer's Disease has not progressed to GDS stage 7 or MMSE or MoCA less than 10.

ST 6MG CAPSULE

02306069 PMS-RIVASTIGMINE	PMS
02417022 RIVASTIGMINE	PDL
02324601 SANDOZ RIVASTIGMINE	SDZ

ST 2MG/ML SOLUTION

02245240 EXELON	NVR
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12:08.08 ANTIMUSCARINICS / ANTISPASMODICS**TRIMEBUTINE MALEATE**

Limited use benefit (prior approval required).

For the treatment and relief of symptoms associated with functional bowel disorders including Irritable Bowel Syndrome (IBS), spastic colon, spastic colitis and mucous colitis; OR

In postoperative paralytic ileus in order to accelerate the resumption of the intestinal transit following abdominal surgery.

100MG TABLET

02349027 AA-TRIMEBUTINE	AAP
02245663 TRIMEBUTINE	AAP

200MG TABLET

02349035 AA-TRIMEBUTINE	AAP
02245664 TRIMEBUTINE	AAP

12:12.08 BETA ADRENERGIC AGONISTS**FLUTICASONE FUROATE, VILANTEROL TRIFENATATE**

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

OR

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry; OR
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA).

100MCG & 25MCG POWDER

02408872 BREO ELLIPTA	GSK
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FLUTICASONE FUROATE, VILANTEROL TRIFENATATE (ASTHMA)

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

200MCG & 25MCG POWDER

02444186 BREO ELLIPTA	GSK
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12:12.08 BETA ADRENERGIC AGONISTS**FORMOTEROL FUMARATE**

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of a rapid-onset, short-duration bronchodilator.

OR

For the treatment of Chronic Obstructive Pulmonary Disease (COPD) in patients not adequately controlled with either ipratropium, tiotropium or a short acting beta-agonist.

12MCG/CAPSULE CAPSULE

02230898 FORADIL

NVR

FORMOTEROL FUMARATE DIHYDRATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of rapid onset, short duration bronchodilator.

6MCG/DOSE POWDER

02237225 OXEZE TURBUHALER

AZC

12MCG/DOSE POWDER

02237224 OXEZE TURBUHALER

AZC

FORMOTEROL FUMARATE DIHYDRATE, BUDESONIDE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

OR

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry; OR
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA).

6MCG & 100MCG/INHALATION POWDER

02245385 SYMBICORT 100 TURBUHALER

AZC

6MCG & 200MCG/INHALATION POWDER

02245386 SYMBICORT 200 TURBUHALER

AZC

FORMOTEROL FUMARATE DIHYDRATE, MOMETASONE FUROATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of rapid onset, short duration bronchodilator.

5MCG & 100MCG/INHALATION AEROSOL

02361752 ZENHALE

FRS

5MCG & 200MCG/INHALATION AEROSOL

02361760 ZENHALE

FRS

5MCG & 50MCG/INHALATION AEROSOL

02361744 ZENHALE

FRS

INDACATEROL MALEATE

Limited use benefit (prior approval required).

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who:

- are not adequately controlled with either ipratropium, tiotropium or a short acting beta-agonist; OR
- have moderate to severe COPD, as defined by spirometry.

75MCG CAPSULE

02376938 ONBREZ BREEZHALER

NVR

12:12.08 BETA ADRENERGIC AGONISTS**SALMETEROL XINAFOATE**

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are using optimal corticosteroid therapy and experiencing breakthrough symptoms requiring regular use of a rapid-onset, short-duration bronchodilator.

OR

For the treatment of Chronic Obstructive Pulmonary Disease (COPD) in patients not adequately controlled with either ipratropium, tiotropium or a short acting beta-agonist.

50MCG/INHALATION POWDER

02231129 SEREVENT DISKUS

GSK

SALMETEROL XINAFOATE, FLUTICASONE PROPIONATE

Limited use benefit (prior approval required).

For the treatment of asthma in patients who are not adequately controlled on medium doses of inhaled corticosteroids (e.g. fluticasone 251-500mcg daily, or the equivalent) as the sole agent and require addition of a long-acting beta agonist. Patients using this combination product must also have access to a short-acting bronchodilator for symptomatic relief.

OR

For the treatment of chronic obstructive pulmonary disease (COPD) in patients who have:

- moderate to severe COPD, as defined by spirometry; OR
- inadequate response to a long-acting beta-2 agonist (LABA) or a long-acting muscarinic antagonist (LAMA).

25MCG & 125MCG AEROSOL

02245126 ADVAIR 125

GSK

25MCG & 250MCG AEROSOL

02245127 ADVAIR 250

GSK

50MCG & 100MCG POWDER

02240835 ADVAIR 100 DISKUS

GSK

50MCG & 250MCG POWDER

02240836 ADVAIR 250 DISKUS

GSK

50MCG & 500MCG POWDER

02240837 ADVAIR 500 DISKUS

GSK

12:20.04 CENTRALL ACTING SKELETAL MUSCLE RELAXANTS**CYCLOBENZAPRINE HYDROCHLORIDE**

Limited use benefit (prior approval is not required).

For relief of muscle spasm associated with acute, painful musculoskeletal conditions. Coverage is limited to 60mg per day for three (3) weeks renewable every two (2) months.

ST 10MG TABLET

02177145 APO-CYCLOBENZAPRINE

APX

02348853 AURO-CYCLOBENZAPRINE

AUR

02220644 CYCLOBENZAPRINE

PDL

02287064 CYCLOBENZAPRINE

SAN

02424584 CYCLOBENZAPRINE

SIV

02238633 DOM-CYCLOBENZAPRINE

DPC

02357127 JAMP-CYCLOBENZAPRINE

JMP

02212048 PMS-CYCLOBENZAPRINE

PMS

02242079 RIVA-CYCLOBENZAPRINE

RIV

02080052 TEVA-CYCLOBENZAPRINE

TEV

TIZANIDINE HYDROCHLORIDE

Limited use benefit (prior approval required).

For treatment of spasticity in patients with multiple sclerosis, who have failed therapy with or are intolerant to baclofen.

4MG TABLET

02239170 PAL-TIZANIDINE

PAL

02259893 TIZANIDINE

AAP

12:92.00 MISCELLANEOUS AUTONOMIC DRUGS**NICOTINE (GUM)**

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 945 pieces during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine gum or lozenges when one year has elapsed from the day the initial prescription was filled.

ST **2MG GUM**

02091933 NICORETTE GUM	KIM
80015240 RUGBY NICOTINE POLACRILEX GUM	ACG
80000396 THRIVE NICOTINELL GUM	GSK

ST **4MG GUM**

02091941 NICORETTE GUM	KIM
80000118 NICOTINE GUM	PER
80000402 THRIVE NICOTINELL GUM	NVC

NICOTINE (INHALER)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 945 doses during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine gum or lozenges when one year has elapsed from the day the initial prescription was filled.

ST **10MG SPRAY**

02241742 NICORETTE INHALER	KIM
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NICOTINE (LOZENGE)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 945 pieces during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine gum or lozenges when one year has elapsed from the day the initial prescription was filled.

ST **1MG LOZENGE**

80007461 THRIVE NICOTINE LOZENGES	NVC
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ST **2MG LOZENGE**

02247347 NICORETTE LOZENGE	KIM
80007464 THRIVE NICOTINE LOZENGES	NVC

ST **4MG LOZENGE**

02247348 NICORETTE LOZENGE	KIM
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NICOTINE (PATCH)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage will be provided for up to the allowable number of patches for one of the following products, during a one-year period. The year starts on the date the first prescription is filled.

NIHB clients are eligible to receive:

- up to 252 nicotine patches of any listed brand in a 12-month period; AND
- ONE course of an as-needed nicotine replacement therapy (NRT) product (i.e. gum, lozenge or inhaler) in a 12-month period; AND
- up to 180 tablets of Zyban in a 12-month period; AND
- up to 165 tablets of Champix in a 12-month period.

Once this quantity has been reached, the client is eligible again for coverage for nicotine patches when one year has elapsed from the day the initial prescription was filled.

ST **2MG GUM**

80025660 CHU NICOTINE ANTI SMOKING AID	UNK
94799974 THRIVE GUM (NS)	NVC

ST **1MG LOZENGE**

80061161 NICHIT	EUR
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ST **2MG LOZENGE**

80059877 NICHIT	EUR
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ST **7MG PATCH**

01943057 HABITROL	NVC
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12:92.00 MISCELLANEOUS AUTONOMIC DRUGS**NICOTINE (PATCH)**

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage will be provided for up to the allowable number of patches for one of the following products, during a one-year period. The year starts on the date the first prescription is filled.

NIHB clients are eligible to receive:

- up to 252 nicotine patches of any listed brand in a 12-month period; AND
- ONE course of an as-needed nicotine replacement therapy (NRT) product (i.e. gum, lozenge or inhaler) in a 12-month period; AND
- up to 180 tablets of Zyban in a 12-month period; AND
- up to 165 tablets of Champix in a 12-month period.

Once this quantity has been reached, the client is eligible again for coverage for nicotine patches when one year has elapsed from the day the initial prescription was filled.

ST 7MG PATCH

80051602 NICOTINE TRANSDERMAL	APX
80044393 TRANSDERMAL NICOTINE	ACG

ST 14MG PATCH

01943065 HABITROL	NVC
80013549 NICOTINE TRANSDERMAL SYSTEM	ADD
80044392 TRANSDERMAL NICOTINE	ACG

ST 18MG PATCH

02241227 TRANSDERMAL NICOTINE PATCHDAY	NVC
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ST 21MG PATCH

01943073 HABITROL	NVC
80051603 NICOTINE TRANSDERMAL	APX
80014250 NICOTINE TRANSDERMAL SYSTEM	ADD
80044389 TRANSDERMAL NICOTINE	ACG

ST 36MG PATCH

02093111 NICODERM	KIM
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ST 53MG PATCH

02241228 TRANSDERMAL NICOTINE PATCHDAY	NVC
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ST 78MG PATCH

02093138 NICODERM	KIM
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ST 114MG PATCH

02093146 NICODERM	KIM
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NICOTINE (SPRAY)

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 3450 sprays during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for nicotine spray when one year has elapsed from the day the initial prescription was filled.

1MG ORAL SPRAY

80038858 NICORETTE QUICKMIST	KIM
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VARENICLINE TARTRATE

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage will be limited to 165 tablets during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for varenicline (Champix®) when one year has elapsed from the day the initial prescription was filled.

ST 0.5MG TABLET

02419882 APO-VARENICLINE	APX
02291177 CHAMPIX	PFI

ST 0.5MG & 1MG TABLET

02298309 CHAMPIX STARTER PACK	PFI
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ST 1MG TABLET

02419890 APO-VARENICLINE	APX
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12:92.00 MISCELLANEOUS AUTONOMIC DRUGS**VARENICLINE TARTRATE**

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage will be limited to 165 tablets during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached, the client is eligible again for coverage for varenicline (Champix®) when one year has elapsed from the day the initial prescription was filled.

ST **1MG TABLET**

02435675 APO-VARENICLINE

APX

02291185 CHAMPIX

PFI

20:00 BLOOD FORMATION COAGULATION AND THROMBOSIS**20:04.04 IRON PREPARATIONS****POLYSACCHARIDE IRON COMPLEX**

Limited use benefit (prior approval not required).

For children 12 years of age or under.

15MG POWDER

80033717 FERAMAX POWDER WATER SOLUBLE POLYSACCHARIDE IRON COMPLEX

BSY

20:12.04 ANTICOAGULANTS**APIXABAN**

Limited use benefit (prior approval required).

For at-risk patients (CHADS2 score ≥ 1) with non-valvular atrial fibrillation who require apixaban for the prevention of stroke and systemic embolism AND in whom:

- Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
- Anticoagulation with warfarin is contraindicated; OR
- Anticoagulation with warfarin is not possible due to inability to regularly monitor via INR testing (i.e., no access to INR testing services at a laboratory, clinic, pharmacy and at home).

OR

For the treatment of venous thromboembolism: Deep Vein Thrombosis (DVT) or Pulmonary Embolism (PE)

ST **2.5MG TABLET**

02377233 ELIQUIS

BMS

ST **5MG TABLET**

02397714 ELIQUIS

BMS

DABIGATRAN ETEXILATE MESILATE

Limited use benefit (prior approval required).

For at-risk patients (CHADS2 score ≥ 1) with non-valvular atrial fibrillation who require dabigatran etexilate for the prevention of stroke and systemic embolism AND in whom:

- Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
- Anticoagulation with warfarin is contraindicated; OR
- Anticoagulation with warfarin is not possible due to inability to regularly monitor via INR testing (i.e., no access to INR testing services at a laboratory, clinic, pharmacy and at home).

ST **110MG CAPSULE**

02312441 PRADAXA

BOE

ST **150MG CAPSULE**

02358808 PRADAXA

BOE

20:12.04 ANTICOAGULANTS**EDOxabAN**

Limited use benefit (prior approval required).

For at-risk patients (CHADS2 score ≥ 1) with non-valvular atrial fibrillation who require apixaban for the prevention of stroke and systemic embolism AND in whom:

- Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
- Anticoagulation with warfarin is contraindicated; OR
- Anticoagulation with warfarin is not possible due to inability to regularly monitor via INR testing (i.e., no access to INR testing services at a laboratory, clinic, pharmacy and at home).

OR

For the treatment of venous thromboembolism: Deep Vein Thrombosis (DVT) or Pulmonary Embolism (PE)

15MG TABLET

02458640 LIXIANA

SEV

30MG TABLET

02458659 LIXIANA

SEV

60MG TABLET

02458667 LIXIANA

SEV

RIVAROXABAN

Limited use benefit (prior approval required).

Criteria for rivaroxaban 15 mg, 20mg tablets (Xarelto) for Stroke Prevention in Atrial Fibrillation (SPAF)

For at-risk patients (CHADS2 score ≥ 1) with non-valvular atrial fibrillation who require rivaroxaban for the prevention of stroke and systemic embolism AND in whom:

- Anticoagulation is inadequate (outside the desired INR range for at least 35% of the tests) with a two-month trial of warfarin; OR
- Anticoagulation with warfarin is contraindicated; OR
- Anticoagulation is not possible due to inability to regularly monitor via International Normalized Ratio (INR) testing (i.e., no access to INR testing service at a laboratory, clinic, pharmacy, and at home)

Criteria for rivaroxaban 15 mg, 20mg tablets (Xarelto)

For the treatment of venous thromboembolism: Deep Vein Thrombosis (DVT) or Pulmonary Embolism (PE).

ST **15MG TABLET**

02378604 XARELTO

BAY

ST **20MG TABLET**

02378612 XARELTO

BAY

RIVAROXABAN (10)

Limited use benefit (prior approval not required).

For the prevention of venous thromboembolism following total knee replacement or total hip replacement surgery, for up to 35 days.

ST **10MG TABLET**

02316986 XARELTO

BAY

20:16.00 HEMATOPOIETIC AGENTS**PEGFILGRASTIM**

Limited use benefit (prior approval required).

CHEMOTHERAPY SUPPORT

Primary Prophylaxis

For use in previously untreated patients receiving a moderate to severely myelosuppressive chemotherapy regimen (i.e. $\geq 40\%$ incidence of febrile neutropenia). Febrile neutropenia is defined as a temperature $\geq 38.5^{\circ}\text{C}$ or $>38.0^{\circ}\text{C}$ three times in a 24 hour period and neutropenia with an absolute neutrophil count (ANC) $<0.5 \times 10^9/\text{L}$.

Secondary Prophylaxis

For use in patients receiving myelosuppressive chemotherapy who have experienced an episode of febrile neutropenic sepsis or profound neutropenia in a previous cycle of chemotherapy; OR

For use in patients who have experienced a dose reduction or treatment delay longer than one week, due to neutropenia.

The recommended dosage of pegfilgrastim is a single subcutaneous injection of 6 mg, administered once per cycle of chemotherapy. Pegfilgrastim should be administered no sooner than 24 hours after the administration of cytotoxic chemotherapy.

10MG/ML SOLUTION

02249790 NEULASTA

AMG

20:16.00 HEMATOPOIETIC AGENTS**PLERIXAFOR**

Limited use benefit (prior approval required).

For use in combination with filgrastim to mobilize hematopoietic stem cells for subsequent autologous transplantation in patients with:

- Non-Hodgkin's lymphoma (NHL); OR
- Multiple myeloma (MM);

AND

- Prescribed by an oncologist or hematologist.

AND if one of the following are met

- A PB^{CD34+} count of < 10 cells/uL after 4 days of filgrastim; OR
- Less than 50% of the target CD34 yield is achieved on the 1st day of apheresis (after being mobilized with filgrastim alone or following chemotherapy); OR
- If a patient has failed a previous stem cell mobilization with filgrastim alone or following chemotherapy.

Reimbursement is limited to a maximum of 4 doses (0.24mg/kg given daily) for a single mobilization attempt.

The dose of Mozobil is limited to a maximum of 40mg per day

20MG SOLUTION

02377225 MOZOBIL

SAC

24:00 CARDIOVASCULAR DRUGS**24:04.92 MISCELLANEOUS CARDIAC DRUGS****IVABRADINE (IVABRADINE HYDROCHLORIDE)**

Limited use benefit (prior approval required).

For the treatment of stable chronic heart failure with New York Heart Association (NYHA) class II or III symptoms in adult patients if the following criteria are met:

- Left ventricular ejection fraction \leq 35%; AND
- 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; AND
- Patient has had at least one hospitalization due to heart failure in the last year; AND
- NYHA class II to III symptoms despite at least four weeks of treatment with an angiotensin converting enzyme inhibitor (ACEI) or an angiotensin II receptor antagonist (ARB) in combination with a beta blocker and, if tolerated, a mineralocorticoid receptor antagonist (MRA).

5MG TABLET

02459973 LANCORA

SEV

7.5MG TABLET

02459981 LANCORA

SEV

24:06.24

ALIROCUMAB

Limited use benefit (prior approval required).

Initial Coverage (12 weeks):

For adult patients with heterozygous familial hypercholesterolemia (HeFH) who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following clinical 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 an LDL-C target of < 2.0 mmol/L for secondary CVD prevention, or at least a 50% reduction in LDL-C from untreated baseline for primary prevention despite:
 - Confirmed adherence to a high-dose statin (e.g., atorvastatin 80 mg or rosuvastatin 40 mg) in combination with ezetimibe for a total of at least 3 months of continuous treatment;
 - OR
 - Patient is unable to tolerate at least two statins, with at least one statin initiated at the lowest daily starting dose; AND
 - For each statin (two statins in total), dose reduction was attempted to resolve intolerable myopathy or creatine kinase > 5 times the upper limit of normal rather than statin discontinuation; AND
 - For each statin (two statins in total), intolerable myopathy or creatine kinase > 5 times the upper limit of normal was reversed upon statin discontinuation, but reoccurred with statin rechallenge where clinically appropriate; AND
 - Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment; AND
 - Other known determinants of intolerable symptoms or abnormal biomarkers have been ruled out;
 - OR
 - Patient developed confirmed and documented rhabdomyolysis;
 - OR
 - Patient has a contraindication to statins; AND
 - Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment.

Continued coverage (6 months):

- Patient is adherent to therapy;
- AND
- Patient has achieved a reduction in LDL-C of at least 40% from baseline.

Note: Annual coverage is limited to 26 prefilled syringes or prefilled pens per year.

75MG SOLUTION

02453754 PRALUENT	SAC
02453819 PRALUENT	SAC

150MG SOLUTION

02453762 PRALUENT	SAC
02453835 PRALUENT	SAC

24:06.24**EVOLOCUMAB**

Limited use benefit (prior approval required).

Initial coverage criteria (Initial approval for 12 weeks):

For adult patients with heterozygous familial hypercholesterolemia (HeFH) who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following clinical 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 an LDL-C target of < 2.0 mmol/L for secondary CVD prevention, or at least a 50% reduction in LDL-C from untreated baseline for primary prevention despite:
 - Confirmed adherence to a high-dose statin (e.g., atorvastatin 80 mg or rosuvastatin 40 mg) in combination with ezetimibe for a total of at least 3 months of continuous treatment;
 - OR
 - Patient is unable to tolerate at least two statins, with at least one statin initiated at the lowest daily starting dose; AND
 - For each statin (two statins in total), dose reduction was attempted to resolve intolerable myopathy or creatine kinase > 5 times the upper limit of normal rather than statin discontinuation; AND
 - For each statin (two statins in total), intolerable myopathy or creatine kinase > 5 times the upper limit of normal was reversed upon statin discontinuation, but reoccurred with statin rechallenge where clinically appropriate; AND
 - Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment; AND
 - Other known determinants of intolerable symptoms or abnormal biomarkers have been ruled out;
- OR
- Patient developed confirmed and documented rhabdomyolysis;
- OR
- Patient has a contraindication to statins; AND
- Confirmed adherence to ezetimibe for a total of at least 3 months continuous treatment.

Note: Annual coverage is limited to 26 prefilled autoinjectors (140 mg every 2 weeks) or 12 automated Mini-Dosers with prefilled cartridges (420 mg once a month).

Renewal coverage criteria (Renewal for 6 months):

- Patient is adherent to therapy;
- AND
- Patient has achieved a reduction in LDL-C of at least 40% from baseline.

Note: Annual coverage is limited to 26 prefilled Autoinjectors (140 mg every 2 weeks) or 12 automated Mini-Dosers with prefilled cartridges (420 mg once a month).

120MG SOLUTION

02459779 REPATHA

AMG

140MG SOLUTION

02446057 REPATHA

AMG

24:12.12 PHOSPHODIESTERASE INHIBITORS**SILDENAFIL CITRATE**

Limited use benefit (prior approval required).

Must be initiated by a Pulmonary Hypertension specialist

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization.

ST **20MG TABLET**

02418118 APO-SILDENAFIL R

APX

02412179 PMS-SILDENAFIL R

PMS

02279401 REVATIO

PFI

02319500 TEVA-SILDENAFIL R

TEV

TADALAFIL

Limited use benefit (prior approval required).

Must be initiated by a Pulmonary Hypertension specialist

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization.

ST **20MG TABLET**

02338327 ADCIRCA

LIL

02421933 APO-TADALAFIL PAH

APX

24:12.92 MISCELLANEOUS VASODILATING AGENTS**AMBRISENTAN**

Limited use benefit (prior approval required).

Maximum dose covered is 10 mg once daily.

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST **5MG TABLET**

02307065 VOLIBRIS

GSK

ST **10MG TABLET**

02307073 VOLIBRIS

GSK

BOSENTAN MONOHYDRATE

Limited use benefit (prior approval required).

Maximum dose covered is 125 mg twice daily

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST **62.5MG TABLET**

02399202 APO-BOSENTAN

APX

02383012 PMS-BOSENTAN

PMS

02386275 SANDOZ BOSENTAN

SDZ

02398400 TEVA-BOSENTAN

TEV

02244981 TRACLEER

JSO

ST **125MG TABLET**

02383020 PMS-BOSENTAN

PMS

02386283 SANDOZ BOSENTAN

SDZ

02244982 TRACLEER

JSO

24:24.00 BETA ADRENERGIC BLOCKING AGENTS**PROPRANOLOL (HEMANGIOL)**

Limited use benefit (prior approval required).

For the treatment of proliferating infantile hemangioma requiring systemic therapy and at least one of the following:

- Life or function-threatening hemangioma, OR
- Ulcerated hemangioma with pain and/or lack of response to simple wound care measures, OR
- Hemangioma with a risk of permanent scarring or disfigurement.

3.75MG SOLUTION

02457857 HEMANGIOL

PFD

24:32.20 MINERALOCORTICOIDE (ALDOSTERONE) RECEPTOR ANTAGONISTS**EPLERENONE**

Limited use benefit (prior approval required).

For the treatment of patients with New York Heart Association (NYHA) class II chronic heart failure with left ventricular systolic dysfunction (with ejection fraction \leq 35%), as an adjunct to standard therapy.

Note: Patients must be on optimal therapy with an angiotensin-converting-enzyme (ACE) inhibitor or an angiotensin-receptor blocker (ARB), and a beta-blocker (unless contraindicated) at the recommended dose or maximal tolerated dose.

25MG TABLET

02323052 INSPRA

PFI

02471442 MINT-EPLERENONE

MIN

50MG TABLET

02323060 INSPRA

PFI

02471450 MINT-EPLERENONE

MIN

24:32.92**VALSARTAN, SACUBITRIL**

Limited use benefit (prior approval required).

For the treatment of New York Heart Association (NYHA) class II or III heart failure if the following criteria are met:

- Must be initiated by a physician experienced in the treatment of heart failure; AND
 - Left ventricular ejection fraction < 40%; AND
 - NYHA class II to III symptoms despite at least four weeks of treatment with an angiotensin-converting enzyme inhibitor (ACEI) or an angiotensin II receptor antagonist (ARB); OR If your patient has a contraindication or intolerance to ACEI or ARBs;
- AND
- Must be used in combination with a beta blocker and an aldosterone antagonist (if tolerated); OR If your patient has a contraindication or intolerance to beta blockers or aldosterone antagonists.

26MG & 24MG TABLET

02446928 ENTRESTO

NVR

51MG & 49MG TABLET

02446936 ENTRESTO

NVR

103MG & 97MG TABLET

02446944 ENTRESTO

NVR

28:00 CENTRAL NERVOUS SYSTEM AGENTS**28:08.04 NONSTEROIDAL ANTI-INFLAMMATORY AGENTS****ACETYLSALICYLIC ACID**

Limited use benefit (prior approval is not required).

ASA 80 mg tablets are a benefit to clients age 21 years and under to allow access for use in pediatric conditions (e.g. Kawasaki Syndrome).

ST 80MG TABLET

02269139 ACETYLSALICYLIC ACID

JMP

02295563 LOWPRIN

EUR

02202360 RIVASA

RIV

ST 80MG TABLET (CHEWABLE)

02009013 ASAPHEN

PMS

02280167 ASATAB

ODN

02250675 EURO-ASA

EUR

02296004 LOWPRIN

SDZ

02429950 M-ASA

MAN

02311518 PRO-AAS

PDL

02202352 RIVASA

RIV

ST 80MG TABLET (DELAYED RELEASE)

02427176 ASA EC

SAN

02238545 ASAPHEN

PMS

02283905 JAMP-ASA

JMP

02311496 PRO-AAS

PDL

DICLOFENAC SODIUM (TOPICAL)

Limited use benefit (prior approval required).

For the treatment of osteoarthritis when:

- pain is inadequately controlled with acetaminophen AND a non-steroidal anti-inflammatory (NSAID); OR
- there is contraindication to acetaminophen and NSAID; OR
- there is intolerance to acetaminophen and NSAID.

ST 1.5% SOLUTION

02354403 APO-DICLOFENAC

APX

02476134 DICLOFENAC SODIUM

TEL

02434571 DICLOFENAC TOPICAL

RAX

02472309 JAMP DICLOFENAC TOPICAL

JMP

02356783 PMS-DICLOFENAC

PMS

02420988 TARO-DICLOFENAC

TAR

28:08.08 OPIATE AGONISTS**ACETAMINOPHEN, CAFFEINE CITRATE, CODEINE PHOSPHATE**

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

300MG & 15MG & 15MG TABLET

00653241	RATIO-LENOLTEC NO 2	TEV
02163934	TYLENOL WITH CODEINE NO.2	JSO

300MG & 15MG & 30MG TABLET

00653276	RATIO-LENOLTEC NO 3	TEV
02163926	TYLENOL WITH CODEINE NO.3	JSO

325MG & 30MG & 15MG TABLET

00293504	ATASOL 15	CHU
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325MG & 30MG & 30MG TABLET

00293512	ATASOL 30	CHU
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ACETAMINOPHEN, CODEINE PHOSPHATE

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

32MG & 1.6MG/ML ELIXIR

00816027	PMS-ACETAMINOPHEN	PMS
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300MG & 30MG TABLET

02232658	PROCET-30	PDL
00608882	TEVA-EMTEC-30	TEV
00789828	TRIATEC-30	RIV

ACETAMINOPHEN, OXYCODONE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

325MG & 5MG TABLET

02324628	APO-OXYCODONE/ACET	APX
02361361	OXYCODONE/ACET	SAN
02327171	OXYCODONE-ACET	PDL
02242468	RIVACOCET	RIV
02307898	SANDOZ OXYCODONE/ACETAMINOPHEN	SDZ
00608165	TEVA-OXYCOCET	TEV

ACETYLSALICYLIC ACID, OXYCODONE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

325MG & 5MG TABLET

00608157	TEVA-OXYCODAN	TEV
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28:08.08 OPIATE AGONISTS**CODEINE MONOHYDRATE, CODEINE SULFATE TRIHYDRATE**

Limited use benefit (prior approval required).

For treatment of:

- chronic pain and palliative care patients as an alternative to products containing codeine in combination with acetaminophen or ASA with or without caffeine; OR
- chronic pain and palliative care patients as an alternative to regular release codeine tablets when large doses are required.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

50MG TABLET (EXTENDED RELEASE)

02230302 CODEINE CONTIN CR PFR

100MG TABLET (EXTENDED RELEASE)

02163748 CODEINE CONTIN CR PFR

150MG TABLET (EXTENDED RELEASE)

02163780 CODEINE CONTIN CR PFR

200MG TABLET (EXTENDED RELEASE)

02163799 CODEINE CONTIN CR PFR

CODEINE PHOSPHATE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

5MG/ML LIQUID

00050024 CODEINE PHOSPHATE ATL

2MG/ML SOLUTION

00380571 LINCTUS CODEINE ATL

15MG TABLET

02009889 CODEINE RIV

00593435 TEVA-CODEINE TEV

30MG TABLET

02009757 CODEINE RIV

00593451 TEVA-CODEINE TEV

FENTANYL

Limited use benefit (prior approval required).

For the management of chronic pain in patients who are unresponsive or intolerant to at least one long-acting oral sustained released product, such as morphine, hydromorphone and oxycodone, despite appropriate dose titration and adjunctive therapy including laxatives and antiemetics.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

12MCG/HR PATCH

02395657 FENTANYL PDL

02341379 PMS-FENTANYL MTX PMS

02330105 RAN-FENTANYL MATRIX RBY

02327112 SANDOZ FENTANYL SDZ

02311925 TEVA-FENTANYL TEV

25MCG/HR PATCH

02395665 FENTANYL PDL

02341387 PMS-FENTANYL MTX PMS

02330113 RAN-FENTANYL MATRIX RBY

02327120 SANDOZ FENTANYL SDZ

02282941 TEVA-FENTANYL TEV

28:08.08 OPIATE AGONISTS**FENTANYL**

Limited use benefit (prior approval required).

For the management of chronic pain in patients who are unresponsive or intolerant to at least one long-acting oral sustained released product, such as morphine, hydromorphone and oxycodone, despite appropriate dose titration and adjunctive therapy including laxatives and antiemetics.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

50MCG/HR PATCH

02395673 FENTANYL	PDL
02341395 PMS-FENTANYL MTX	PMS
02330121 RAN-FENTANYL MATRIX	RBY
02327147 SANDOZ FENTANYL	SDZ
02282968 TEVA-FENTANYL	TEV

75MCG/HR PATCH

02395681 FENTANYL	PDL
02341409 PMS-FENTANYL MTX	PMS
02330148 RAN-FENTANYL MATRIX	RBY
02327155 SANDOZ FENTANYL	SDZ
02282976 TEVA-FENTANYL	TEV

100MCG/HR PATCH

02395703 FENTANYL	PDL
02341417 PMS-FENTANYL MTX	PMS
02330156 RAN-FENTANYL MATRIX	RBY
02327163 SANDOZ FENTANYL	SDZ
02282984 TEVA-FENTANYL	TEV

HYDROMORPHONE HYDROCHLORIDE

Limited use benefit.

Prior approval required for controlled release capsules only. Regular release dosage forms are full benefits and do not require prior approval.

For treatment of moderate to severe chronic pain when other opioids such as morphine have been ineffective in controlling pain or in patients experiencing intolerable side effects.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

3MG CAPSULE (EXTENDED RELEASE)

02476614 APO-HYDROMORPHONE	APX
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4.5MG CAPSULE (EXTENDED RELEASE)

02476622 APO-HYDROMORPHONE	APX
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6MG CAPSULE (EXTENDED RELEASE)

02476630 APO-HYDROMORPHONE	APX
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9MG CAPSULE (EXTENDED RELEASE)

02476649 APO-HYDROMORPHONE	APX
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12MG CAPSULE (EXTENDED RELEASE)

02476657 APO-HYDROMORPHONE	APX
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18MG CAPSULE (EXTENDED RELEASE)

02476665 APO-HYDROMORPHONE	APX
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24MG CAPSULE (EXTENDED RELEASE)

02476673 APO-HYDROMORPHONE	APX
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30MG CAPSULE (EXTENDED RELEASE)

02476681 APO-HYDROMORPHONE	APX
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3MG CAPSULE (SUSTAINED RELEASE)

02125323 HYDROMORPH CONTIN	PFR
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28:08.08 OPIATE AGONISTS**HYDROMORPHONE HYDROCHLORIDE**

Limited use benefit.

Prior approval required for controlled release capsules only. Regular release dosage forms are full benefits and do not require prior approval.

For treatment of moderate to severe chronic pain when other opioids such as morphine have been ineffective in controlling pain or in patients experiencing intolerable side effects.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

4.5MG CAPSULE (SUSTAINED RELEASE)

02359502 HYDROMORPH CONTIN PFR

6MG CAPSULE (SUSTAINED RELEASE)

02125331 HYDROMORPH CONTIN PFR

9MG CAPSULE (SUSTAINED RELEASE)

02359510 HYDROMORPH CONTIN PFR

12MG CAPSULE (SUSTAINED RELEASE)

02125366 HYDROMORPH CONTIN PFR

18MG CAPSULE (SUSTAINED RELEASE)

02243562 HYDROMORPH CONTIN PFR

24MG CAPSULE (SUSTAINED RELEASE)

02125382 HYDROMORPH CONTIN PFR

30MG CAPSULE (SUSTAINED RELEASE)

02125390 HYDROMORPH CONTIN PFR

1MG/ML LIQUID

01916386 PMS HYDROMORPHONE PMS

3MG SUPPOSITORY

01916394 PMS HYDROMORPHONE PMS

1MG TABLET

02364115 APO-HYDROMORPHONE APX

00705438 DILAUDID PFR

00885444 PMS-HYDROMORPHONE PMS

02319403 TEVA-HYDROMORPHONE TEV

2MG TABLET

02364123 APO-HYDROMORPHONE APX

00125083 DILAUDID PFR

00885436 PMS-HYDROMORPHONE PMS

02319411 TEVA-HYDROMORPHONE TEV

4MG TABLET

02364131 APO-HYDROMORPHONE APX

00125121 DILAUDID PFR

00885401 PMS-HYDROMORPHONE PMS

02319438 TEVA-HYDROMORPHONE TEV

8MG TABLET

02364158 APO-HYDROMORPHONE APX

00786543 DILAUDID PFR

00885428 PMS-HYDROMORPHONE PMS

02319446 TEVA-HYDROMORPHONE TEV

28:08.08 OPIATE AGONISTS**METHADONE HYDROCHLORIDE (METADOL)**

Limited use benefit (prior approval required) with the following criteria:

Prescriber is registered with Health Canada and is eligible to prescribe methadone for the management of pain; AND
For the management of moderate to severe cancer pain or chronic non-cancer pain, as an alternative to other opioids; OR
For the management of pain for palliative care patients. Pharmacists may only dispense a maximum supply of 30 days at one time.

1MG/ML SOLUTION

02247694 METADOL PAL

10MG/ML SOLUTION

02241377 METADOL PAL

1MG TABLET

02247698 METADOL PAL

5MG TABLET

02247699 METADOL PAL

10MG TABLET

02247700 METADOL PAL

25MG TABLET

02247701 METADOL PAL

MORPHINE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

1MG/ML SYRUP

00614491 DOLORAL 1 ATL

5MG/ML SYRUP

00614505 DOLORAL 5 ATL

MORPHINE SULFATE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

10MG CAPSULE (EXTENDED RELEASE)

02019930 M-ESLON ETH

15MG CAPSULE (EXTENDED RELEASE)

02177749 M-ESLON ETH

30MG CAPSULE (EXTENDED RELEASE)

02019949 M-ESLON ETH

60MG CAPSULE (EXTENDED RELEASE)

02019957 M-ESLON ETH

100MG CAPSULE (EXTENDED RELEASE)

02019965 M-ESLON ETH

200MG CAPSULE (EXTENDED RELEASE)

02177757 M-ESLON ETH

5MG SUPPOSITORY

00632228 STATEX PAL

10MG SUPPOSITORY

00632201 STATEX PAL

20MG SUPPOSITORY

00596965 STATEX PAL

28:08.08 OPIATE AGONISTS**MORPHINE SULFATE**

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

1MG/ML SYRUP

00591467 STATEX PAL

5MG TABLET

00594652 STATEX PAL

10MG TABLET

00594644 STATEX PAL

25MG TABLET

00594636 STATEX PAL

50MG TABLET

00675962 STATEX PAL

15MG TABLET (EXTENDED RELEASE)

02350815 MORPHINE SR SAN

02015439 MS CONTIN SR PFR

02244790 SANDOZ MORPHINE SR SDZ

02302764 TEVA-MORPHINE SR TEV

30MG TABLET (EXTENDED RELEASE)

02350890 MORPHINE SR SAN

02014297 MS CONTIN SR PFR

02244791 SANDOZ MORPHINE SR SDZ

02302772 TEVA-MORPHINE SR TEV

60MG TABLET (EXTENDED RELEASE)

02350912 MORPHINE SR SAN

02014300 MS CONTIN SR PFR

02244792 SANDOZ MORPHINE SR SDZ

02302780 TEVA-MORPHINE SR TEV

100MG TABLET (EXTENDED RELEASE)

02014319 MS CONTIN SR PFR

02302799 TEVA-MORPHINE SR TEV

200MG TABLET (EXTENDED RELEASE)

02014327 MS CONTIN SR PFR

02302802 TEVA-MORPHINE SR TEV

5MG TABLET (IMMEDIATE RELEASE)

02014203 MS IR PFR

10MG TABLET (IMMEDIATE RELEASE)

02014211 MS IR PFR

20MG TABLET (IMMEDIATE RELEASE)

02014238 MS IR PFR

30MG TABLET (IMMEDIATE RELEASE)

02014254 MS IR PFR

28:08.08 OPIATE AGONISTS**MORPHINE SULFATE (KADIAN)**

Limited use benefit (prior approval required).

- For the treatment of opioid dependence where methadone and Suboxone are not available or not appropriate; OR
- For the treatment of chronic pain.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

10MG CAPSULE (SUSTAINED RELEASE)

02242163 KADIAN	BGP
09991310 KADIAN	MAY

20MG CAPSULE (SUSTAINED RELEASE)

02184435 KADIAN	BGP
09991311 KADIAN	MAY

50MG CAPSULE (SUSTAINED RELEASE)

02184443 KADIAN	BGP
09991312 KADIAN	MAY

100MG CAPSULE (SUSTAINED RELEASE)

02184451 KADIAN	BGP
09991313 KADIAN	MAY

OXYCODONE HYDROCHLORIDE

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented an opioid dose limit of 200 mg morphine equivalents per day for non-cancer, non-palliative pain. This limit will be calculated based on the total dose of all opioids a client is receiving from NIHB within a 30-day period (i.e. 6000 morphine equivalents over 30 days).

10MG SUPPOSITORY

00392480 SUPEUDOL	SDZ
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20MG SUPPOSITORY

00392472 SUPEUDOL	SDZ
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5MG TABLET

02325950 OXYCODONE	PDL
02231934 OXY-IR	PFR
02319977 PMS-OXYCODONE	PMS
00789739 SUPEUDOL	SDZ

10MG TABLET

02325969 OXYCODONE	PDL
02240131 OXY-IR	PFR
02319985 PMS-OXYCODONE	PMS
00443948 SUPEUDOL	SDZ

20MG TABLET

02325977 OXYCODONE	PDL
02319993 PMS-OXYCODONE	PMS
02262983 SUPEUDOL	SDZ

20MG TABLET (IMMEDIATE RELEASE)

02240132 OXY-IR	PFR
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28:08.12 OPIATE PARTIAL AGONISTS**BUPRENORPHINE HYDROCHLORIDE, NALOXONE HYDROCHLORIDE**

Limited use benefit (prior approval required).

For the treatment of opioid dependence when:

- The client must be 16 years or older.
- In cases where the client lives in a remote or isolated location, confirmation is required that the community has the ability to support buprenorphine/naloxone administration. These supports include the safe daily witnessing, storage and handling of the buprenorphine/naloxone doses. After this confirmation, NIHB will approve the buprenorphine/naloxone for the client.

2MG & 0.5MG TABLET

02453908	ACT BUPRENORPHINE/NALOXONE	ACG
02408090	MYLAN-BUPRENORPHINE/NALOXONE	MYL
02424851	PMS-BUPRENORPHINE-NALOXONE	PMS
02295695	SUBOXONE	IND

8MG & 2MG TABLET

02453916	ACT BUPRENORPHINE/NALOXONE	ACG
02408104	MYLAN-BUPRENORPHINE/NALOXONE	MYL
02424878	PMS-BUPRENORPHINE-NALOXONE	PMS
02295709	SUBOXONE	IND

28:08.92 MISCELLANEOUS ANALGESICS AND ANTIPYRETICS**ACETAMINOPHEN**

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

ST 80MG/ML DROP

01904140	ACETAMINOPHEN	TAN
01905864	ACETAMINOPHEN	TLI
02263793	PEDIAPHEN	EUR
02027801	PEDIATRIX	TEV
00875988	TEMPRA INFANT	PAL
02046059	TYLENOL	MCL

ST 16MG/ML LIQUID

01905848	ACETAMINOPHEN	TLI
00792713	PDP-ACETAMINOPHEN	PED
02263807	PEDIAPHEN	EUR
00884553	TEMPRA CHILDREN'S	PAL

ST 32MG/ML LIQUID

01901389	ACETAMINOPHEN	JMP
01958836	ACETAMINOPHEN	TLI
00792691	PDP-ACETAMINOPHEN	PED
02263831	PEDIAPHEN	EUR
02027798	PEDIATRIX	TEV
00875996	TEMPRA CHILDREN'S DOUBLE STRENGTH	PAL
02046040	TYLENOL	MCL

325MG SUPPOSITORY

01919393	ABENOL	PED
02230436	ACET 325	PED
02046687	PMS-ACETAMINOPHEN	PMS

650MG SUPPOSITORY

02230437	ACET 650	PED
02046695	PMS-ACETAMINOPHEN	PMS

ST 80MG TABLET

02015676	ACETAMINOPHEN	TAN
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28:08.92 MISCELLANEOUS ANALGESICS AND ANTIPYRETICS**ACETAMINOPHEN**

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on acetaminophen. The limit accumulates against the amount of acetaminophen claimed to the program from plain acetaminophen and/or acetaminophen in combination with opioids such as codeine (i.e. Tylenol® #3) or oxycodone (i.e. Percocet®). A total of 360 grams of acetaminophen is permitted in a 100-day period, for a total daily dose of 3600mg/day.

ST 80MG TABLET

02263815 PEDIAPHEN EUR

ST 160MG TABLET

02230934 ACETAMINOPHEN TAN

ST 325MG TABLET

00605751 ACETAMINOPHEN VTH

00743542 ACETAMINOPHEN PMT

00789801 ACETAMINOPHEN TLI

01938088 ACETAMINOPHEN JMP

02022214 ACÉTAMINOPHÈNE RIV

02362198 ACÉTAMINOPHÈNE RIV

00544981 APO ACETAMINOPHEN APX

02229873 APO-ACETAMINOPHEN APX

00389218 NOVO-GESIC TEV

00559393 TYLENOL MCL

00723894 TYLENOL MCL

ST 500MG TABLET

00549703 ACETAMINOPHEN PMT

00605778 ACETAMINOPHEN VTH

00789798 ACETAMINOPHEN TLI

01939122 ACETAMINOPHEN JMP

01962353 ACETAMINOPHEN TAN

02252813 ACETAMINOPHEN PMT

02255251 ACETAMINOPHEN PMT

02022222 ACÉTAMINOPHÈNE RIV

02362228 ACÉTAMINOPHÈNE RIV

02362201 ACÉTAMINOPHÈNE BLASON SHIELD RIV

00545007 APO ACETAMINOPHEN APX

02229977 APO-ACETAMINOPHEN APX

00013668 ATASOL FORTE CHU

02355299 JAMP ACETAMINOPHEN BLAZON JMP

00482323 NOVO-GESIC FORTE TEV

00892505 PMS-ACETAMINOPHEN PMS

00723908 TYLENOL MCL

00559407 TYLENOL EXTRA STRENGTH MCL

ST 80MG TABLET (CHEWABLE)

01905856 ACETAMINOPHEN TLI

02017458 ACETAMINOPHEN RIV

02129957 ACETAMINOPHEN VTH

ST 160MG TABLET (CHEWABLE)

02017431 ACETAMINOPHEN RIV

02142805 ACETAMINOPHEN VTH

02263823 PEDIAPHEN EUR

02347792 TYLENOL JR STRENGTH FASTMELTS MCL

02241361 TYLENOL JUNIOR STRENGTH MCL

28:12.08 ANTICONVULSANTS - BENZODIAZEPINES**CLONAZEPAM**

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 0.25MG TABLET

02442027 CLONAZEPAM	SIV
02179660 PMS-CLONAZEPAM	PMS

ST 0.5MG TABLET

02177889 APO-CLONAZEPAM	APX
02230366 CLONAPAM	VAE
02442035 CLONAZEPAM	SIV
02048701 PMS-CLONAZEPAM	PMS
02207818 PMS-CLONAZEPAM-R	PMS
02311593 PRO-CLONAZEPAM	PDL
02242077 RIVA-CLONAZEPAM	RIV
00382825 RIVOTRIL	HLR
02239024 TEVA-CLONAZEPAM	TEV

ST 1MG TABLET

02230368 CLONAPAM	VAE
02442043 CLONAZEPAM	SIV
02048728 PMS-CLONAZEPAM	PMS
02311607 PRO-CLONAZEPAM	PDL

ST 2MG TABLET

02177897 APO-CLONAZEPAM	APX
02230369 CLONAPAM	VAE
02442051 CLONAZEPAM	SIV
02048736 PMS-CLONAZEPAM	PMS
02311615 PRO-CLONAZEPAM	PDL
02242078 RIVA-CLONAZEPAM	RIV
00382841 RIVOTRIL	HLR
02239025 TEVA-CLONAZEPAM	TEV

28:12.92 MISCELLANEOUS ANTICONVULSANTS**BRIVARACETAM**

Limited use benefit (prior approval required).

For adjunctive therapy in adult patients with refractory partial-onset seizures who meet all of the following criteria:

- Are under the care of a physician experienced in the treatment of epilepsy; AND
- Are currently receiving two or more antiepileptic medications; AND
- Have failed or demonstrated intolerance to at least two other antiepileptic medications; AND
- Are not receiving concurrent therapy with levetiracetam.

10MG TABLET

02452936 BRIVLERA	UCB
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25MG TABLET

02452944 BRIVLERA	UCB
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50MG TABLET

02452952 BRIVLERA	UCB
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75MG TABLET

02452960 BRIVLERA	UCB
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100MG TABLET

02452979 BRIVLERA	UCB
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28:12.92 MISCELLANEOUS ANTICONVULSANTS**ESLICARBAZEPINE ACETATE**

Limited use benefit (prior approval required).

For adjunctive therapy in adult patients with refractory partial-onset seizures who meet all of the following criteria:

- Are under the care of a physician experienced in the treatment of epilepsy; AND
- Are currently receiving two or more antiepileptic medications; AND
- Have failed or demonstrated intolerance to at least two other antiepileptic medications.

ST **200MG TABLET**

02426862 APTIOM

SPC

ST **400MG TABLET**

02426870 APTIOM

SPC

ST **600MG TABLET**

02426889 APTIOM

SPC

ST **800MG TABLET**

02426897 APTIOM

SPC

GABAPENTIN

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on gabapentin. The limit accumulates against the amount of gabapentin claimed to the program. A total of 400 grams of gabapentin is permitted in a 100-day period, for a total daily dose of 4000 mg/day.

ST **100MG CAPSULE**

02244304 APO-GABAPENTIN

APX

02321203 AURO-GABAPENTIN

AUR

02243743 DOM-GABAPENTIN

DPC

02246314 GABAPENTIN

SIV

02353245 GABAPENTIN

SAN

02416840 GABAPENTIN

ACC

02285819 GD-GABAPENTIN

PFI

02361469 JAMP-GABAPENTIN

JMP

02391473 MAR-GABAPENTIN

MAR

02084260 NEURONTIN

PFI

02243446 PMS-GABAPENTIN

PMS

02310449 PRO-GABAPENTIN

PDL

02319055 RAN-GABAPENTIN

RBY

02251167 RIVA-GABAPENTIN

RIV

02244513 TEVA-GABAPENTIN

TEV

ST **300MG CAPSULE**

02244305 APO-GABAPENTIN

APX

02321211 AURO-GABAPENTIN

AUR

02243744 DOM-GABAPENTIN

DPC

02246315 GABAPENTIN

SIV

02353253 GABAPENTIN

SAN

02416859 GABAPENTIN

ACC

02285827 GD-GABAPENTIN

PFI

02361485 JAMP-GABAPENTIN

JMP

02391481 MAR-GABAPENTIN

MAR

02084279 NEURONTIN

PFI

02243447 PMS-GABAPENTIN

PMS

02310457 PRO-GABAPENTIN

PDL

02319063 RAN-GABAPENTIN

RBY

02251175 RIVA-GABAPENTIN

RIV

02244514 TEVA-GABAPENTIN

TEV

ST **400MG CAPSULE**

02244306 APO-GABAPENTIN

APX

28:12.92 MISCELLANEOUS ANTICONVULSANTS**GABAPENTIN**

Limited use benefit (prior approval is not required).

For safety reasons NIHB has implemented a dose limit on gabapentin. The limit accumulates against the amount of gabapentin claimed to the program. A total of 400 grams of gabapentin is permitted in a 100-day period, for a total daily dose of 4000 mg/day.

ST 400MG CAPSULE

02321238	AURO-GABAPENTIN	AUR
02243745	DOM-GABAPENTIN	DPC
02246316	GABAPENTIN	SIV
02353261	GABAPENTIN	SAN
02416867	GABAPENTIN	ACC
02361493	JAMP-GABAPENTIN	JMP
02391503	MAR-GABAPENTIN	MAR
02084287	NEURONTIN	PFI
02243448	PMS-GABAPENTIN	PMS
02310465	PRO-GABAPENTIN	PDL
02319071	RAN-GABAPENTIN	RBY
02251183	RIVA-GABAPENTIN	RIV
02244515	TEVA-GABAPENTIN	TEV

ST 600MG TABLET

02293358	APO-GABAPENTIN	APX
02388200	GABAPENTIN	SIV
02392526	GABAPENTIN	ACC
02431289	GABAPENTIN	SAN
02285843	GD-GABAPENTIN	PFI
02402289	JAMP-GABAPENTIN	JMP
02239717	NEURONTIN	PFI
02255898	PMS-GABAPENTIN	PMS
02310473	PRO-GABAPENTIN	PDL
02259796	RIVA-GABAPENTIN	RIV
02248457	TEVA-GABAPENTIN	TEV

ST 800MG TABLET

02293366	APO-GABAPENTIN	APX
02388219	GABAPENTIN	SIV
02392534	GABAPENTIN	ACC
02431297	GABAPENTIN	SAN
02402297	JAMP-GABAPENTIN	JMP
02239718	NEURONTIN	PFI
02255901	PMS-GABAPENTIN	PMS
02310481	PRO-GABAPENTIN	PDL
02259818	RIVA-GABAPENTIN	RIV
02247346	TEVA-GABAPENTIN	TEV

ST 600MG TABLET (IMMEDIATE RELEASE)

02410990	GLN-GABAPENTIN	GLK
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ST 800MG TABLET (IMMEDIATE RELEASE)

02411008	GLN-GABAPENTIN	GLK
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28:12.92 MISCELLANEOUS ANTICONVULSANTS**LACOSAMIDE**

Limited use benefit (prior approval required).

For adjunctive therapy in adult patients with refractory partial-onset seizures who meet all of the following criteria:

- Are under the care of a physician experienced in the treatment of epilepsy; AND
- Are currently receiving two or more antiepileptic medications; AND
- Have failed or demonstrated intolerance to at least two other antiepileptic medications.

ST 50MG TABLET

02475332 AURO-LACOSAMIDE	AUR
02478196 PHARMA-LACOSAMIDE	PMS
02474670 SANDOZ LACOSAMIDE	SDZ
02472902 TEVA-LACOSAMIDE	TEV
02357615 VIMPAT	UCB

ST 100MG TABLET

02475340 AURO-LACOSAMIDE	AUR
02478218 PHARMA-LACOSAMIDE	PMS
02474689 SANDOZ LACOSAMIDE	SDZ
02472910 TEVA-LACOSAMIDE	TEV
02357623 VIMPAT	UCB

ST 150MG TABLET

02475359 AURO-LACOSAMIDE	AUR
02478226 PHARMA-LACOSAMIDE	PMS
02474697 SANDOZ LACOSAMIDE	SDZ
02472929 TEVA-LACOSAMIDE	TEV
02357631 VIMPAT	UCB

ST 200MG TABLET

02475367 AURO-LACOSAMIDE	AUR
02478234 PHARMA-LACOSAMIDE	PMS
02474700 SANDOZ LACOSAMIDE	SDZ
02472937 TEVA-LACOSAMIDE	TEV
02357658 VIMPAT	UCB

OXCARBAZEPINE (SUSPENSION)

Limited use benefit (prior approval is not required).

For patients 19 years of age or over who are unable to swallow the tablet formulation due to:

- Tube feeding; OR
- Severe dysphagia

Note:

Trileptal (oxcarbazepine) suspension is an open benefit for patients 18 years of age and under and does not require prior approval for these patients.

Oxcarbazepine tablets are an open benefit for patients of all ages and do not require prior approval.

60MG SUSPENSION

02244673 TRILEPTAL	NVR
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PREGABALIN

Limited use benefit (prior approval required).

For the treatment of neuropathic pain in patients who have failed to effectively treat their pain with a tricyclic antidepressant (TCA);
OR

For the treatment of neuropathic pain in patients who have a contraindication or intolerance to a tricyclic antidepressant (TCA).

Coverage is limited to a maximum of 600mg per day.

ST 25MG CAPSULE

02402912 ACT PREGABALIN	ACG
02394235 APO-PREGABALIN	APX
02433869 AURO-PREGABALIN	AUR
02402556 DOM-PREGABALIN	DPC
02435977 JAMP-PREGABALIN	JMP

28:12.92 MISCELLANEOUS ANTICONVULSANTS**PREGABALIN**

Limited use benefit (prior approval required).

For the treatment of neuropathic pain in patients who have failed to effectively treat their pain with a tricyclic antidepressant (TCA);

OR

For the treatment of neuropathic pain in patients who have a contraindication or intolerance to a tricyclic antidepressant (TCA).

Coverage is limited to a maximum of 600mg per day.

ST **25MG CAPSULE**

02268418	LYRICA	PFI
02417529	MAR-PREGABALIN	MAR
02423804	MINT-PREGABALIN	MIN
02467291	M-PREGABALIN	MAN
02479117	NRA-PREGABALIN	UNK
02359596	PMS-PREGABALIN	PMS
02396483	PREGABALIN	PDL
02403692	PREGABALIN	SIV
02405539	PREGABALIN	SAN
02476304	PREGABALIN	RIV
02392801	RAN-PREGABALIN	RBY
02377039	RIVA-PREGABALIN	RIV
02390817	SANDOZ PREGABALIN	SDZ
02361159	TEVA-PREGABALIN	TEV

ST **50MG CAPSULE**

02402920	ACT PREGABALIN	ACG
02394243	APO-PREGABALIN	APX
02433877	AURO-PREGABALIN	AUR
02402564	DOM-PREGABALIN	DPC
02435985	JAMP-PREGABALIN	JMP
02268426	LYRICA	PFI
02417537	MAR-PREGABALIN	MAR
02423812	MINT-PREGABALIN	MIN
02467305	M-PREGABALIN	MAN
02479125	NRA-PREGABALIN	UNK
02359618	PMS-PREGABALIN	PMS
02396505	PREGABALIN	PDL
02403706	PREGABALIN	SIV
02405547	PREGABALIN	SAN
02476312	PREGABALIN	RIV
02392828	RAN-PREGABALIN	RBY
02377047	RIVA-PREGABALIN	RIV
02390825	SANDOZ PREGABALIN	SDZ
02361175	TEVA-PREGABALIN	TEV

ST **75MG CAPSULE**

02402939	ACT PREGABALIN	ACG
02394251	APO-PREGABALIN	APX
02433885	AURO-PREGABALIN	AUR
02402572	DOM-PREGABALIN	DPC
02435993	JAMP-PREGABALIN	JMP
02268434	LYRICA	PFI
02417545	MAR-PREGABALIN	MAR
02424185	MINT-PREGABALIN	MIN
02467313	M-PREGABALIN	MAN
02479133	NRA-PREGABALIN	UNK

28:12.92 MISCELLANEOUS ANTICONVULSANTS**PREGABALIN**

Limited use benefit (prior approval required).

For the treatment of neuropathic pain in patients who have failed to effectively treat their pain with a tricyclic antidepressant (TCA);

OR

For the treatment of neuropathic pain in patients who have a contraindication or intolerance to a tricyclic antidepressant (TCA).

Coverage is limited to a maximum of 600mg per day.

ST **75MG CAPSULE**

02359626	PMS-PREGABALIN	PMS
02396513	PREGABALIN	PDL
02403714	PREGABALIN	SIV
02405555	PREGABALIN	SAN
02476320	PREGABALIN	RIV
02392836	RAN-PREGABALIN	RBY
02377055	RIVA-PREGABALIN	RIV
02390833	SANDOZ PREGABALIN	SDZ
02361183	TEVA-PREGABALIN	TEV

ST **150MG CAPSULE**

02402955	ACT PREGABALIN	ACG
02394278	APO-PREGABALIN	APX
02433907	AURO-PREGABALIN	AUR
02402580	DOM-PREGABALIN	DPC
02436000	JAMP-PREGABALIN	JMP
02268450	LYRICA	PFI
02417561	MAR-PREGABALIN	MAR
02424207	MINT-PREGABALIN	MIN
02467321	M-PREGABALIN	MAN
02479168	NRA-PREGABALIN	UNK
02359634	PMS-PREGABALIN	PMS
02396521	PREGABALIN	PDL
02403722	PREGABALIN	SIV
02405563	PREGABALIN	SAN
02476347	PREGABALIN	RIV
02392844	RAN-PREGABALIN	RBY
02377063	RIVA-PREGABALIN	RIV
02390841	SANDOZ PREGABALIN	SDZ
02361205	TEVA-PREGABALIN	TEV

ST **300MG CAPSULE**

02402998	ACT PREGABALIN	ACG
02394294	APO-PREGABALIN	APX
02436019	JAMP-PREGABALIN	JMP
02268485	LYRICA	PFI
02359642	PMS-PREGABALIN	PMS
02396548	PREGABALIN	PDL
02403730	PREGABALIN	SIV
02405598	PREGABALIN	SAN
02476371	PREGABALIN	RIV
02392860	RAN-PREGABALIN	RBY
02377071	RIVA-PREGABALIN	RIV
02390868	SANDOZ PREGABALIN	SDZ
02361248	TEVA-PREGABALIN	TEV

28:12.92 MISCELLANEOUS ANTICONVULSANTS**RUFINAMIDE**

Limited use benefit (prior approval required).

- For the adjunctive treatment of seizures associated with Lennox-Gastaut syndrome in adults and children 4 years and older when prescribed by a neurologist or experienced specialist.
- Patient has failed or is intolerant to or has contraindications to at least two adjunctive antiepileptic drugs.

ST **100MG TABLET**

02369613 BANZEL

EIS

ST **200MG TABLET**

02369621 BANZEL

EIS

ST **400MG TABLET**

02369648 BANZEL

EIS

28:16.04 ANTIDEPRESSANTS**BUPROPION HYDROCHLORIDE (ZYBAN)**

Limited use benefit with quantity and frequency limits (prior approval is not required).

For smoking cessation:

Coverage is limited to 180 tablets during a one-year period. The year starts on the date the first prescription is filled. Once this quantity has been reached the client is eligible again for coverage for bupropion hydrochloride when one year has elapsed from the day the initial prescription was filled.

ST **150MG TABLET (EXTENDED RELEASE)**

02238441 ZYBAN

VAE

28:16.08 ANTIPSYCHOTIC AGENTS**ASENAPINE MALEATE**

Limited use benefit (prior approval required).

For the acute treatment of manic or mixed episodes associated with bipolar I disorder as either:

- Monotherapy, after a trial of lithium or divalproex sodium has failed or is contraindicated, and trials of two atypical antipsychotic agents have failed due to intolerance or lack of response; OR
- Co-therapy with lithium or divalproex sodium, after trials of two atypical antipsychotic agents have failed due to intolerance or lack of response.

ST **5MG TABLET**

02374803 SAPHRIS

FRS

ST **10MG TABLET**

02374811 SAPHRIS

FRS

LURASIDONE HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of schizophrenia and schizoaffective disorders in patients:

- who have intolerance or lack of response to an adequate trial of another antipsychotic agent; OR
- a contraindication to another antipsychotic agent.

ST **20MG TABLET**

02422050 LATUDA

SPC

ST **40MG TABLET**

02387751 LATUDA

SPC

ST **60MG TABLET**

02413361 LATUDA

SPC

ST **80MG TABLET**

02387778 LATUDA

SPC

ST **120MG TABLET**

02387786 LATUDA

SPC

28:20.04 AMPHETAMINES**AMPHETAMINE, DEXTROAMPHETAMINE**

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST 5MG CAPSULE (EXTENDED RELEASE)

02439239	ACT AMPHETAMINE XR	TEV
02248808	ADDERALL XR	UNK
02440369	PMS-AMPHETAMINES XR	PMS
02457288	SANDOZ AMPHETAMINE XR	SDZ

ST 10MG CAPSULE (EXTENDED RELEASE)

02439247	ACT AMPHETAMINE XR	TEV
02248809	ADDERALL XR	UNK
02440377	PMS-AMPHETAMINES XR	PMS
02457296	SANDOZ AMPHETAMINE XR	SDZ

ST 15MG CAPSULE (EXTENDED RELEASE)

02439255	ACT AMPHETAMINE XR	TEV
02248810	ADDERALL XR	UNK
02440385	PMS-AMPHETAMINES XR	PMS
02457318	SANDOZ AMPHETAMINE XR	SDZ

ST 20MG CAPSULE (EXTENDED RELEASE)

02439263	ACT AMPHETAMINE XR	TEV
02248811	ADDERALL XR	UNK
02440393	PMS-AMPHETAMINES XR	PMS
02457326	SANDOZ AMPHETAMINE XR	SDZ

ST 25MG CAPSULE (EXTENDED RELEASE)

02439271	ACT AMPHETAMINE XR	TEV
02248812	ADDERALL XR	UNK
02440407	PMS-AMPHETAMINES XR	PMS
02457334	SANDOZ AMPHETAMINE XR	SDZ

ST 30MG CAPSULE (EXTENDED RELEASE)

02439298	ACT AMPHETAMINE XR	TEV
02248813	ADDERALL XR	UNK
02440415	PMS-AMPHETAMINES XR	PMS
02457342	SANDOZ AMPHETAMINE XR	SDZ

DEXTROAMPHETAMINE SULFATE

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST 10MG CAPSULE (SUSTAINED RELEASE)

02448319	ACT DEXTROAMPHETAMINE SR	ACG
01924559	DEXEDRINE SPANSULE	PAL

ST 15MG CAPSULE (SUSTAINED RELEASE)

02448327	ACT DEXTROAMPHETAMINE SR	ACG
01924567	DEXEDRINE SPANSULE	PAL

ST 5MG TABLET

01924516	DEXEDRINE	PAL
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28:20.04 AMPHETAMINES**DEXTROAMPHETAMINE SULFATE**

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST **5MG TABLET**

02443236 DEXTROAMPHETAMINE

AAP

LISDEXAMFETAMINE DIMESYLATE

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST **10MG CAPSULE**

02439603 VYVANSE

SHI

ST **20MG CAPSULE**

02347156 VYVANSE

SHI

ST **30MG CAPSULE**

02322951 VYVANSE

SHI

ST **40MG CAPSULE**

02347164 VYVANSE

SHI

ST **50MG CAPSULE**

02322978 VYVANSE

SHI

ST **60MG CAPSULE**

02347172 VYVANSE

SHI

28:20.32 CNS STIMULANTS**METHYLPHENIDATE HYDROCHLORIDE**

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST **5MG TABLET**

02273950 APO-METHYLPHENIDATE

APX

02326221 METHYLPHENIDATE

PDL

02234749 PMS-METHYLPHENIDATE

PMS

ST **10MG TABLET**

02249324 APO-METHYLPHENIDATE

APX

02326248 METHYLPHENIDATE

PDL

00584991 PMS-METHYLPHENIDATE

PMS

ST **20MG TABLET**

02249332 APO-METHYLPHENIDATE

APX

02326256 METHYLPHENIDATE

PDL

00585009 PMS-METHYLPHENIDATE

PMS

ST **18MG TABLET (EXTENDED RELEASE)**

02441934 ACT METHYLPHENIDATE ER

ACG

02452731 APO-METHYLPHENIDATE ER

APX

28:20.32 CNS STIMULANTS**METHYLPHENIDATE HYDROCHLORIDE**

Limited use benefit (prior approval is not required).

The NIHB Program introduced a dose coverage limit for stimulants on February 25, 2015 as part of a strategy to deal with the potential misuse and abuse of these medications. The stimulant dose coverage limit is set at 100 mg of methylphenidate equivalents* per day for adults and children. This limit is calculated based on the total dose of all stimulants that patients are receiving from NIHB. The Program will continue to monitor the utilization of stimulants and adjust the eligible dose limit as required.

* To convert to methylphenidate equivalents, 1 mg of METHYLPHENIDATE, or LISDEXAMFETAMINE is equal to 0.5 mg DEXTROAMPHETAMINE

ST 18MG TABLET (EXTENDED RELEASE)

02247732	CONCERTA	JSO
02413728	PMS-METHYLPHENIDATE ER	PMS
02315068	TEVA-METHYLPHENIDATE	TEV

ST 20MG TABLET (EXTENDED RELEASE)

02266687	APO-METHYLPHENIDATE SR	APX
02320312	SANDOZ METHYLPHENIDATE SR	SDZ

ST 27MG TABLET (EXTENDED RELEASE)

02441942	ACT METHYLPHENIDATE ER	ACG
02452758	APO-METHYLPHENIDATE ER	APX
02250241	CONCERTA	JSO
02413736	PMS-METHYLPHENIDATE ER	PMS
02315076	TEVA-METHYLPHENIDATE	TEV

ST 36MG TABLET (EXTENDED RELEASE)

02441950	ACT METHYLPHENIDATE ER	ACG
02452766	APO-METHYLPHENIDATE ER	APX
02247733	CONCERTA	JSO
02413744	PMS-METHYLPHENIDATE ER	PMS
02315084	TEVA-METHYLPHENIDATE	TEV

ST 54MG TABLET (EXTENDED RELEASE)

02441969	ACT METHYLPHENIDATE ER	ACG
02330377	APO-METHYLPHENIDATE ER	APX
02247734	CONCERTA	JSO
02413752	PMS-METHYLPHENIDATE ER	PMS
02315092	TEVA-METHYLPHENIDATE	TEV

28:20.92 MISC ANOREXIGENIC AGENTS & RESPIRATORY & CEREBRAL STIMULANT**CAFFEINE CITRATE**

Limited use benefit (prior approval not required).

For children up to 1 year of age

POWDER

00972037	CAFFEINE CITRATE	MDS
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28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES**ALPRAZOLAM**

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 0.25MG TABLET

01908189	ALPRAZOLAM	PDL
02349191	ALPRAZOLAM	SAN
00865397	APO-ALPRAZ	APX
02400111	JAMP-ALPRAZOLAM	JMP

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES**ALPRAZOLAM**

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 0.25MG TABLET

01913484 TEVA-ALPRAZOLAM	TEV
00548359 XANAX	PFI

ST 0.5MG TABLET

01908170 ALPRAZOLAM	PDL
02349205 ALPRAZOLAM	SAN
00865400 APO-ALPRAZ	APX
02400138 JAMP-ALPRAZOLAM	JMP
01913492 TEVA-ALPRAZOLAM	TEV
00548367 XANAX	PFI

ST 1MG TABLET

02248706 ALPRAZOLAM	PDL
02243611 APO-ALPRAZ	APX
02400146 JAMP-ALPRAZOLAM	JMP
00723770 XANAX	PFI

ST 2MG TABLET

02243612 APO-ALPRAZ	APX
02400154 JAMP-ALPRAZOLAM	JMP
00813958 XANAX TS	PFI

BROMAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 1.5MG TABLET

02177153 APO-BROMAZEPAM	APX
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ST 3MG TABLET

02177161 APO-BROMAZEPAM	APX
02220520 BROMAZEPAM	PDL
02230584 TEVA-BROMAZEPAM	TEV

ST 6MG TABLET

02177188 APO-BROMAZEPAM	APX
02220539 BROMAZEPAM	PDL
02230585 TEVA-BROMAZEPAM	TEV

DIAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 1MG/ML SOLUTION

00891797 PMS-DIAZEPAM	PMS
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28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES**DIAZEPAM**

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **2MG TABLET**

00405329 DIAZEPAM	AAP
02247490 PMS-DIAZEPAM	PMS

ST **5MG TABLET**

00313580 DIAZEPAM	PDL
00362158 DIAZEPAM	AAP
02247491 PMS-DIAZEPAM	PMS
00013285 VALIUM	HLR

ST **10MG TABLET**

00405337 DIAZEPAM	AAP
02247492 PMS-DIAZEPAM	PMS

DIAZEPAM (DIASTAT)

Limited use benefit (prior approval not required).

For children 12 years of age or under.

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **5MG/ML GEL**

02238162 DIASTAT	VAE
09853340 DIASTAT 2X10MG RECTAL PACK	ELN
09853430 DIASTAT 2X15MG RECTAL PACK	ELN

LORAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **0.5MG TABLET**

00655740 APO-LORAZEPAM	APX
02041413 ATIVAN	PFI
02041456 ATIVAN SUBLINGUAL	PFI
02351072 LORAZEPAM	SAN
02410745 LORAZEPAM SUBLINGUAL	AAP
00728187 PMS-LORAZEPAM	PMS
00655643 PRO-LORAZEPAM	PDL
00711101 TEVA-LORAZEPAM	TEV

ST **1MG TABLET**

00655759 APO-LORAZEPAM	APX
02041421 ATIVAN	PFI
02041464 ATIVAN SUBLINGUAL	PFI
02351080 LORAZEPAM	SAN
02410753 LORAZEPAM SUBLINGUAL	AAP
00728195 PMS-LORAZEPAM	PMS
00655651 PRO-LORAZEPAM	PDL

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES**LORAZEPAM**

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 1MG TABLET

00637742 TEVA-LORAZEPAM TEV

ST 2MG TABLET

00655767 APO-LORAZEPAM APX

02041448 ATIVAN PFI

02041472 ATIVAN SUBLINGUAL PFI

02351099 LORAZEPAM SAN

02410761 LORAZEPAM SUBLINGUAL AAP

00728209 PMS-LORAZEPAM PMS

00655678 PRO-LORAZEPAM PDL

00637750 TEVA-LORAZEPAM TEV

NITRAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 5MG TABLET

00511528 MOGADON AAP

ST 10MG TABLET

00511536 MOGADON AAP

OXAZEPAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST 10MG TABLET

00402680 APO OXAZEPAM APX

00497754 OXAZEPAM PDL

00414247 OXPAM BMI

00568392 RIVA OXAZEPAM RIV

ST 15MG TABLET

00402745 APO OXAZEPAM APX

00497762 OXAZEPAM PDL

00568406 RIVA OXAZEPAM RIV

ST 30MG TABLET

00402737 APO OXAZEPAM APX

00497770 OXAZEPAM PDL

00414263 OXPAM BMI

00568414 RIVA OXAZEPAM RIV

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES**TEMAZEPAM**

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **15MG CAPSULE**

00604453 RESTORIL	AAP
02225964 TEMAZEPAM	APX
02229760 TEMAZEPAM	PDL
02230095 TEVA-TEMAZEPAM	TEV

ST **30MG CAPSULE**

00604461 RESTORIL	AAP
02225972 TEMAZEPAM	APX
02229761 TEMAZEPAM	PDL
02230102 TEVA-TEMAZEPAM	TEV

TRIAZOLAM

Limited use benefit (prior approval is not required).

To promote safe, therapeutically effective and efficient use of drug therapy NIHB has implemented a benzodiazepine dose limit of 40 mg diazepam equivalents per day. This limit will be calculated based on the total dose of all benzodiazepines a client is receiving from NIHB within a 100-day period (i.e. 4 000 diazepam equivalents over 100 days). According to the product monograph for diazepam, the recommended usual adult dosage is up to 40 mg per day.

The benzodiazepine dose limit will be further reduced to 30 mg DEQ per day. The new limit will be implemented region-by-region.

ST **0.25MG TABLET**

00808571 TRIAZOLAM	AAP
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28:32.28 SELECTIVE SEROTONIN AGONISTS**ALMOTRIPTAN MALATE**

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

6.25MG TABLET

02405792 APO-ALMOTRIPTAN	APX
02248128 AXERT	MCL
02398435 MYLAN-ALMOTRIPTAN	MYL

12.5MG TABLET

02424029 ALMOTRIPTAN	PDL
02466821 ALMOTRIPTAN	SAN
02405806 APO-ALMOTRIPTAN	APX
02248129 AXERT	MCL
02398443 MYLAN-ALMOTRIPTAN	MYL
02405334 SANDOZ ALMOTRIPTAN	SDZ
02434849 TEVA-ALMOTRIPTAN	TEV

NARATRIPTAN HYDROCHLORIDE

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

1MG TABLET

02237820 AMERGE	GSK
02314290 TEVA-NARATRIPTAN	TEV

2.5MG TABLET

02237821 AMERGE	GSK
02322323 SANDOZ NARATRIPTAN	SDZ

28:32.28 SELECTIVE SEROTONIN AGONISTS**NARATRIPTAN HYDROCHLORIDE**

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

2.5MG TABLET

02314304 TEVA-NARATRIPTAN

TEV

RIZATRIPTAN BENZOATE

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

5MG TABLET

02393468 APO-RIZATRIPTAN

APX

02380455 JAMP-RIZATRIPTAN

JMP

02429233 JAMP-RIZATRIPTAN IR

JMP

02379651 MAR-RIZATRIPTAN

MAR

02428512 VAN-RIZATRIPTAN

VAN

10MG TABLET

02381702 ACT RIZATRIPTAN

ACG

02393476 APO-RIZATRIPTAN

APX

02441144 AURO-RIZATRIPTAN

AUR

02380463 JAMP-RIZATRIPTAN

JMP

02429241 JAMP-RIZATRIPTAN IR

JMP

02379678 MAR-RIZATRIPTAN

MAR

02240521 MAXALT

FRS

02428520 VAN-RIZATRIPTAN

VAN

5MG TABLET (ORALLY DISINTEGRATING)

02374730 ACT RIZATRIPTAN ODT

ACG

02393484 APO-RIZATRIPTAN RPD

APX

02465086 JAMP-RIZATRIPTAN ODT

JMP

02462788 MAR-RIZATRIPTAN ODT

MAR

02240518 MAXALT RPD

FRS

02379198 MYLAN-RIZATRIPTAN ODT

MYL

02436604 NAT-RIZATRIPTAN ODT

NPH

02393360 PMS-RIZATRIPTAN RDT

PMS

02442906 RIZATRIPTAN ODT

SAN

02446111 RIZATRIPTAN ODT

SIV

02415798 RIZATRIPTAN RDT

PDL

02351870 SANDOZ RIZATRIPTAN ODT

SDZ

02396661 TEVA-RIZATRIPTAN ODT

TEV

10MG TABLET (ORALLY DISINTEGRATING)

02374749 ACT RIZATRIPTAN ODT

ACG

02393492 APO-RIZATRIPTAN RPD

APX

02396203 DOM-RIZATRIPTAN RDT

DPC

02465094 JAMP-RIZATRIPTAN ODT

JMP

02462796 MAR-RIZATRIPTAN ODT

MAR

02240519 MAXALT RPD

FRS

02379201 MYLAN-RIZATRIPTAN ODT

MYL

02436612 NAT-RIZATRIPTAN ODT

NPH

02393379 PMS-RIZATRIPTAN RDT

PMS

02442914 RIZATRIPTAN ODT

SAN

02446138 RIZATRIPTAN ODT

SIV

02415801 RIZATRIPTAN RDT

PDL

02351889 SANDOZ RIZATRIPTAN ODT

SDZ

28:32.28 SELECTIVE SEROTONIN AGONISTS**RIZATRIPTAN BENZOATE**

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

10MG TABLET (ORALLY DISINTEGRATING)

02396688	TEVA-RIZATRIPTAN ODT	TEV
02448505	VAN-RIZATRIPTAN ODT	VAN

SUMATRIPTAN SUCCINATE

Limited use benefit (prior approval is not required).

A total of 12 tablets (or injections) are permitted in a 30-day period.

6MG/0.5ML INJECTION

99000598	IMITREX STAT DOSE KIT	GSK
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12MG/ML SOLUTION

02212188	IMITREX	GSK
02361698	TARO-SUMATRIPTAN	TAR

25MG TABLET

02270749	DOM-SUMATRIPTAN	DPC
02268906	MYLAN-SUMATRIPTAN	MYL
02256428	PMS-SUMATRIPTAN	PMS
02286815	TEVA-SUMATRIPTAN DF	TEV

50MG TABLET

02268388	APO-SUMATRIPTAN	APX
02270757	DOM-SUMATRIPTAN	DPC
02212153	IMITREX DF	GSK
02268914	MYLAN-SUMATRIPTAN	MYL
02256436	PMS-SUMATRIPTAN	PMS
02263025	SANDOZ SUMATRIPTAN	SDZ
02286521	SUMATRIPTAN	SAN
02324652	SUMATRIPTAN	PDL
02385570	SUMATRIPTAN DF	SIV
02286823	TEVA-SUMATRIPTAN DF	TEV

100MG TABLET

02257904	ACT SUMATRIPTAN	ACG
02268396	APO-SUMATRIPTAN	APX
02270765	DOM-SUMATRIPTAN	DPC
02212161	IMITREX DF	GSK
02268922	MYLAN-SUMATRIPTAN	MYL
02256444	PMS-SUMATRIPTAN	PMS
02263033	SANDOZ SUMATRIPTAN	SDZ
02286548	SUMATRIPTAN	SAN
02324660	SUMATRIPTAN	PDL
02385589	SUMATRIPTAN DF	SIV
02239367	TEVA-SUMATRIPTAN	TEV
02286831	TEVA-SUMATRIPTAN DF	TEV

ZOLMITRIPTAN

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

2.5MG TABLET

02380951	APO-ZOLMITRIPTAN	APX
02389525	DOM-ZOLMITRIPTAN	DPC
02421623	JAMP-ZOLMITRIPTAN	JMP

28:32.28 SELECTIVE SEROTONIN AGONISTS**ZOLMITRIPTAN**

Limited use benefit (prior approval is not required).

A total of 12 tablets are permitted in a 30-day period.

2.5MG TABLET

02399458	MAR-ZOLMITRIPTAN	MAR
02419521	MINT-ZOLMITRIPTAN	MIN
02421534	NAT-ZOLMITRIPTAN	NPH
02324229	PMS-ZOLMITRIPTAN	PMS
02401304	RIVA-ZOLMITRIPTAN	RIV
02362988	SANDOZ ZOLMITRIPTAN	SDZ
02313960	TEVA-ZOLMITRIPTAN	TEV
02379929	ZOLMITRIPTAN	PDL
02238660	ZOMIG	AZC

2.5MG TABLET (ORALLY DISINTEGRATING)

02438453	AG-ZOLMITRIPTAN ODT	ANG
02381575	APO-ZOLMITRIPTAN RAPID	APX
02428237	JAMP-ZOLMITRIPTAN ODT	JMP
02324768	PMS-ZOLMITRIPTAN ODT	PMS
02362996	SANDOZ ZOLMITRIPTAN ODT	SDZ
02428474	SEPTA-ZOLMITRIPTAN-ODT	SPT
02342545	TEVA-ZOLMITRIPTAN OD	TEV
02438763	VAN-ZOLMITRIPTAN ODT	VAN
02379988	ZOLMITRIPTAN ODT	PDL
02243045	ZOMIG RAPIMELT	AZC

28:36.20 ANTIPARKINSONIAN AGENTS - DOPAMINE RECEPTOR AGONISTS**CABERGOLINE**

Limited use benefit (prior approval required).

For treatment of hyperprolactinemia in patients who have failed therapy with or are intolerant to bromocriptine.

0.5MG TABLET

02301407	ACT CABERGOLINE	ACG
02455897	APO-CABERGOLINE	APX
02242471	DOSTINEX	PFI

ROTIGOTINE

Limited use benefit (prior approval required).

As an adjunct to levodopa for the treatment of patients with advanced stage Parkinson's disease; AND Patient is currently receiving treatment with levodopa.

2MG PATCH

02403900	NEUPRO	UCB
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4MG PATCH

02403927	NEUPRO	UCB
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6MG PATCH

02403935	NEUPRO	UCB
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8MG PATCH

02403943	NEUPRO	UCB
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28:92.00 MISCELLANEOUS CENTRAL NERVOUS SYSTEM AGENTS**ACAMPROSATE CALCIUM**

Limited use benefit (prior approval required).

For patients who have been abstinent from alcohol for at least four days and where available, are currently enrolled in an alcohol addiction treatment program.

333MG TABLET (DELAYED RELEASE)

02293269 CAMPRAL

MYL

ATOMOXETINE HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with Attention Deficit Hyperactivity Disorder (ADHD) who meet one of the following criteria:

- Failure or intolerance to methylphenidate or amphetamine; OR
- Contraindication to stimulant medication; OR
- Potential risk of stimulant misuse or diversion; OR
- Prescribed or recommended by a pediatrician or a psychiatrist.

10MG CAPSULE

02318024 APO-ATOMOXETINE	APX
02358190 ATOMOXETINE	AAP
02396904 ATOMOXETINE	PDL
02445883 ATOMOXETINE	SIV
02467747 ATOMOXETINE	SAN
02390469 DOM-ATOMOXETINE	DPC
02381028 PMS-ATOMOXETINE	PMS
02405962 RIVA-ATOMOXETINE	RIV
02386410 SANDOZ ATOMOXETINE	SDZ
02262800 STRATTERA	LIL
02314541 TEVA-ATOMOXETINE	TEV

18MG CAPSULE

02318032 APO-ATOMOXETINE	APX
02358204 ATOMOXETINE	AAP
02396912 ATOMOXETINE	PDL
02445905 ATOMOXETINE	SIV
02467755 ATOMOXETINE	SAN
02390477 DOM-ATOMOXETINE	DPC
02381036 PMS-ATOMOXETINE	PMS
02405970 RIVA-ATOMOXETINE	RIV
02386429 SANDOZ ATOMOXETINE	SDZ
02262819 STRATTERA	LIL
02314568 TEVA-ATOMOXETINE	TEV

25MG CAPSULE

02318040 APO-ATOMOXETINE	APX
02358212 ATOMOXETINE	AAP
02396920 ATOMOXETINE	PDL
02445913 ATOMOXETINE	SIV
02467763 ATOMOXETINE	SAN
02390485 DOM-ATOMOXETINE	DPC
02381044 PMS-ATOMOXETINE	PMS
02405989 RIVA-ATOMOXETINE	RIV
02386437 SANDOZ ATOMOXETINE	SDZ
02262827 STRATTERA	LIL
02314576 TEVA-ATOMOXETINE	TEV

40MG CAPSULE

02318059 APO-ATOMOXETINE	APX
02358220 ATOMOXETINE	AAP
02396939 ATOMOXETINE	PDL

28:92.00 MISCELLANEOUS CENTRAL NERVOUS SYSTEM AGENTS**ATOMOXETINE HYDROCHLORIDE**

Limited use benefit (prior approval required).

For the treatment of patients with Attention Deficit Hyperactivity Disorder (ADHD) who meet one of the following criteria:

- Failure or intolerance to methylphenidate or amphetamine; OR
- Contraindication to stimulant medication; OR
- Potential risk of stimulant misuse or diversion; OR
- Prescribed or recommended by a pediatrician or a psychiatrist.

40MG CAPSULE

02445948	ATOMOXETINE	SIV
02467771	ATOMOXETINE	SAN
02390493	DOM-ATOMOXETINE	DPC
02381052	PMS-ATOMOXETINE	PMS
02405997	RIVA-ATOMOXETINE	RIV
02386445	SANDOZ ATOMOXETINE	SDZ
02262835	STRATTERA	LIL
02314584	TEVA-ATOMOXETINE	TEV

60MG CAPSULE

02318067	APO-ATOMOXETINE	APX
02358239	ATOMOXETINE	AAP
02396947	ATOMOXETINE	PDL
02445956	ATOMOXETINE	SIV
02467798	ATOMOXETINE	SAN
02390515	DOM-ATOMOXETINE	DPC
02381060	PMS-ATOMOXETINE	PMS
02406004	RIVA-ATOMOXETINE	RIV
02386453	SANDOZ ATOMOXETINE	SDZ
02262843	STRATTERA	LIL
02314592	TEVA-ATOMOXETINE	TEV

80MG CAPSULE

02318075	APO-ATOMOXETINE	APX
02358247	ATOMOXETINE	AAP
02467801	ATOMOXETINE	SAN
02404664	PMS-ATOMOXETINE	PMS
02422824	RIVA-ATOMOXETINE	RIV
02386461	SANDOZ ATOMOXETINE	SDZ
02279347	STRATTERA	LIL
02362511	TEVA-ATOMOXETINE	TEV

100MG CAPSULE

02318083	APO-ATOMOXETINE	APX
02358255	ATOMOXETINE	AAP
02467828	ATOMOXETINE	SAN
02404672	PMS-ATOMOXETINE	PMS
02422832	RIVA-ATOMOXETINE	RIV
02386488	SANDOZ ATOMOXETINE	SDZ
02279355	STRATTERA	LIL
02362538	TEVA-ATOMOXETINE	TEV

28:92.00 MISCELLANEOUS CENTRAL NERVOUS SYSTEM AGENTS**DIMETHYL FUMARATE**

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

120MG CAPSULE (DELAYED RELEASE)

02404508 TECFIDERA

UNK

240MG CAPSULE (DELAYED RELEASE)

02420201 TECFIDERA

UNK

32:00 CONTRACEPTIVES (NON-ORAL)**32:00.00 CONTRACEPTIVES (NON-ORAL)****INTRAUTERINE DEVICE**

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 1 device every 12 months.

DEVICE

00970328 FLEXI-T +300 IUD

TSN

00970336 FLEXI-T +380 IUD

TSN

98099999 FLEXI-TD

TSN

99401085 LIBERTE UT380 SHORT IUD

MSF

99401086 LIBERTE UT380 STANDARD IUD

MSF

99400482 NOVA-T

BEX

36:00 DIAGNOSTIC AGENTS (DX)**36:26.00 DX - DIABETES MELLITUS****GLUCOSE OXIDASE, PEROXIDASE**

Limited use benefit (prior approval not required).

The number of test strips that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 500 test strips per 100 days. A client can test up to five times per day.
- Clients managing diabetes with diabetes medication with a high risk of causing low blood sugar will be allowed 400 test strips per 365 days. A client can test once daily.
- Clients managing diabetes with diabetes medication with a low risk of causing low blood sugar will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Clients managing diabetes with diet/lifestyle therapy only (no insulin or diabetes medications) will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Non-diabetic clients with rare conditions leading to symptomatic hypo or hyperglycemia will be allowed 500 test strips per 100 days.

STRIP

09857563 ACCU-CHEK GUIDE (ON)

ROD

97799177 ACCU-CHEK GUIDE (SK)

ROD

ACCU-CHEK ADVANTAGE STRIP

09853626 ACCU-CHEK ADVANTAGE

ROD

97799824 ACCU-CHEK ADVANTAGE

ROD

ACCU-CHEK AVIVA STRIP

09857178 ACCU-CHEK AVIVA

ROD

97799814 ACCU-CHEK AVIVA

ROD

ACCU-CHEK COMPACT STRIP

09854282 ACCU-CHEK COMPACT

ROD

97799962 ACCU-CHEK COMPACT

ROD

ACCU-CHEK MOBILE STRIP

09857452 ACCU-CHEK MOBILE BG

ROD

36:26.00 DX - DIABETES MELLITUS**GLUCOSE OXIDASE, PEROXIDASE**

Limited use benefit (prior approval not required).

The number of test strips that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 500 test strips per 100 days. A client can test up to five times per day.
- Clients managing diabetes with diabetes medication with a high risk of causing low blood sugar will be allowed 400 test strips per 365 days. A client can test once daily.
- Clients managing diabetes with diabetes medication with a low risk of causing low blood sugar will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Clients managing diabetes with diet/lifestyle therapy only (no insulin or diabetes medications) will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Non-diabetic clients with rare conditions leading to symptomatic hypo or hyperglycemia will be allowed 500 test strips per 100 days.

ACCU-CHEK MOBILE STRIP

97799497 ACCU-CHEK MOBILE CASSETT ROD

ACCUTREND STRIP

09853162 ACCUTREND ROD

97799959 ACCUTREND ROD

ASCENSIA BREEZE 2 STRIP

97799748 ASCENSIA BREEZE 2 BAY

09857293 BREEZE 2 BG (ON) BAY

ASCENSIA CONTOUR STRIP

97799702 ASCENSIA CONTOUR BAY

09857127 CONTOUR BG (ON) BAY

BG STAR STRIP

97799465 BG STAR SAC

09857422 BG STAR (ON) SAC

CONTOUR NEXT STRIP

97799459 CONTOUR NEXT BAY

09857453 CONTOUR NEXT (ON) BAY

EZ HEALTH STRIP

09857357 EZ HEALTH ORACLE TRE

97799564 EZ HEALTH ORACLE TRE

FREESTYLE STRIP

97799829 FREESTYLE ABB

09857141 FREESTYLE (ON) ABB

FREESTYLE LITE STRIP

97799597 FREESTYLE LITE ABB

09857297 FREESTYLE LITE (ON) ABB

FREESTYLE PRECISION STRIP

97799346 FREESTYLE PRECISION ABB

09857502 FREESTYLE PRECISION (ON) ABB

GE200 STRIP

97799373 GE200 AUC

09857525 GE200 (ON) AUC

ITEST STRIP

09857348 ITEST AUC

97799692 ITEST AUC

MEDI+SURE STRIP

97799403 MEDI+SURE MEC

09857432 MEDI+SURE (ON) MEC

NOVA MAX STRIP

09857313 NOVA MAX NCA

36:26.00 DX - DIABETES MELLITUS**GLUCOSE OXIDASE, PEROXIDASE**

Limited use benefit (prior approval not required).

The number of test strips that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 500 test strips per 100 days. A client can test up to five times per day.
- Clients managing diabetes with diabetes medication with a high risk of causing low blood sugar will be allowed 400 test strips per 365 days. A client can test once daily.
- Clients managing diabetes with diabetes medication with a low risk of causing low blood sugar will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Clients managing diabetes with diet/lifestyle therapy only (no insulin or diabetes medications) will be allowed 200 test strips per 365 days. A client can test three to four times per week.
- Non-diabetic clients with rare conditions leading to symptomatic hypo or hyperglycemia will be allowed 500 test strips per 100 days.

ONE TOUCH ULTRA STRIP

09854290 ONE TOUCH ULTRA	JAJ
97799985 ONE TOUCH ULTRA	JAJ

ONE TOUCH VERIO STRIP

97799475 ONETOUCH VERIO	JAJ
09857392 ONETOUCH VERIO (ON)	JAJ

PRECISION XTRA STRIP

09854070 PRECISION XTRA	ABB
97799840 PRECISION XTRA	AUC

SIDEKICK STRIP

97799601 SIDEKICK	HOD
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SPIRIT STRIP

97799291 FIRST CANHEALTH SPIRIT	ARA
09857547 SPIRIT TEST STRIP (ON)	ARA

SURE STEP STRIP

97799355 SURE STEP	SKY
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SURETEST STRIP

09857522 SURETEST (ON)	SKY
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TRUETEST STRIP

97799532 TRUETEST	HOD
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TRUETRACK STRIP

09857283 TRUE TRACK	AUC
97799602 TRUE TRACK	HOD

40:00 ELECTROLYTIC, CALORIC, AND WATER BALANCE**40:18.19 PHOSPHATE - REMOVING AGENTS****LANTHANUM CARBONATE HYDRATE**

Limited use benefit (prior approval required).

For patients with elevated phosphate levels OR elevated phosphate X calcium product despite dietary restriction of phosphate and use of calcium-based phosphate binders (short term elevations should be managed with aluminium based binders).

For patients with elevated calcium levels despite discontinuation of calcium binder, and Vitamin D analogue and/or modification of dialysate calcium.

For patients with adynamic bone disease and low PTH levels (<100 pg/ml or <0.9 pmol/L) with normal or elevated calcium levels.

250MG TABLET (CHEWABLE)

02287145 FOSRENOL	UNK
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500MG TABLET (CHEWABLE)

02287153 FOSRENOL	UNK
-------------------	-----

750MG TABLET (CHEWABLE)

02287161 FOSRENOL	UNK
-------------------	-----

1000MG TABLET (CHEWABLE)

02287188 FOSRENOL	UNK
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40:18.19 PHOSPHATE - REMOVING AGENTS**SEVELAMER CARBONATE**

Limited use benefit (prior approval required).

For patients with elevated phosphate levels OR elevated phosphate X calcium product despite dietary restriction of phosphate and use of calcium-based phosphate binders (short term elevations should be managed with aluminium based binders).

For patients with elevated calcium levels despite discontinuation of calcium binder, and Vitamin D analogue and/or modification of dialysate calcium.

For patients with adynamic bone disease and low PTH levels (<100 pg/ml or <0.9 pmol/L) with normal or elevated calcium levels.

800MG TABLET

02461501 ACCEL-SEVELAMER

ACP

02354586 RENVELA

SAC

SEVELAMER HYDROCHLORIDE

Limited use benefit (prior approval required).

For patients with elevated phosphate levels OR elevated phosphate X calcium product despite dietary restriction of phosphate and use of calcium-based phosphate binders (short term elevations should be managed with aluminium based binders).

For patients with elevated calcium levels despite discontinuation of calcium binder, and Vitamin D analogue and/or modification of dialysate calcium.

For patients with adynamic bone disease and low PTH levels (<100 pg/ml or <0.9 pmol/L) with normal or elevated calcium levels.

800MG TABLET

02244310 RENAGEL

SAC

40:20.00 CALORIC AGENTS**LEVOCARNITINE**

Limited use benefit (prior approval required).

For treatment of carnitine deficiency.

100MG/ML SOLUTION

02144336 CARNITOR

UNK

200MG/ML SOLUTION

02144344 CARNITOR

UNK

330MG TABLET

02144328 CARNITOR

UNK

48:00 RESPIRATORY TRACT AGENTS**48:02.00 ANTIFIBROTIC AGENTS****NINTEDANIB ESILATE**

Limited use benefit (prior approval required).

Initial Request - Coverage is provided for a period of 7 months (6 months plus a 4 week allowance for repeat pulmonary function tests):

For the treatment of adult patients with a diagnosis of mild to moderate idiopathic pulmonary fibrosis (IPF) who meet the following criteria:

- Diagnosis confirmed by a respirologist and a high-resolution CT scan within the previous 24 months; AND
- All other causes of restrictive lung disease (e.g. collagen vascular disorder or hypersensitivity pneumonitis) should be excluded; AND
- Mild to moderate IPF is defined as forced vital capacity (FVC) greater than or equal to 50% of predicted; AND
- Patient is under the care of a physician with experience in IPF.

Renewal at 6 months - Coverage is provided for a period of 6 months:

- Patients must NOT demonstrate progression of disease defined as an absolute decline in percent predicted 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 Renewals at 12 months and thereafter - Coverage is provided for a period of 12 months:

- 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.

Combination use of Ofev (nintedanib) and Esbriet (pirfenidone) will not be provided.

100MG CAPSULE

02443066 OFEV

BOE

150MG CAPSULE

02443074 OFEV

BOE

48:02.00 ANTIFIBROTIC AGENTS**PIRFENIDONE**

Limited use benefit (prior approval required).

Initial Request - Coverage is provided for a period of 7 months (6 months plus a 4 weeks allowance for repeat pulmonary function tests):

For the treatment of adult patients with a diagnosis of mild to moderate idiopathic pulmonary fibrosis (IPF) who meet the following criteria:

- Diagnosis confirmed by a respirologist and a high-resolution CT scan within the previous 24 months; AND
- All other causes of restrictive lung disease (e.g. collagen vascular disorder or hypersensitivity pneumonitis) should be excluded; AND
- Mild to moderate IPF is defined as forced vital capacity (FVC) greater than or equal to 50% of predicted; AND
- Patient is under the care of a physician with experience in IPF.

Renewal at 6 months - Coverage is provided for a period of 6 months:

- Patients must NOT demonstrate progression of disease defined as an absolute decline in percent predicted 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 Renewals at 12 months and thereafter - Coverage is provided for a period of 12 months:

- 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.

Combination use of Ofev (nintedanib) and Esbriet (pirfenidone) will not be provided.

267MG CAPSULE

02393751 ESBRIET

HLR

267MG TABLET

02464489 ESBRIET

HLR

801MG TABLET

02464500 ESBRIET

HLR

48:10.24 LEUKOTRIENE MODIFIERS**MONTELUKAST SODIUM**

Limited use benefit (prior approval required).

For treatment of:

- asthma when used in patients on concurrent steroid therapy; OR
- asthma patients not well controlled with or intolerant to inhaled corticosteroids.

Montelukast is open benefit for children up to 17 years of age.

ST 4MG GRANULES

02358611 SANDOZ MONTELUKAST

SDZ

02247997 SINGULAIR

FRS

ST 10MG TABLET

02374609 APO-MONTELUKAST

APX

02401274 AURO-MONTELUKAST

AUR

02445735 BIO-MONTELUKAST

UNK

02376695 DOM-MONTELUKAST

DPC

02391422 JAMP-MONTELUKAST

JMP

02399997 MAR-MONTELUKAST

MAR

02408643 MINT-MONTELUKAST

MIN

02379333 MONTELUKAST

SAN

02379856 MONTELUKAST

PDL

02382474 MONTELUKAST

SIV

02379236 MONTELUKAST SODIUM

ACC

02373947 PMS-MONTELUKAST

PMS

02389517 RAN-MONTELUKAST

RBV

02398826 RIVA-MONTELUKAST

RIV

02328593 SANDOZ MONTELUKAST

SDZ

02238217 SINGULAIR

FRS

02355523 TEVA-MONTELUKAST

TEV

4MG TABLET (CHEWABLE)

02377608 APO-MONTELUKAST

APX

48:10.24 LEUKOTRIENE MODIFIERS**MONTELUKAST SODIUM**

Limited use benefit (prior approval required).

For treatment of:

- asthma when used in patients on concurrent steroid therapy; OR
- asthma patients not well controlled with or intolerant to inhaled corticosteroids.

Montelukast is open benefit for children up to 17 years of age.

4MG TABLET (CHEWABLE)

02422867 AURO-MONTELUKAST	AUR
02442353 JAMP-MONTELUKAST	JMP
02399865 MAR-MONTELUKAST	MAR
02408627 MINT-MONTELUKAST	MIN
02379317 MONTELUKAST	SAN
02379821 MONTELUKAST	PDL
02382458 MONTELUKAST	SIV
02354977 PMS-MONTELUKAST	PMS
02402793 RAN-MONTELUKAST	RBY
02330385 SANDOZ MONTELUKAST	SDZ
02243602 SINGULAIR	FRS
02355507 TEVA-MONTELUKAST	TEV

ST 5MG TABLET (CHEWABLE)

02377616 APO-MONTELUKAST	APX
02422875 AURO-MONTELUKAST	AUR
02442361 JAMP-MONTELUKAST	JMP
02399873 MAR-MONTELUKAST	MAR
02408635 MINT-MONTELUKAST	MIN
02379325 MONTELUKAST	SAN
02379848 MONTELUKAST	PDL
02382466 MONTELUKAST	SIV
02354985 PMS-MONTELUKAST	PMS
02402807 RAN-MONTELUKAST	RBY
02330393 SANDOZ MONTELUKAST	SDZ
02238216 SINGULAIR	FRS
02355515 TEVA-MONTELUKAST	TEV

48:48.00 VASODILATING AGENTS**AMBRISENTAN**

Limited use benefit (prior approval required).

Maximum dose covered is 10 mg once daily.

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST 5MG TABLET

02475375 APO-AMBRISENTAN	APX
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ST 10MG TABLET

02475383 APO-AMBRISENTAN	APX
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48:48.00 VASODILATING AGENTS**BOSENTAN MONOHYDRATE**

Limited use benefit (prior approval required).

Maximum dose covered is 125 mg twice daily

Patients with World Health Organization (WHO) class III pulmonary artery hypertension (PAH), either idiopathic (i.e. primary) or associated with a congenital or systemic condition (e.g. connective tissue disease) and confirmed by right heart catheterization; AND

- who have failed to respond to sildenafil OR tadalafil; OR
- who have contraindications to sildenafil OR tadalafil.

ST **125MG TABLET**

02399210 APO-BOSENTAN

APX

RIOCIQUAT

Limited use benefit (prior approval required).

For the treatment of patients 18 years of age or older with chronic thromboembolic pulmonary hypertension (CTEPH) with World Health Organization (WHO) Functional Class 2 or 3 pulmonary hypertension with:

- Inoperable CTEPH, World Health Organization (WHO) Group 4;
- OR
- Persistent or recurrent CTEPH after surgical treatment; AND
- Prescriber experienced in the diagnosis and treatment of CTEPH.

0.5MG TABLET

02412764 ADEMPAS

BAY

1MG TABLET

02412772 ADEMPAS

BAY

1.5MG TABLET

02412799 ADEMPAS

BAY

2MG TABLET

02412802 ADEMPAS

BAY

2.5MG TABLET

02412810 ADEMPAS

BAY

SELEXIPAG

Limited use benefit (prior approval required).

For the treatment of adult patients with World Health Organization (WHO) functional class (FC) II to III pulmonary arterial hypertension (PAH), including idiopathic PAH, heritable PAH, PAH associated with connective tissue disorders or PAH associated with congenital heart disease:

- Patient is under the care of a physician with experience in the diagnosis and treatment of PAH; AND
- Patient has failed to respond to first- and second-line PAH therapies; OR
- Patient has contraindications/intolerance to first- and second-line PAH therapies.

200MCG TABLET

02451158 UPTRAVI

JSO

400MCG TABLET

02451166 UPTRAVI

JSO

600MCG TABLET

02451174 UPTRAVI

JSO

800MCG TABLET

02451182 UPTRAVI

JSO

1000MCG TABLET

02451190 UPTRAVI

JSO

1200MCG TABLET

02451204 UPTRAVI

JSO

1400MCG TABLET

02451212 UPTRAVI

JSO

1600MCG TABLET

02451220 UPTRAVI

JSO

48:92.00 MISCELLANEOUS RESPIRATORY TRACT AGENTS**OMALIZUMAB**

Limited use benefit (prior approval required).

Coverage is provided for an initial period of 24 weeks at a maximum dose of 300 mg every 4 weeks (6 injections over a 24 week period).

1. For the treatment of adults and adolescents (12 years of age or older) with moderate to severe chronic idiopathic urticaria (CIU) who remain symptomatic (presence of hives and/or associated itching) despite optimum management with H1 antihistamines; AND Prescriber is experienced in the treatment of CIU (Allergist, Dermatologist, Immunologist, OR other authorized prescriber experienced in the treatment of CIU).

Treatment cessation could be considered for patients who experience complete symptom control (UAS-7 = 0) for at least 12 consecutive weeks at the end of a 24-week treatment period.

Renewal coverage is provided for 24 weeks at a maximum dose of 300 mg every 4 weeks (6 injections/24 weeks).

2. For the treatment of adults and adolescents (12 years of age or older) with moderate to severe chronic idiopathic urticaria (CIU); AND

Patient stopped omalizumab after achieving complete symptom control (UAS-7 = 0) for at least 12 weeks while on treatment, but has experienced symptom relapse; OR

Patient achieved complete symptom control, but for a period of less than 12 consecutive weeks; OR

Patient achieved a partial response to treatment, defined as a ≥ 9.5 -point reduction in baseline urticaria activity score over 7 days (UAS-7).

In patients where treatment is discontinued due to temporary symptom control, treatment re-initiation may be considered should CIU symptoms reappear.

150MG POWDER FOR SOLUTION

02260565 XOLAIR

NVR

52:00 EYE, EAR, NOSE AND THROAT (EENT) PREPARATIONS**52:28.00 EENT - MOUTHWASHES AND GARGLES****BENZYDAMINE HYDROCHLORIDE**

Limited use benefit (prior approval required).

- For the treatment of radiation mucositis and oral ulcerative complications of chemotherapy.
- For use in immunocompromised patients who are at risk of mucosal breakdown.

0.15% MOUTHWASH

02239044 APO-BENZYDAMINE

APX

02229777 PHARIXIA

PED

02239537 PMS-BENZYDAMINE

PMS

52:92.00 MISCELLANEOUS EENT DRUGS**AFLIBERCEPT**

Limited use benefit (prior approval required).

For the treatment of:
 Diabetic Macular Edema (DME)
 Wet Age-Related Macular Degeneration (w-AMD)
 Retinal Vein Occlusion (RVO)

Criteria for coverage of aflibercept (Eylea) for DME, RVO and w-AMD:

- Administered by a qualified ophthalmologist experienced in intravitreal injections
- Interval between doses not shorter than 1 month

Note: Coverage will be limited to a maximum of 1 vial of Eylea per eye treated every 30 days

1. For the treatment of diabetic macular edema (DME) for patients who meet the following:

- Clinically significant diabetic macular edema for whom laser photocoagulation is also indicated; AND
- Have a hemoglobin A1c of less than 12%

2. Initial Coverage for the treatment of neovascular wet age-related macular degeneration (wAMD) where all of the following apply to the eye to be treated:

- Best Corrected Visual Acuity (BCVA) is between 6/12 and 6/96
- The lesion size is less than or equal to 12 disc areas in greatest linear dimension
- There is evidence of recent (< 3 months) presumed disease progression (blood vessel growth, as indicated by fluorescein angiography, or optical coherence tomography (OCT))

Note: Coverage will not be approved for patients:

- With permanent retinal damage as defined by the Royal College of Ophthalmology guidelines.
- Receiving concurrent treatment with verteporfin

Continued Coverage:

Treatment with Eylea for wAMD should be continued only in people who maintain adequate response to therapy

Treatment with Eylea should be permanently discontinued if any one of the following occurs:

- Reduction in BCVA in the treated eye to less than 15 letters (absolute) on two (2) consecutive visits in the treated eye, attributed to AMD in the absence of other pathology
- Reductions in BCVA of 30 letters or more compared to either baseline and/or best recorded level since baseline as this may indicate either poor treatment effect, adverse events or both.
- There is evidence of deterioration of the lesion morphology despite optimum treatment over three (3) consecutive visits.

3. For the treatment of RVO for patients who meet one of the following:

- Clinically significant macular edema secondary to branch retinal vein occlusion (BRVO); OR
- Central retinal vein occlusion (CRVO).
- It is recommended that Eylea be administered once every month. The interval between two doses should not be shorter than one month. The treatment interval may be extended up to 3 months based on visual and anatomic outcomes. Prescribers are advised to periodically assess (every 1 to 2 months) the need for continued therapy.
- Treatment with anti-VEGF agents should only be continued in patients who maintain adequate response to therapy.

40MG SOLUTION

02415992 EYLEA

BAY

52:92.00 MISCELLANEOUS EENT DRUGS**RANIBIZUMAB**

Limited use benefit (prior approval required).

For the treatment of:
 Diabetic Macular Edema (DME)
 Wet Age-Related Macular Degeneration (w-AMD)
 Retinal Vein Occlusion (RVO)
 Choroidal Neovascularization secondary to pathologic myopia (mCNV)

Criteria for coverage of ranibizumab (Lucentis) for DME, RVO, mCNV and w-AMD:

- Administered by a qualified ophthalmologist experienced in intravitreal injections
- Interval between doses not shorter than 1 month

Note: Coverage will be limited to a maximum of 1 vial of Lucentis per eye treated every 30 days

1. For the treatment of diabetic macular edema (DME) for patients who meet the following:

- Clinically significant diabetic macular edema for whom laser photocoagulation is also indicated; AND
- Have a hemoglobin A1c of less than 11%

2. Initial Coverage for the treatment of neovascular wet age-related macular degeneration (wAMD) where all of the following apply to the eye to be treated:

- Best Corrected Visual Acuity (BCVA) is between 6/12 and 6/96
- The lesion size is less than or equal to 12 disc areas in greatest linear dimension
- There is evidence of recent (< 3 months) presumed disease progression (blood vessel growth, as indicated by fluorescein angiography, or optical coherence tomography (OCT))

Note: Coverage will not be approved for patients:

- With permanent retinal damage as defined by the Royal College of Ophthalmology guidelines.
- Receiving concurrent treatment with verteporfin

Continued Coverage:

Treatment with Lucentis for wAMD should be continued only in people who maintain adequate response to therapy

Treatment with Lucentis should be permanently discontinued if any one of the following occurs:

- Reduction in BCVA in the treated eye to less than 15 letters (absolute) on two (2) consecutive visits in the treated eye, attributed to AMD in the absence of other pathology
- Reductions in BCVA of 30 letters or more compared to either baseline and/or best recorded level since baseline as this may indicate either poor treatment effect, adverse events or both.
- There is evidence of deterioration of the lesion morphology despite optimum treatment over three (3) consecutive visits.

3. For the treatment of RVO for patients who meet one of the following:

- Clinically significant macular edema secondary to branch retinal vein occlusion (BRVO); OR
- Central retinal vein occlusion (CRVO).
- Treatment to be given monthly and continued until maximum visual acuity is achieved, confirmed by stable visual acuity for three consecutive monthly assessments performed while on ranibizumab treatment. Thereafter patients should be monitored monthly for visual acuity.
- Treatment is resumed with monthly injections when monitoring indicates a loss of visual acuity due to macular edema secondary to retinal vein occlusion and continued until stable visual acuity is reached again for three consecutive monthly assessments.
- Treatment with anti-VEGF agents should only be continued in patients who maintain adequate response to therapy.

4. For the treatment of mCNV for patients who meet the following:

- Visual impairment due to choroidal neovascularization secondary to pathologic myopia (mCNV).

Treatment is initiated with a single intravitreal injection. Monitoring is recommended monthly for the first two months and at least every three months thereafter during the first year. If monitoring reveals signs of disease activity (e.g. reduced visual acuity and/or signs of lesion activity), further treatment is recommended at a frequency of 1 injection per month until no disease activity is seen.

10MG/ML SOLUTION

02296810 LUCENTIS

NVR

02425629 LUCENTIS PFS

NVR

VERTEPORFIN

Limited use benefit (prior approval required).

For treatment of age related macular degeneration for patients with this diagnosis who are being treated by a certified ophthalmologist.

15MG/VIAL POWDER FOR SOLUTION

02242367 VISUDYNE

VAE

56:00 GASTROINTESTINAL DRUGS**56:04.00 ANTACIDS AND ADSORBENTS****BISMUTH SUBSALICYLATE**

Limited use benefit (prior approval not required).

Coverage will be limited to 8 tablets a day every 14 days, followed by a 28 day lockout;
OR

Coverage will be limited to 120mL a day every 14 days, followed by a 28 day lockout.

262MG CAPLET

00245730 BISMUTH

JMP

17.6MG/ML SUSPENSION

02097079 PEPTO BISMOL

PGI

262MG TABLET

02326582 BISMUTH SUBSALICYLATE

UNK

02177994 PEPTO BISMOL

PGI

56:22.08 ANTIHISTAMINES**DIMENHYDRINATE**

Limited use benefit (prior approval not required).

The NIHB Program implemented a dose coverage limit for DIMENHYDRINATE in June 2017 as part of a strategy to address safety concerns and potential misuse.

The dimenhydrinate dose limit is currently 400 mg per day for a total of 12,000 mg of dimenhydrinate in a 30-day period.

This limit applies only to the 15 mg and 50 mg tablets. Dimenhydrinate in liquid, suppository and injectable forms are not included in this limit.

50MG TABLET

02241532 ANTI-NAUSEANT

VTH

00363766 APO DIMENHYDRINATE

APX

00013803 GRAVOL

CHU

02245416 JAMP-DIMENHYDRINATE

JMP

02377179 MOTION SICKNESS

APX

00586331 PMS-DIMENHYDRINATE

PMS

00605786 TRAVEL

VTH

00021423 TRAVEL ON

NOP

56:22.32 MISCELLANEOUS ANTIEMETICS**APREPITANT**

Limited use benefit (prior approval required).

When used in combination with a 5-HT3 antagonist and dexamethasone for the prevention of acute and delayed nausea and vomiting due to highly emetogenic cancer chemotherapy (e.g. Cisplatin > 70mg/m2).

ST **80MG CAPSULE**

02298791 EMEND

FRS

ST **125MG CAPSULE**

02298805 EMEND

FRS

ST **125MG & 80MG CAPSULE**

02298813 EMEND TRI-PACK

FRS

56:22.92 MISCELLANEOUS ANTIEMETICS**NABILONE**

Limited use benefit (prior approval required).

For patients who are experiencing nausea and vomiting due to cancer chemotherapy or radiation;

OR

Patient is palliative (diagnosed with a terminal illness or disease which is expected to be the primary cause of death within six months or less).

0.25MG CAPSULE

02312263 CESAMET

VAE

02358077 RAN-NABILONE

RBY

56:22.92 MISCELLANEOUS ANTIEMETICS**NABILONE**

Limited use benefit (prior approval required).

For patients who are experiencing nausea and vomiting due to cancer chemotherapy or radiation;

OR

Patient is palliative (diagnosed with a terminal illness or disease which is expected to be the primary cause of death within six months or less).

0.25MG CAPSULE

02392925 TEVA-NABILONE TEV

0.5MG CAPSULE

02393581 ACT NABILONE ACG

02256193 CESAMET VAE

02380900 PMS-NABILONE PMS

02358085 RAN-NABILONE RBY

02384884 TEVA-NABILONE TEV

1MG CAPSULE

02393603 ACT NABILONE ACG

00548375 CESAMET VAE

02380919 PMS-NABILONE PMS

02358093 RAN-NABILONE RBY

02384892 TEVA-NABILONE TEV

56:28.36 PROTON-PUMP INHIBITORS**LANSOPRAZOLE**

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

The following PPI status change is primarily based on the Canadian Optimal Medication Prescribing and Utilization Service (COMPUS) report on optimal PPI therapy. The report concluded that:

- All PPIs are equally efficacious
- Double dose PPI is not necessary for initial therapy
- Double dose PPI is effective in H. Pylori eradication; however, treatment is not needed beyond 14 days.

PPI use has been associated with increased risk of hip fracture, community-acquired pneumonia and Clostridium difficile associated diarrhea. Although further study is needed to establish clinical significance, it is prudent to use the lowest dose and shortest duration of therapy required to control symptoms.

All proton pump inhibitors (open benefit and limited use (LU) PPIs) have a maximum quantity limit of 400 tablets/capsules per 180 day period. This quantity limit will be in effect for the entire class of PPIs.

- For example, if a patient fills 30 tablets of rabeprazole, then switch to 30 tablets of omeprazole, then switch to 30 capsules of lansoprazole, this will count as 90 PPI tablets/capsules towards the quantity limit.
- Patients taking two rabeprazole 10mg tablets a day can be switched to one rabeprazole 20mg tablet a day to avoid reaching the quantity limit
- Patients taking two omeprazole 10mg tablets/capsules a day can be switched to one omeprazole 20mg tablet/capsule a day to avoid reaching the quantity limit

Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST 15MG CAPSULE (DELAYED RELEASE)

02293811 APO-LANSOPRAZOLE APX

02357682 LANSOPRAZOLE SAN

02385767 LANSOPRAZOLE SIV

02433001 LANSOPRAZOLE PMS

02353830 MYLAN-LANSOPRAZOLE MYL

02395258 PMS-LANSOPRAZOLE PMS

02165503 PREVACID TAK

02402610 RAN-LANSOPRAZOLE RBY

02422808 RIVA-LANSOPRAZOLE RIV

02385643 SANDOZ LANSOPRAZOLE SDZ

02280515 TEVA-LANSOPRAZOLE TEV

ST 30MG CAPSULE (DELAYED RELEASE)

02293838 APO-LANSOPRAZOLE APX

02414775 DOM-LANSOPRAZOLE DPC

56:28.36 PROTON-PUMP INHIBITORS**LANSOPRAZOLE**

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

The following PPI status change is primarily based on the Canadian Optimal Medication Prescribing and Utilization Service (COMPUS) report on optimal PPI therapy. The report concluded that:

- All PPIs are equally efficacious
- Double dose PPI is not necessary for initial therapy
- Double dose PPI is effective in H. Pylori eradication; however, treatment is not needed beyond 14 days.

PPI use has been associated with increased risk of hip fracture, community-acquired pneumonia and Clostridium difficile associated diarrhea. Although further study is needed to establish clinical significance, it is prudent to use the lowest dose and shortest duration of therapy required to control symptoms.

All proton pump inhibitors (open benefit and limited use (LU) PPIs) have a maximum quantity limit of 400 tablets/capsules per 180 day period. This quantity limit will be in effect for the entire class of PPIs.

- For example, if a patient fills 30 tablets of rabeprazole, then switch to 30 tablets of omeprazole, then switch to 30 capsules of lansoprazole, this will count as 90 PPI tablets/capsules towards the quantity limit.
- Patients taking two rabeprazole 10mg tablets a day can be switched to one rabeprazole 20mg tablet a day to avoid reaching the quantity limit
- Patients taking two omeprazole 10mg tablets/capsules a day can be switched to one omeprazole 20mg tablet/capsule a day to avoid reaching the quantity limit

Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST 30MG CAPSULE (DELAYED RELEASE)

02357690	LANSOPRAZOLE	SAN
02366282	LANSOPRAZOLE	PDL
02410389	LANSOPRAZOLE	SIV
02433028	LANSOPRAZOLE	PMS
02353849	MYLAN-LANSOPRAZOLE	MYL
02395266	PMS-LANSOPRAZOLE	PMS
02165511	PREVACID	TAK
02402629	RAN-LANSOPRAZOLE	RBY
02422816	RIVA-LANSOPRAZOLE	RIV
02280523	TEVA-LANSOPRAZOLE	TEV

ST 30MG TABLET (DELAYED RELEASE)

02385651	SANDOZ LANSOPRAZOLE	SDZ
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LANSOPRAZOLE ODT

Limited use benefit (prior approval required).

Coverage will be limited to 400 tablets/capsules every 180 days.

For children 12 years of age or under who are unable to swallow the capsule formulation; OR
For patients with dysphagia or a feeding tube when the use of the capsule formulation is not possible.

The following PPI status change is primarily based on the Canadian Optimal Medication Prescribing and Utilization Service (COMPUS) report on optimal PPI therapy. The report concluded that:

- All PPIs are equally efficacious
- Double dose PPI is not necessary for initial therapy
- Double dose PPI is effective in H. Pylori eradication; however, treatment is not needed beyond 14 days.

PPI use has been associated with increased risk of hip fracture, community-acquired pneumonia and Clostridium difficile associated diarrhea. Although further study is needed to establish clinical significance, it is prudent to use the lowest dose and shortest duration of therapy required to control symptoms.

All proton pump inhibitors (open benefit and limited use (LU) PPIs) have a maximum quantity limit of 400 tablets/capsules per 180 day period. This quantity limit will be in effect for the entire class of PPIs.

- For example, if a patient fills 30 tablets of rabeprazole, then switch to 30 tablets of omeprazole, then switch to 30 capsules of lansoprazole, this will count as 90 PPI tablets/capsules towards the quantity limit.
- Patients taking two rabeprazole 10mg tablets a day can be switched to one rabeprazole 20mg tablet a day to avoid reaching the quantity limit
- Patients taking two omeprazole 10mg tablets/capsules a day can be switched to one omeprazole 20mg tablet/capsule a day to avoid reaching the quantity limit

Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST 15MG TABLET (DELAYED RELEASE)

02249464	PREVACID FASTAB	TAK
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ST 30MG TABLET (DELAYED RELEASE)

02249472	PREVACID FASTAB	TAK
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56:28.36 PROTON-PUMP INHIBITORS**OMEPRAZOLE MAGNESIUM**

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

The following PPI status change is primarily based on the Canadian Optimal Medication Prescribing and Utilization Service (COMPUS) report on optimal PPI therapy. The report concluded that:

- All PPIs are equally efficacious
- Double dose PPI is not necessary for initial therapy
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Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST 20MG CAPSULE (DELAYED RELEASE)

02245058	APO-OMEPRAZOLE	APX
00846503	LOSEC	AZC
02339927	OMEPRAZOLE	PDL
02348691	OMEPRAZOLE	SAN
02411857	OMEPRAZOLE-20	SIV
02320851	PMS-OMEPRAZOLE	PMS
02403617	RAN-OMEPRAZOLE	RBY
02296446	SANDOZ OMEPRAZOLE	SDZ

20MG TABLET (DELAYED RELEASE)

02449927	BIO-OMEPRAZOLE	BMI
02420198	JAMP-OMEPRAZOLE DR	JMP
02190915	LOSEC	AZC
02439549	NAT-OMEPRAZOLE DR	NPH
02416549	OMEPRAZOLE	ACC
02374870	RAN-OMEPRAZOLE	RBY
02402416	RIVA-OMEPRAZOLE DR	RIV
02295415	TEVA-OMEPRAZOLE	TEV
02432404	VAN-OMEPRAZOLE	VAN

56:28.36 PROTON-PUMP INHIBITORS**PANTOPRAZOLE MAGNESIUM**

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

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Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST 40MG TABLET (DELAYED RELEASE)

02466147 PANTOPRAZOLE T SAN

ST 40MG TABLET (ENTERIC COATED)

02408570 MYLAN-PANTOPRAZOLE T MYL

02441853 PANTOPRAZOLE MAGNESIUM UNK

02267233 TECTA TAK

02440628 TEVA-PANTOPRAZOLE MAGNESIUM TEV

PANTOPRAZOLE SODIUM

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

The following PPI status change is primarily based on the Canadian Optimal Medication Prescribing and Utilization Service (COMPUS) report on optimal PPI therapy. The report concluded that:

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Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST 40MG TABLET (DELAYED RELEASE)

02292920 APO-PANTOPRAZOLE APX

02415208 AURO-PANTOPRAZOLE AUR

02445867 BIO-PANTOPRAZOLE BMI

02310007 DOM-PANTOPRAZOLE DPC

02357054 JAMP-PANTOPRAZOLE JMP

02416565 MAR-PANTOPRAZOLE MAR

02417448 MINT-PANTOPRAZOLE MIN

02467372 M-PANTOPRAZOLE MAN

02229453 PANTOLOC TAK

02318695 PANTOPRAZOLE PDL

02370808 PANTOPRAZOLE SAN

02431327 PANTOPRAZOLE RIV

56:28.36 PROTON-PUMP INHIBITORS**PANTOPRAZOLE SODIUM**

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

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Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST **40MG TABLET (DELAYED RELEASE)**

02437945	PANTOPRAZOLE	PMS
02439107	PANTOPRAZOLE	DPC
02428180	PANTOPRAZOLE-40	SIV
02307871	PMS-PANTOPRAZOLE	PMS
02425378	PRIVA-PANTOPRAZOLE	PHA
02305046	RAN-PANTOPRAZOLE	RBY
02316463	RIVA-PANTOPRAZOLE	RIV
02301083	SANDOZ PANTOPRAZOLE	SDZ
02285487	TEVA-PANTOPRAZOLE	TEV

RABEPRAZOLE SODIUM

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

The following PPI status change is primarily based on the Canadian Optimal Medication Prescribing and Utilization Service (COMPUS) report on optimal PPI therapy. The report concluded that:

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Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST **10MG TABLET (ENTERIC COATED)**

02345579	APO-RABEPRAZOLE	APX
02243796	PARIET	JSO
02310805	PMS-RABEPRAZOLE	PMS
02315181	PRO-RABEPRAZOLE	PDL
02385449	RABEPRAZOLE	SIV
02356511	RABEPRAZOLE EC	SAN
02298074	RAN-RABEPRAZOLE	RBY
02330083	RIVA-RABEPRAZOLE EC	RIV
02314177	SANDOZ RABEPRAZOLE	SDZ

56:28.36 PROTON-PUMP INHIBITORS**RABEPRAZOLE SODIUM**

Limited use benefit (prior approval not required).

Coverage will be limited to 400 tablets/capsules every 180 days.

The following PPI status change is primarily based on the Canadian Optimal Medication Prescribing and Utilization Service (COMPUS) report on optimal PPI therapy. The report concluded that:

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Patients with Zollinger Ellison Syndrome, Barrett's esophagus, erosive esophagitis and those who remain symptomatic on a single dose PPI will be eligible for additional doses above 400 tablets/capsules per 180 days through the prior approval process.

ST 10MG TABLET (ENTERIC COATED)

02296632 TEVA-RABEPRAZOLE

TEV

ST 20MG TABLET (ENTERIC COATED)

02345587 APO-RABEPRAZOLE

APX

02320460 DOM-RABEPRAZOLE EC

DPC

02243797 PARIET

JSO

02310813 PMS-RABEPRAZOLE

PMS

02315203 PRO-RABEPRAZOLE

PDL

02385457 RABEPRAZOLE

SIV

02356538 RABEPRAZOLE EC

SAN

02298082 RAN-RABEPRAZOLE

RBY

02330091 RIVA-RABEPRAZOLE

RIV

02314185 SANDOZ RABEPRAZOLE

SDZ

02296640 TEVA-RABEPRAZOLE

TEV

56:92.00 MISCELLANEOUS GI DRUGS**OBETICHOLIC ACID**

Limited use benefit (prior approval required).

Criteria for initial 12-month coverage:

The patient has a confirmed diagnosis of primary biliary cholangitis (PBC), defined as:

- Positive antimitochondrial antibodies (AMA); OR
- Liver biopsy results consistent with PBC.

AND

The patient is under the care of a gastroenterologist, hepatologist or internal medicine specialist with experience in the treatment of PBC.

AND

The patient has received ursodeoxycholic acid (UDCA) for a minimum of 12 months and has experienced an inadequate response to UDCA and can benefit from the addition of obeticholic acid. An inadequate response is defined as:

- Alkaline phosphatase (ALP) $\geq 1.67 \times$ upper limit of normal (ULN); AND/OR
- Bilirubin $> \text{ULN}$ and $< 2 \times \text{ULN}$; AND/OR
- Evidence of compensated cirrhosis by fibroscan or biopsy.

OR

The patient has experienced documented and unmanageable intolerance to UDCA.

Criteria for renewal every 12 months:

The patient continues to benefit from treatment with obeticholic acid as evidenced by:

- A reduction in the ALP level to less than $1.67 \times \text{ULN}$; OR
- A 15% reduction in the ALP level compared with values before beginning treatment with obeticholic acid.

5MG TABLET

02463121 OCALIVA

UNK

10MG TABLET

02463148 OCALIVA

UNK

56:92.00 MISCELLANEOUS GI DRUGS**PINAVERIUM BROMIDE**

Limited use benefit (prior approval required).

For the treatment and relief of symptoms associated with functional bowel disorders including Irritable Bowel Syndrome (IBS), spastic colon, spastic colitis and mucous colitis; OR
In postoperative paralytic ileus in order to accelerate the resumption of the intestinal transit following abdominal surgery.

50MG CAPSULE

00465240 DICETEL

SPH

50MG TABLET

01950592 DICETEL

BGP

100MG TABLET

02230684 DICETEL

BGP

68:00 HORMONES AND SYNTHETIC SUBSTITUTES**68:08.00 ANDROGENS****TESTOSTERONE (TOPICAL)**

Limited use benefit (prior approval required).

The NIH Program covers topical testosterone for the treatment of the following in adult males above 18 years old.

- Orchiectomy, undescended testes, Klinefelter's; OR
- Pituitary tumour or post-pituitary surgery with low testosterone; OR
- AIDS-wasting syndrome with low testosterone; OR
- Gender affirming hormone therapy.

Note: Older males with non-specific symptoms such as, but not limited to, fatigue, malaise, or depression who have a low random testosterone level do not meet coverage criteria.

1% GEL

02245345 ANDROGEL

BGP

02245346 ANDROGEL

BGP

02463792 TARO-TESTOSTERONE

TAR

02463806 TARO-TESTOSTERONE

TAR

02280248 TESTIM

PAL

12.5MG GEL

02249499 ANDROGEL

BGP

2.5MG PATCH

02239653 ANDRODERM

ALL

5MG PATCH

02245972 ANDRODERM

ALL

68:12.00 CONTRACEPTIVES**LEVONORGESTREL INTRAUTERINE INSERT**

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 1 device every 2 years.

52MG INSERT (EXTENDED-RELEASE)

02243005 MIRENA

BAY

ULIPRISTAL ACETATE

Limited use benefit (prior approval not required).

Coverage will be limited to 90 tablets, benefits only for women age 18 to 60 years.

ST 5MG TABLET

02408163 FIBRISTAL

ALL

68:16.12 ESTROGEN AGONISTS-ANTAGONISTS**RALOXIFENE HYDROCHLORIDE**

Limited use benefit (prior approval required).

For secondary prevention of osteoporosis in women who experience failure on bisphosphonates.

For secondary prevention of osteoporosis in women who have a personal history or a first degree relative with a history of breast cancer.

60MG TABLET

02358840	ACT RALOXIFENE	ACG
02279215	APO-RALOXIFENE	APX
02239028	EVISTA	LIL
02358921	PMS-RALOXIFENE	PMS

68:20.05 DIPEPTIDYL PEPTIDASE-4 (DPP-4) INHIBITORS**LINAGLIPTIN**

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **5MG TABLET**

02370921	TRAJENTA	BOE
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LINAGLIPTIN, METFORMIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **2.5MG & 1000MG TABLET**

02403277	JENTADUETO	BOE
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ST **2.5MG & 500MG TABLET**

02403250	JENTADUETO	BOE
----------	------------	-----

ST **2.5MG & 850MG TABLET**

02403269	JENTADUETO	BOE
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SAXAGLIPTIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who: did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **2.5MG TABLET**

02375842	ONGLYZA	AZC
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ST **5MG TABLET**

02333554	ONGLYZA	AZC
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SAXAGLIPTIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **2.5MG & 1000MG TABLET**

02389185	KOMBOGLYZE	AZC
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ST **2.5MG & 500MG TABLET**

02389169	KOMBOGLYZE	AZC
----------	------------	-----

ST **2.5MG & 850MG TABLET**

02389177	KOMBOGLYZE	AZC
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68:20.05 DIPEPTIDYL PEPTIDASE-4 (DPP-4) INHIBITORS**SITAGLIPTIN PHOSPHATE MONOHYDRATE**

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **25MG TABLET**

02388839 JANUVIA

FRS

ST **50MG TABLET**

02388847 JANUVIA

FRS

ST **100MG TABLET**

02303922 JANUVIA

FRS

SITAGLIPTIN PHOSPHATE MONOHYDRATE, METFORMIN HYDROCHLORIDE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **50MG & 1000MG TABLET**

02333872 JANUMET

FRS

ST **50MG & 500MG TABLET**

02333856 JANUMET

FRS

ST **50MG & 850MG TABLET**

02333864 JANUMET

FRS

ST **50MG & 1000MG TABLET (EXTENDED RELEASE)**

02416794 JANUMET XR

FRS

ST **50MG & 500MG TABLET (EXTENDED RELEASE)**

02416786 JANUMET XR

FRS

ST **100MG & 1000MG TABLET (EXTENDED RELEASE)**

02416808 JANUMET XR

FRS

68:20.18 SODIUM-GLUCOSE CONTRANSPORTER 2 (SGLT2) INHIBITORS**CANAGLIFLOZIN**

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **100MG TABLET**

02425483 INVOKANA

JSO

ST **300MG TABLET**

02425491 INVOKANA

JSO

DAPAGLIFLOZIN PROPANEDIOL MONOHYDRATE

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonylurea.

ST **5MG TABLET**

02435462 FORXIGA

AZC

ST **10MG TABLET**

02435470 FORXIGA

AZC

68:20.18 SODIUM-GLUCOSE CONTRANSPORTER 2 (SGLT2) INHIBITORS**EMPAGLIFLOZIN**

Open benefit with therapeutic notes (prior approval is not required).

For the treatment of type 2 diabetes mellitus:

- in patients who did not achieve glycemic control with an adequate trial of metformin AND a sulfonyleurea

OR

- to reduce the incidence of cardiovascular death in patients with established cardiovascular disease who did not achieve adequate glycemic control despite an appropriate trial of metformin

Established cardiovascular disease is defined as one of the following:

- history of myocardial infarction
- 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 either a positive non-invasive stress test or discharged from hospital with a documented diagnosis of unstable angina
- unstable angina with confirmed evidence of coronary multi-vessel or single-vessel disease
- history of ischemic or hemorrhagic stroke
- occlusive peripheral artery disease.

ST **10MG TABLET**

02443937 JARDIANCE

BOE

ST **25MG TABLET**

02443945 JARDIANCE

BOE

METFORMIN HYDROCHLORIDE, DAPAGLIFLOZIN

Limited use benefit (prior approval required).

For the treatment of patients with type 2 diabetes mellitus who did not achieve glycemic control or who demonstrated intolerance to an adequate trial of metformin AND a sulfonyleurea.

ST **850MG & 5MG TABLET**

02449935 XIGDUO

AZC

ST **1000MG & 5MG TABLET**

02449943 XIGDUO

AZC

METFORMIN HYDROCHLORIDE, EMPAGLIFLOZIN

Open benefit with therapeutic notes (prior approval is not required).

For the treatment of patients with type 2 diabetes mellitus in patients who are eligible to receive metformin and empagliflozin, to replace the individual components.

500MG & 12.5MG TABLET

02456605 SYNJARDY

BOE

500MG & 5MG TABLET

02456575 SYNJARDY

BOE

850MG & 12.5MG TABLET

02456613 SYNJARDY

BOE

850MG & 5MG TABLET

02456583 SYNJARDY

BOE

1000MG & 12.5MG TABLET

02456621 SYNJARDY

BOE

1000MG & 5MG TABLET

02456591 SYNJARDY

BOE

68:32.00 PROGESTINS**DIENOGEST**

Limited use benefit (prior approval required).

For the management of pelvic pain associated with endometriosis.

ST **2MG TABLET**

02374900 VISANNE

BAY

68:32.00 PROGESTINS**PROGESTERONE**

Limited use benefit (prior approval required).

For the treatment of women:

- With postmenopausal symptoms who are intolerant to medroxyprogesterone acetate (MPA); OR
- Who are at risk of preterm birth; OR
- Who are using the medication to prevent miscarriage.

In adults:

- For use as Gender Affirming Hormone Therapy.

100MG CAPSULE

02476576 PMS-PROGESTERONE

PMS

02166704 PROMETRIUM

FRS

02463113 REDDY-PROGESTERONE

REC

02439913 TEVA-PROGESTERONE

TEV

84:00 SKIN AND MUCOUS MEMBRANE AGENTS (SMMA)**84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS****BRODALUMAB**

Limited use benefit (prior approval required).

For PSORIASIS, coverage is provided for an initial period of 12 weeks at a dose of 210 mg at week 0, 1, and 2, followed by 210 mg every 2 weeks.

- Prescribed by a dermatologist

For the treatment of patients with moderate to severe PSORIASIS who meet all of the following criteria:

- Body surface area (BSA) involvement greater than 10% and/or significant involvement of the face, hands, feet or genital region; AND
- Intolerance or lack of response to phototherapy; OR
- Inability to access phototherapy; AND
- Intolerance or lack of response to methotrexate (MTX) weekly oral or parenteral at 20 mg or greater (15 mg or greater if patient is > 65 years of age) for more than 8 weeks; AND
- Intolerance or lack of response to cyclosporine; OR
- A contraindication to methotrexate or cyclosporine.

Coverage beyond 12 to 16 weeks will be based on a significant reduction in the Body Surface Area (BSA) involved and improvements in the Psoriasis Area Severity Index (PASI) score and the Dermatology Life Quality Index (DLQI).

- A 75% reduction in Psoriasis Area Severity Index (PASI) score; OR
- A ≥ 50% reduction in the Psoriasis Area Severity Index (PASI) score with a ≥ 5-point improvement in the Dermatology Life Quality Index (DLQI); OR
- A significant reduction in Body Surface Area (BSA) involved, with consideration of important areas such as the face, hands, feet or genital regions.

210MG SOLUTION

02473623 SILIQ

VAE

IMIQUIMOD

Limited use benefit (prior approval required).

For the treatment of condylomata acuminata (genital warts) in patients who have failed:

- self-applied podophyllotoxin (podofilox 0.5% solution); OR
- provider-applied podophyllum resin (10%-25%).

5% CREAM

02239505 ALDARA P

VAE

02407825 APO-IMIQUIMOD

APX

02482983 TARO-IMIQUIMOD PUMP

TAR

84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS**IXEKIZUMAB**

Limited use benefit (prior approval required).

1. For the treatment of moderate to severe PSORIATIC ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 160 mg at Week 0, followed by 80 mg every 4 weeks. For psoriatic arthritis patients with coexistent moderate-to-severe psoriasis, coverage is provided for psoriasis dosing: 160 mg at week 0, followed by 80 mg at weeks 2, 4, 6, 8, 10, and 12, then 80 mg every 4 weeks. For psoriatic arthritis patients with coexistent mild plaque psoriasis, coverage is provided for psoriatic arthritis dosing: 160 mg at Week 0, followed by 80 mg every 4 weeks.

- Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;
- PLUS a minimum of any two of the following:
 - methotrexate weekly (weekly oral or parenteral) at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
 - leflunomide: 20mg daily for 10 weeks; OR
 - sulfasalazine at least 2g daily for 3 months; OR
 - cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) \geq 4; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

Coverage beyond one year will be based on improvement in at least 2 of 4 Psoriatic Arthritis Response Criteria (PsARC).

- Improvement in at least two of the four PsARC criteria, one of which has to be joint tenderness or swelling score, with no worsening in any of the four criteria. A response in joint count is determined by a reduction of \geq 30%. A response in the Physician or Patient Global Assessment scale is determined by a reduction of 1 point.

2. For PSORIASIS ONLY, coverage is provided for an initial period of 12 weeks at a dose of 160mg at week 0, followed by 80mg at weeks 2, 4, 6, 8, 10, and 12, then 80mg every 4 weeks.

- Prescribed by a dermatologist

For the treatment of patients with moderate to severe PSORIASIS who meet all of the following criteria:

- Body surface area (BSA) involvement greater than 10% and/or significant involvement of the face, hands, feet or genital region; AND
- Intolerance or lack of response to phototherapy; OR
- Inability to access phototherapy; AND
- Intolerance or lack of response to methotrexate (MTX) weekly oral or parenteral at 20 mg or greater (15 mg or greater if patient is > 65 years of age) for more than 8 weeks; AND
- Intolerance or lack of response to cyclosporine; OR
- A contraindication to methotrexate or cyclosporine.

Coverage beyond 12 weeks will be based on a significant reduction in the Body Surface Area (BSA) involved and improvements in the Psoriasis Area Severity Index (PASI) score and the Dermatology Life Quality Index (DLQI):

- A 75% reduction in Psoriasis Area Severity Index (PASI) score; OR
- A \geq 50% reduction in the Psoriasis Area Severity Index (PASI) score with a \geq 5-point improvement in the Dermatology Life Quality Index (DLQI); OR
- A significant reduction in Body Surface Area (BSA) involved, with consideration of important areas such as the face, hands, feet or genital regions.

80MG SOLUTION

02455102 TALTZ

LIL

02455110 TALTZ

LIL

PIMECROLIMUS

Limited use benefit (prior approval required).

For patients who have failed topical corticosteroid therapy or have experienced side effects from such treatment.

Note: Contraindicated in children less than 2 years of age.

1% CREAM

02247238 ELIDEL

VAE

84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS**SECUKINUMAB**

Limited use benefit (prior approval required).

1. For the treatment of moderate to severe PSORIASIS

Coverage is provided for an initial period of 12 weeks at a dose of 300mg at Weeks 0, 1, 2 and 3, followed by 300mg per month starting at Week 4.
 • Prescribed by a dermatologist

For the treatment of patients with moderate to severe PSORIASIS who meet all of the following criteria:

- Body surface area (BSA) involvement greater than 10% and/or significant involvement of the face, hands, feet or genital region; AND
- Intolerance or lack of response to phototherapy; OR
- Inability to access phototherapy; AND
- Intolerance or lack of response to methotrexate (MTX) weekly oral or parenteral at 20 mg or greater (15 mg or greater if patient is > 65 years of age) for more than 8 weeks; AND
- Intolerance or lack of response to cyclosporine; OR
- A contraindication to methotrexate or cyclosporine.

Coverage beyond 12 weeks will be based on a significant reduction in the Body Surface Area (BSA) involved and improvements in the Psoriasis Area Severity Index (PASI) score and the Dermatology Life Quality Index (DLQI):

- A 75% reduction in Psoriasis Area Severity Index (PASI) score; OR
- A $\geq 50\%$ reduction in the Psoriasis Area Severity Index (PASI) score with a ≥ 5 -point improvement in the Dermatology Life Quality Index (DLQI); OR
- A significant reduction in Body Surface Area (BSA) involved, with consideration of important areas such as the face, hands, feet or genital regions.

2. For the treatment of moderate to severe PSORIATIC ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 150 mg at weeks 0, 1, 2 and 3, followed by 150 mg per month starting at week 4. If patient is an anti-TNF inadequate responder and continues to have active psoriatic arthritis or has co-existent severe plaque psoriasis, 300 mg per month will be considered.
 • Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;

PLUS a minimum of any two of the following:

- methotrexate weekly (weekly oral or parenteral) at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
- leflunomide: 20mg daily for 10 weeks; OR
- sulfasalazine at least 2g daily for 3 months; OR
- cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) ≥ 4 ; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

Coverage beyond one year will be based on improvement according to the Psoriatic Arthritis Response Criteria (PsARC).

- Improvement in at least two of the four PsARC criteria, one of which has to be joint tenderness or swelling score, with no worsening in any of the four criteria. A response in joint count is determined by a reduction of $\geq 30\%$. A response in the Physician or Patient Global Assessment scale is determined by a reduction of 1 point.

3. For the treatment of ANKYLOSING SPONDYLITIS

Coverage is provided for an initial period of one year at a dose of 150 mg at weeks 0, 1, 2 and 3, followed by 150 mg per month starting at week 4.

- Prescribed by a rheumatologist
- BASDAI > 4; AND
- Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks; AND for peripheral joint involvement, patient is refractory:
 - Methotrexate (MTX) (weekly oral or parenteral) weekly at 20 mg or greater (15 mg or greater if patient is >65 years of age) for more than 8 weeks; AND
 - Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

Coverage beyond one year will be based on improvement in the BASDAI score.

- Improvement of at least 50% or 2 units in the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) score.

150MG/ML INJECTION

99101215 COSENTYX (STYLO)

NVC

09857548 COSENTYX PEN (ON)

NVC

150MG SOLUTION

02438070 COSENTYX

NVR

84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS**TACROLIMUS (PROTOPIC)**

Limited use benefit (prior approval required).

For patients who have failed topical corticosteroid therapy or have experienced side effects from such treatment.

Note: Contraindicated in children less than 2 years of age.

0.03% OINTMENT

02244149 PROTOPIC

LEO

0.1% OINTMENT

02244148 PROTOPIC

LEO

86:00 SMOOTH MUSCLE RELAXANTS**86:12.04 ANTIMUSCARINICS****DARIFENACIN HYDROBROMIDE**

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:

- with symptoms of urinary frequency, urgency or urge incontinence; AND
- who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

7.5MG TABLET (EXTENDED RELEASE)

02273217 ENABLEX

UNK

15MG TABLET (EXTENDED RELEASE)

02273225 ENABLEX

UNK

FESOTERODINE FUMARATE

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:

- with symptoms of urinary frequency, urgency or urge incontinence; AND
- who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

ST 4MG TABLET (EXTENDED RELEASE)

02380021 TOVIAZ

PFI

ST 8MG TABLET (EXTENDED RELEASE)

02380048 TOVIAZ

PFI

TROSPIUM CHLORIDE

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:

- with symptoms of urinary frequency, urgency or urge incontinence; AND
- who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

ST 20MG TABLET

02275066 TROSEC

SPC

86:12.08 BETA-ADRENERGIC AGONISTS**MIRABEGRON**

Limited use benefit (prior approval required).

For the symptomatic relief of overactive bladder in patients:

- with symptoms of urinary frequency, urgency or urge incontinence; AND
- who have failed on or are intolerant to therapy with immediate-release oxybutynin OR solifenacin OR tolterodine ER.

ST 25MG TABLET (EXTENDED RELEASE)

02402874 MYRBETRIQ

AST

ST 50MG TABLET (EXTENDED RELEASE)

02402882 MYRBETRIQ

AST

88:00 VITAMINS**88:20.00 VITAMIN E****VITAMIN E**

Limited use benefit (prior approval required).

For use in malabsorption

ST 100IU CAPSULE (SOFTGEL)		
00122823 VITAMIN E		JAM
ST 200IU CAPSULE (SOFTGEL)		
00122831 VITAMIN E		JAM
ST 400IU CAPSULE (SOFTGEL)		
00122858 VITAMIN E		JAM
ST 800IU CAPSULE (SOFTGEL)		
00330191 VITAMIN E		JAM
ST 20U/ML LIQUID		
09991656 AQUA-E/ML		UNK
ST 75U/ML LIQUID		
09991652 AQUA-E		UNK
ST 50IU ORAL LIQUID		
00480215 AQUASOL E		NVC
ST 50IU/ML ORAL LIQUID		
02162075 AQUASOL E VITAMIN E		CLC

88:28.00 MULTIVITAMIN PREPARATIONS**MULTIVITAMINS (CHILDREN AND YOUTH)**

Limited use benefit (prior approval is not required).

Multivitamins are benefits for children up to 19 years of age.

ST DROP		
00762946 ENFAMIL POLYVISOL		MJO
ST 450MG & 10MG & 30MG LIQUID		
80008471 JAMP VITAMIN A, D AND C		JMP
ST 2,500IU & 666.67IU & 50MG/ML LIQUID		
00762903 ENFAMIL TRIVISOL		MJO
02229790 PEDIAVIT		EUR
0MG TABLET		
02246362 CENTRUM		PFI
80021452 CENTRUM		PFI
80024482 CENTRUM FOR WOMEN		PFI
2MG TABLET		
80045908 ONE A DAY WOMEN		BAY
10MG TABLET		
80039441 STRESSTABS FOR WOMEN		PFI
ST TABLET (CHEWABLE)		
80011134 CENTRUM JUNIOR COMPLETE		PFI
80020794 CENTRUM JUNIOR COMPLETE		PFI
02247995 FLINTSTONES MULTIPLE VITAMINS PLUS IRON		BAY
02247975 FLINTSTONES MULTIPLE VITAMINS WITH EXTRA C		BAY

88:28.00 MULTIVITAMIN PREPARATIONS**MULTIVITAMINS (PRENATAL)**

Limited use benefit (prior approval is not required.).

Prenatal and postnatal vitamins are benefits only for women of childbearing age (12 to 50 years).

ST **CAPSULE**

80042704 CENTRUM DHA PFI

ST **TABLET**

80045822 CENTRUM PRENATAL PFI

80080882 MATERNA NES

80082297 MATERNA NES

80001842 NESTL MATERNA NES

02241235 PRENATAL AND POSTPARTUM VITAMINS AND MINERALS VTH

80005770 PRENATAL AND POSTPARTUM VITAMINS AND MINERALS PMT

02229535 WAMPOLE COMPLETE MULT-PRE AND POST NATAL WITH FOLIC ACID WAM

2MG TABLET

80004919 NATURES BOUNTY PRENATAL VITAMINS VTH

92:00 UNCLASSIFIED THERAPEUTIC AGENTS**92:00.00 UNCLASSIFIED THERAPEUTIC AGENTS****EXTEMPORANEOUS MIXTURE (GENDER AFFIRMING)**

Limited use benefit (prior approval required).

For gender affirming hormone therapy.

INJECTION

00915312 GENDER AFFIRMING HORMONES UNK

LIQUID

00915311 GENDER AFFIRMING TOPICAL HORMONES UNK

EXTEMPORANEOUS MIXTURE (LU)

Limited use benefit (prior approval required).

INJECTION

99506021 MISCELLANEOUS COMPOUNDED INJECTION/INFUSION UNK

MISCELLANEOUS

99504001 MISC LIMITED USE EXTERNAL COMPOUND MIXTURE UNK

OPHTHALMIC AND OTIC SOLUTION

99507000 MISCELLANEOUS COMPOUNDED EYE/EAR DROP UNK

ORAL LIQUID

99503033 MISC LIMITED USE COMPOUND INTERNAL UNK

99503032 OPIOID COMPOUNDED UNK

POWDER

99504000 MISCELLANEOUS COMPOUNDED EXTERNAL POWDER UNK

SUPPOSITORY

99508000 MISCELLANEOUS COMPOUNDED SUPPOSITORY UNK

92:00.00 UNCLASSIFIED THERAPEUTIC AGENTS**USTEKINUMAB**

Limited use benefit (prior approval required).

Coverage is provided for an initial period of 16 weeks. For patients \leq 100 kg, the initial dose is 45 mg at week 0, followed by 45 mg at weeks 4 and 16. Alternatively, ustekinumab 90 mg may be used in patients weighing more than 100 kg. Response must be assessed prior to a fourth dose and further doses will be provided only for responders.

- Prescribed by a dermatologist

For the treatment of patients with moderate to severe PSORIASIS who meet all of the following criteria:

- Body surface area (BSA) involvement greater than 10% and/or significant involvement of the face, hands, feet or genital region; AND
- Intolerance or lack of response to phototherapy; OR
- Inability to access phototherapy; AND
- Intolerance or lack of response to methotrexate (MTX) weekly oral or parenteral at 20 mg or greater (15 mg or greater if patient is $>$ 65 years of age) for more than 8 weeks; AND
- Intolerance or lack of response to cyclosporine; OR
- A contraindication to methotrexate or cyclosporine.

Coverage beyond 16 weeks will be based on a significant reduction in the Body Surface Area (BSA) involved and improvements in the Psoriasis Area Severity Index (PASI) score and the Dermatology Life Quality Index (DLQI):

- A 75% reduction in Psoriasis Area Severity Index (PASI) score; OR
- A \geq 50% reduction in the Psoriasis Area Severity Index (PASI) score with a \geq 5-point improvement in the Dermatology Life Quality Index (DLQI); OR
- A significant reduction in Body Surface Area (BSA) involved, with consideration of important areas such as the face, hands, feet or genital regions.

45MG/0.5ML SOLUTION

02320673 STELARA

JSO

90MG/ML SOLUTION

02320681 STELARA

JSO

92:16.00 ANTIGOUT AGENTS**FEBUXOSTAT**

Limited use benefit (prior approval required).

For patients with symptomatic gout who have documented hypersensitivity to allopurinol.

ST **80MG TABLET**

02357380 ULORIC

TAK

92:20.00 IMMUNOMODULATORY AGENTS**GLATIRAMER ACETATE**

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

20MG SOLUTION

02245619 COPAXONE

TEV

02460661 GLATECT

PMS

INTERFERON BETA-1A

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

30MCG INJECTION

09857395 AVONEX PEN

UNK

99100763 AVONEX PEN

UNK

60MCG POWDER FOR SOLUTION

02267594 AVONEX

UNK

92:20.00 IMMUNOMODULATORY AGENTS**INTERFERON BETA-1A**

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

22MCG SOLUTION

02237319 REBIF

SRO

30MCG SOLUTION

02269201 AVONEX

UNK

44MCG SOLUTION

02237318 REBIF

SRO

02237320 REBIF

SRO

66MCG SOLUTION

02318253 REBIF

SRO

132MCG SOLUTION

02318261 REBIF

SRO

02318288 REBIF

SRO

INTERFERON BETA-1B

Limited use benefit (prior approval required).

- As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

0.3MG INJECTION

99100555 BETASERON INITIATION KIT

BAY

0.3MG POWDER FOR SOLUTION

02169649 BETASERON

BAY

02337819 EXTAVIA

NVR

92:20.00 IMMUNOMODULATORY AGENTS**OCRELIZUMAB**

Limited use benefit (prior approval required).

1. For the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence who meet all of the following criteria:

- Prescribed by a neurologist experienced in the management of RRMS; AND
- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids. Expanded Disability Status Scale score (EDSS) of 5.5 or less.
- Patient is 18 years of age or older.

a. A clinical relapse is defined as the appearance of new symptoms or worsening of old symptoms, lasting at least 24 hours in the absence of fever, preceded by stability for at least one month.

b. MRI activity is defined as any new multiple sclerosis lesion/s, expanding lesion/s, and/or enhancing lesion/s.

OR

2. For the treatment of primary progressive multiple sclerosis (PPMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence who meet all of the following criteria:

Initial Coverage (one year)

- Prescribed by a neurologist experienced in the management of PPMS; AND
- Expanded Disability Status Scale (EDSS) between 3.0 and 6.5; AND
- Score of at least 2.0 on the Functional Systems scale (FSS) for the pyramidal system due to lower extremity findings; AND
- Disease duration of less than 15 years for those with an EDSS greater than 5.0 or less than 10 years for those with an EDSS of 5.0 or less; AND
- Patient is 18 years of age or older.

Renewal Coverage for PPMS (one year):

- EDSS of less than 7.0.

30MG SOLUTION

02467224 OCREVUS

HLR

TERIFLUNOMIDE

Limited use benefit (prior approval required).

• As a first-line therapy for the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence, when prescribed by a neurologist experienced in the management of RRMS.

AND for patients who meet ALL of the following criteria:

- Patient has had a clinical relapse and/or new MRI activity in the last two years; AND
- Patient is fully ambulatory for 100 meters without aids; AND
- Patient is 18 years of age or older.

14MG TABLET

02416328 AUBAGIO

GEE

92:24.00 BONE RESORPTION INHIBITORS**DENOSUMAB (PROLIA)**

Limited use benefit (prior approval required).

For the treatment of osteoporosis in patients who have a significant fracture risk defined as either:

- moderate 10-year fracture risk (10% to 20%) with a prior fragility fracture; OR
- high 10-year fracture risk ($\geq 20\%$);

AND

- Have a contraindication to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment); OR
- Have failed or have an intolerance to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment).

60MG/ML SOLUTION

02343541 PROLIA

AMG

DENOSUMAB (XGEVA)

Limited use benefit (prior approval required).

For the prevention of skeletal-related events (SREs) in patients with castrate-resistant prostate cancer (CRPC) with:

- one or more documented bone metastases; AND
- good performance status (ECOG performance status score of 0, 1, or 2).

120MG/1.7ML SOLUTION

02368153 XGEVA

AMG

92:24.00 BONE RESORPTION INHIBITORS**ZOLEDRONIC ACID MONOHYDRATE**

Limited use benefit (prior approval required).

Maximum dose covered is 5mg per 12-month period

For the treatment of Paget's disease;

OR

For the treatment of osteoporosis in patients who have a significant fracture risk defined as either:

- moderate 10-year fracture risk (10% to 20%) with a prior fragility fracture; OR

- high 10-year fracture risk ($\geq 20\%$)

AND

- Have a contraindication to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment);OR

- Have failed or have an intolerance to oral bisphosphonates (e.g. hypersensitivity, esophageal abnormality, renal impairment).

5MG/100ML SOLUTION

02269198 ACLASTA

NVR

02415100 TARO-ZOLEDRONIC ACID

TAR

02408082 ZOLEDRONIC ACID

TEV

02422433 ZOLEDRONIC ACID

REC

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**ABATACEPT**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 500mg IV for patients weighting <60kg; 750mg IV for patients weighting 60kg to 100kg; and 1000mg IV for patients weighing >100kg. Initial IV doses are given at 0, 2, and 4 weeks, then every 4 weeks. Alternatively, a single weight-based IV loading dose is covered (if required), followed by 125mg SC weekly.

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX; AND

- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment;

OR, if the patient has a contraindication, failure, or intolerance to MTX:

- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

AND (FOR IV FORMULATION ONLY):

- etanercept (sc) OR adalimumab (sc) OR golimumab (sc) OR certolizumab (sc) OR abatacept (sc) OR tocilizumab OR tofacitinib (po) OR Inflectra (iv) OR Renflexis (iv): for a minimum trial of 12 weeks.

Coverage beyond one year will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- >20% reduction in number of tender and swollen joints; PLUS
- >20% improvement in Physician Global Assessment scale; PLUS either
- >20% improvement in Patient Global Assessment scale; OR
- >20% reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of severely active polyarticular JUVENILE IDIOPATHIC ARTHRITIS

Coverage is provided for an initial period of 16 weeks at a dose of 10mg/kg for children weighing < 75kg; Pediatric patients weighing 75kg or more should be dosed according to the adult regimen, not to exceed a maximum dose of 1000mg. Doses are given at 0, 2, and 4 weeks, then every 4 weeks.

- Prescribed by a rheumatologist

In patients six to seventeen years of age who meet the following criteria:

- ≥ 5 swollen joints; AND
- ≥ 3 joints with limited range of motion and/or pain/tenderness; AND
- Condition is refractory to an adequate trial of a therapeutic dose of methotrexate.

Coverage beyond 16 weeks is based on a >30% improvement in 3 of 6 baseline clinical parameters

Patient has experienced 3 of 6 of the following variables:

- >30% reduction in the number of active joints
- >30% reduction in the number of joints with loss of range of motion
- >30% improvement in the Physician Global Assessment scale
- >30% improvement in the Patient or Parent Global Assessment scale
- >30% improvement in the Child Health Assessment Questionnaire (CHAQ)
- >30% reduction in ESR; AND
- No more than one of these variables has worsened by greater than 30%

250MG POWDER FOR SOLUTION

02282097 ORENCIA

BMS

125MG SOLUTION

02402475 ORENCIA

BMS

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**ADALIMUMAB**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 40 mg every two weeks.

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX; AND
- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond one year will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- $>20\%$ reduction in number of tender and swollen joints; PLUS
- $>20\%$ improvement in Physician Global Assessment scale; PLUS either
- $>20\%$ improvement in Patient Global Assessment scale; OR
- $>20\%$ reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of moderate to severe PSORIATIC ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 40 mg every two weeks.

- Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;
- PLUS a minimum of any two of the following:
- methotrexate weekly (weekly oral or parenteral) at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
- leflunomide: 20mg daily for 10 weeks; OR
- sulfasalazine at least 2g daily for 3 months; OR
- cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) ≥ 4 ; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

Coverage beyond one year will be based on improvement according to the Psoriatic Arthritis Response Criteria (PsARC).

- Improvement in at least two of the four PsARC criteria, one of which has to be joint tenderness or swelling score, with no worsening in any of the four criteria. A response in joint count is determined by a reduction of $\geq 30\%$. A response in the Physician or Patient Global Assessment scale is determined by a reduction of 1 point.

3. For the treatment of ANKYLOSING SPONDYLITIS

Coverage is provided for an initial period of one year at a dose of 40 mg every two weeks.

- Prescribed by a rheumatologist
- BASDAI > 4 ; AND
- Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks; AND for peripheral joint involvement, patient is refractory:
- Methotrexate (MTX) (weekly oral or parenteral) weekly at 20 mg or greater (15 mg or greater if patient is >65 years of age) for more than 8 weeks; AND
- Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

Coverage beyond one year will be based on improvement in the BASDAI score.

- Improvement of at least 50% or 2 units in the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) score.

4. For the treatment of patients with moderate to severe PSORIASIS who meet all of the following criteria:

Coverage is provided for an initial period of 16 weeks at a dose of 80 mg as an initial dose, followed by 40 mg every 2 weeks, starting one week after the initial dose.

- Prescribed by a dermatologist
- Body surface area (BSA) involvement greater than 10% and/or significant involvement of the face, hands, feet or genital region; AND
- Intolerance or lack of response to phototherapy; OR
- Inability to access phototherapy; AND
- Intolerance or lack of response to methotrexate (MTX) (weekly oral or parenteral) at 20 mg or greater (15 mg or greater if patient is > 65 years of age) for more than 8 weeks; AND

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- Intolerance or lack of response to cyclosporine; OR
- A contraindication to methotrexate or cyclosporine.

Coverage beyond 16 weeks will be based on a significant reduction in the Body Surface Area (BSA) involved and improvements in the Psoriasis Area Severity Index (PASI) score and the Dermatology Life Quality Index (DLQI):

- A 75% reduction in Psoriasis Area Severity Index (PASI) score; OR
- A $\geq 50\%$ reduction in the Psoriasis Area Severity Index (PASI) score with a ≥ 5 -point improvement in the Dermatology Life Quality Index (DLQI); OR
- A significant reduction in Body Surface Area (BSA) involved, with consideration of important areas such as the face, hands, feet or genital regions.

5. For the treatment of moderately to severely active CROHN'S DISEASE

Coverage is provided for an initial period of 12 weeks at an induction dose of 160 mg, followed by 80 mg two weeks later. Maintenance therapy is provided at a dose not exceeding 40 mg every two weeks.

- Prescribed by a gastroenterology specialist

Patient meets the following criteria:

- Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication; PLUS
- Azathioprine 2 mg/kg/day for a minimum of 12 weeks; OR
- 6-mercaptopurine 1 mg/day for a minimum of 12 weeks; OR
- MTX (oral or parenteral) 15 mg per week for a minimum of 12 weeks..

Coverage beyond the initial twelve-week period will be based on improvement in the CDAI or HBI scores.

- At least a 100-point reduction in the Crohn's Disease Activity Index (CDAI) OR at least a 3-point reduction in the Harvey Bradshaw Index (HBI).

6. For the treatment of severely active polyarticular JUVENILE IDIOPATHIC ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 24 mg/m² body surface area up to a maximum single dose of 40 mg every other week.

- Prescribed by a rheumatologist

In patients two years of age and older who meet the following criteria:

- ≥ 5 swollen joints; AND
- ≥ 3 joints with limited range of motion and/or pain/tenderness; AND
- Condition is refractory to an adequate trial of a therapeutic dose of methotrexate.

Coverage beyond the initial one-year period will be based on a 30% improvement in 3 of 6 clinical parameters

Patient has experienced 3 of 6 of the following variables:

- $>30\%$ reduction in the number of active joints
- $>30\%$ reduction in the number of joints with loss of range of motion
- $>30\%$ improvement in the Physician Global Assessment scale
- $>30\%$ improvement in the Patient or Parent Global Assessment scale
- $>30\%$ improvement in the Child Health Assessment Questionnaire (CHAQ)
- $>30\%$ reduction in ESR; AND
- No more than one of these variables has worsened by greater than 30%

7. For the treatment of adult patients with moderately to severely active ULCERATIVE COLITIS who meet the following:

Coverage is provided for an initial period of 12 weeks at a dose of 160 mg at week 0, followed by 80 mg two weeks later and then 40 mg every two weeks thereafter.

- Prescribed by expert in gastroenterology
- Partial Mayo score > 4
- Inadequate response to conventional therapies:
 - 5-ASA 4grams/day for 6 weeks; PLUS
 - Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication.

Coverage beyond the initial 12 week period will be based on improvement in the partial Mayo score of ≥ 2 points.

8. For the treatment of adult patients with active moderate to severe HIDRADENITIS SUPPURATIVA

Coverage is provided for an initial period of 12 weeks at a dose of 160 mg at week 0, followed by 80 mg two weeks later, and then 40 mg every week beginning 4 weeks after the initial dose.

- Prescribed by a dermatologist

For the treatment of adult patients with active moderate to severe HIDRADENITIS SUPPURATIVA who meet all of the following criteria:

- Total inflammatory lesion (abscess and nodule) count of 3 or greater; AND
- Lesions in at least two distinct anatomic areas, one of which must be Hurley Stage II or III*; AND
- Inadequate response to a 90-day trial of oral antibiotics.

* Hurley Stage II and III defined as:

Stage II :One or more widely separated recurrent abscesses with tract formation and scars

Stage III: Multiple interconnected tracts and abscesses throughout an entire area

Coverage beyond the initial 12-week period will be based on decreases in inflammatory nodule and abscess counts:

- At least a 50% reduction in abscesses and inflammatory nodule count from baseline; AND
- No increase in abscess count; AND
- No increase in draining fistula count.

40MG/VIAL SOLUTION

02258595 HUMIRA

ABV

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**CERTOLIZUMAB PEGOL**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 400mg at weeks 0, 2, and 4, followed by 200mg every other week or 400mg every 4 weeks.
 • Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients \geq 18 years who have failed:

- MTX (oral or parenteral) at a dose \geq 20 mg weekly (\geq 15 mg weekly if patient is \geq 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX; AND
- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond the initial three doses will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- >20% reduction in number of tender and swollen joints; PLUS
- >20% improvement in Physician Global Assessment scale; PLUS either
- >20% improvement in Patient Global Assessment scale; OR
- >20% reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of moderate to severe PSORIATIC ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 400mg at weeks 0, 2, and 4, followed by 200mg every other week or 400mg every 4 weeks.
 • Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;
- PLUS a minimum of any two of the following:
- methotrexate weekly parenteral (SC or IM) at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
- leflunomide: 20mg daily for 10 weeks; OR
- sulfasalazine at least 2g daily for 3 months; OR
- cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) \geq 4; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

Coverage beyond one year will be based on improvement according to the Psoriatic Arthritis Response Criteria (PsARC).

- Improvement in at least two of the four PsARC criteria, one of which has to be joint tenderness or swelling score, with no worsening in any of the four criteria. A response in joint count is determined by a reduction of \geq 30%. A response in the Physician or Patient Global Assessment scale is determined by a reduction of 1 point.

3. For the treatment of ANKYLOSING SPONDYLITIS

Coverage is provided for an initial period of one year at a dose of 400mg at weeks 0, 2, and 4, followed by 200mg every other week or 400mg every 4 weeks.

- Prescribed by a rheumatologist
- BASDAI > 4; AND
- Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks;
- AND for peripheral joint involvement, patient is refractory:
- Methotrexate (MTX) weekly at 20 mg or greater (15 mg or greater if patient is >65 years of age) for more than 8 weeks; AND
- Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

Coverage beyond the initial three doses will be based on improvement in the BASDAI score.

- Improvement of at least 50% or 2 units in the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) score.

200MG SOLUTION

02465574 CIMZIA

UCB

200MG/ML SOLUTION

02331675 CIMZIA

UCB

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**ETANERCEPT**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 50 mg weekly.

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX; AND
- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond the initial three doses will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- $>20\%$ reduction in number of tender and swollen joints; PLUS
- $>20\%$ improvement in Physician Global Assessment scale; PLUS either
- $>20\%$ improvement in Patient Global Assessment scale; OR
- $>20\%$ reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of moderate to severe PSORIATIC ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 50 mg weekly.

- Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;
- PLUS a minimum of any two of the following:
- methotrexate (oral or parenteral) at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
- leflunomide: 20mg daily for 10 weeks; OR
- sulfasalazine at least 2g daily for 3 months; OR
- cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) ≥ 4 ; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

Coverage beyond one year will be based on improvement according to the Psoriatic Arthritis Response Criteria (PsARC).

- Improvement in at least two of the four PsARC criteria, one of which has to be joint tenderness or swelling score, with no worsening in any of the four criteria. A response in joint count is determined by a reduction of $\geq 30\%$. A response in the Physician or Patient Global Assessment scale is determined by a reduction of 1 point.

3. For the treatment of ANKYLOSING SPONDYLITIS

Coverage is provided for an initial period of one year at a dose of 50 mg weekly.

- Prescribed by a rheumatologist
- BASDAI > 4 ; AND
- Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks; AND for peripheral joint involvement, patient is refractory:
- Methotrexate (MTX) weekly (oral or parenteral) at 20 mg or greater (15 mg or greater if patient is >65 years of age) for more than 8 weeks; AND
- Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

Coverage beyond one year will be based on improvement in the BASDAI score.

- Improvement of at least 50% or 2 units in the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) score.

4. For the treatment of severely active polyarticular JUVENILE IDIOPATHIC ARTHRITIS

Coverage is provided for children age 4 to 17, for an initial period of one year at a dose of 0.8 mg/kg/week body surface area up to a maximum single dose of 50 mg/week.

- Prescribed by a rheumatologist

In patients four to seventeen years of age and older who meet the following criteria:

- ≥ 5 swollen joints; AND
- ≥ 3 joints with limited range of motion and/or pain/tenderness; AND

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- Condition is refractory to an adequate trial of a therapeutic dose of methotrexate.

Coverage beyond the initial one-year period will be based on a 30% improvement in 3 of 6 clinical parameters.

Patient has experienced 3 of 6 of the following variables:

- >30% reduction in the number of active joints
- >30% reduction in the number of joints with loss of range of motion
- >30% improvement in the Physician Global Assessment scale
- >30% improvement in the Patient or Parent Global Assessment scale
- >30% improvement in the Child Health Assessment Questionnaire (CHAQ)
- >30% reduction in ESR

AND

- No more than one of these variables has worsened by greater than 30%

25MG/VIAL INJECTION

02242903 ENBREL

PED

50MG/ML INJECTION

02274728 ENBREL

PED

99100373 ENBREL SURECLICK

AMG

ETANERCEPT (BRENZYS)

Limited use benefit (prior approval required).

Coverage for BRENZYS will be approved indefinitely.

1. For the treatment of severely active RHEUMATOID ARTHRITIS

- Prescribed by a rheumatologist.

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients \geq 18 years who have failed:

- MTX (oral or parenteral) at a dose \geq 20 mg weekly (\geq 15 mg weekly if patient is \geq 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX; AND
- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

2. For the treatment of ANKYLOSING SPONDYLITIS

- Prescribed by a rheumatologist
- BASDAI $>$ 4; AND
- Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks; AND for peripheral joint involvement, patient is refractory:
- Methotrexate (MTX) weekly (oral or parenteral) at 20 mg or greater (15 mg or greater if patient is $>$ 65 years of age) for more than 8 weeks; AND
- Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

50MG SOLUTION

02455323 BRENZYS

UNK

02455331 BRENZYS

UNK

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**ETANERCEPT (ERELZI)**

Limited use benefit (prior approval required).

Coverage for ERELZI will be approved indefinitely.

1. For the treatment of severely active RHEUMATOID ARTHRITIS

- Prescribed by a rheumatologist.

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX; AND
- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

2. For the treatment of moderate to severe PSORIATIC ARTHRITIS

- Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;
- PLUS a minimum of any two of the following:
- methotrexate (oral or parenteral) at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
 - leflunomide: 20mg daily for 10 weeks; OR
 - sulfasalazine at least 2g daily for 3 months; OR
 - cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) ≥ 4 ; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

Coverage beyond one year will be based on improvement according to the Psoriatic Arthritis Response Criteria (PsARC).

- Improvement in at least two of the four PsARC criteria, one of which has to be joint tenderness or swelling score, with no worsening in any of the four criteria. A response in joint count is determined by a reduction of $\geq 30\%$. A response in the Physician or Patient Global Assessment scale is determined by a reduction of 1 point.

3. For the treatment of ANKYLOSING SPONDYLITIS

- Prescribed by a rheumatologist
 - BASDAI > 4; AND
 - Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks;
- AND for peripheral joint involvement, patient is refractory:
- Methotrexate (MTX) weekly (oral or parenteral) at 20 mg or greater (15 mg or greater if patient is >65 years of age) for more than 8 weeks; AND
 - Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

4. For the treatment of severely active polyarticular JUVENILE IDIOPATHIC ARTHRITIS

- Prescribed by a rheumatologist

In children 4 years or older who meet the following criteria:

- ≥ 5 swollen joints; AND
- ≥ 3 joints with limited range of motion and/or pain/tenderness; AND
- Condition is refractory to an adequate trial of a therapeutic dose of methotrexate.

25MG SOLUTION

02462877 ERELZI

SDZ

50MG SOLUTION

02462850 ERELZI

SDZ

02462869 ERELZI

SDZ

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**GOLIMUMAB**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 50 mg every month.

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX;

AND

- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX;
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond one year will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- $>20\%$ reduction in number of tender and swollen joints; PLUS
- $>20\%$ improvement in Physician Global Assessment scale; PLUS either
- $>20\%$ improvement in Patient Global Assessment scale; OR
- $>20\%$ reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of moderate to severe PSORIATIC ARTHRITIS

Coverage is provided for an initial period of one year at a dose of 50 mg every month.

- Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;

PLUS a minimum of any two of the following:

- methotrexate weekly parenteral at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
- leflunomide: 20mg daily for 10 weeks; OR
- sulfasalazine at least 2g daily for 3 months; OR
- cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) ≥ 4 ; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

Coverage beyond one year will be based on improvement according to the Psoriatic Arthritis Response Criteria (PsARC).

- Improvement in at least two of the four PsARC criteria, one of which has to be joint tenderness or swelling score, with no worsening in any of the four criteria. A response in joint count is determined by a reduction of $\geq 30\%$. A response in the Physician or Patient Global Assessment scale is determined by a reduction of 1 point.

3. For the treatment of ANKYLOSING SPONDYLITIS

Coverage is provided for an initial period of one year at a dose of 50 mg every month.

- Prescribed by a rheumatologist
 - BASDAI > 4 ; AND
 - Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks;
- AND for peripheral joint involvement, patient is refractory:
- Methotrexate (MTX) (oral or parenteral) weekly at 20 mg or greater (15 mg or greater if patient is >65 years of age) for more than 8 weeks; AND
 - Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

Coverage beyond one year will be based on improvement in the BASDAI score.

- Improvement of at least 50% or 2 units in the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) score.

4. For the treatment of adult patients with moderately to severely active ULCERATIVE COLITIS who meet the following:

Coverage is provided for an initial period of three months at a dose of 200 mg at week 0, followed by 100 mg at week 2 and then 50 mg every four weeks thereafter.

- Prescribed by expert in gastroenterology
- Partial Mayo score > 4
- Inadequate response to conventional therapies:
 - 5-ASA 4grams/day for 6 weeks; PLUS

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- Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication.

The treating physician may utilize 100 mg every four weeks as a maintenance dose if necessary.

Coverage beyond one year will be based on a decrease in the partial Mayo score of ≥ 2 points and patients should be off corticosteroids.

50MG/0.5ML SOLUTION

02324776 SIMPONI

JSO

02324784 SIMPONI

JSO

100MG/ML SOLUTION

02413175 SIMPONI

JSO

02413183 SIMPONI

JSO

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**INFLIXIMAB (INFLECTRA)**

Limited use benefit (prior approval required).

Coverage for INFLECTRA will be approved indefinitely.

1. For the treatment of severely active RHEUMATOID ARTHRITIS

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX;

AND

- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX;
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

2. For the treatment of moderate to severe PSORIATIC ARTHRITIS

- Prescribed by a rheumatologist

Client who meet at least 2 of the following criteria:

- 5 or more swollen joints
- if less than 5 swollen joints, at least one joint proximal to, or including wrist or ankle
- more than one joint with erosion on imaging study
- dactylitis of two or more digits
- tenosynovitis refractory to oral NSAIDs and steroid injections
- enthesitis refractory to oral NSAIDs and steroid injections (not required for Achilles tendon)
- inflammatory spinal symptoms refractory to two NSAIDs (minimum four weeks trial each) and has a BASDAI greater than 4
- daily use of corticosteroids
- use of opioids > 12 hours per day for pain resulting from inflammation

AND patient is refractory to:

- a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks;

PLUS a minimum of any two of the following:

- methotrexate weekly parenteral (SC or IM) at 20mg or greater (15mg or greater if patient is >65 years of age) for more than 8 weeks; OR
- leflunomide: 20mg daily for 10 weeks; OR
- sulfasalazine at least 2g daily for 3 months; OR
- cyclosporine

OR Axial disease with both of the following:

- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) ≥ 4 ; AND
- Patient is refractory to a trial of at least two different NSAIDs at maximum tolerated doses for a combined total duration of four weeks.

3. For the treatment of ANKYLOSING SPONDYLITIS

- Prescribed by a rheumatologist
- BASDAI > 4 ; AND
- Patient is refractory to a trial of two different NSAIDs at maximum tolerated doses for a combined total duration of at least 4 weeks; AND for peripheral joint involvement, patient is refractory:
- Methotrexate (MTX) weekly at 20 mg or greater (15 mg or greater if patient is >65 years of age) for more than 8 weeks; AND
- Sulfasalazine 2 g/day for at least 3 months.

NOTE: For axial involvement, patient does not need to be tried on methotrexate or sulfasalazine.

4. For the treatment of patients with moderate to severe PSORIASIS who meet all of the following criteria:

- Prescribed by a dermatologist
 - Body surface area (BSA) involvement greater than 10% and/or significant involvement of the face, hands, feet or genital region;
- AND
- Intolerance or lack of response to phototherapy; OR
 - Inability to access phototherapy;
- AND
- Intolerance or lack of response to methotrexate (MTX) weekly oral or parenteral (SC or IM) at 20 mg or greater (15 mg or greater if patient is > 65 years of age) for more than 8 weeks;
- AND
- Intolerance or lack of response to cyclosporine; OR
 - A contraindication to methotrexate or cyclosporine.

5. For the treatment of moderately to severely active CROHN'S DISEASE

- Prescribed by a gastroenterology specialist

Patient meets the following criteria:

- Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication;
- PLUS
- Azathioprine 2 mg/kg/day for a minimum of 12 weeks; OR
 - 6-mercaptopurine 1 mg/day for a minimum of 12 weeks; OR

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS

- MTX (oral or parenteral) 15 mg per week for a minimum of 12 weeks.

6. For the treatment of FISTULIZING CROHN'S DISEASE

- Prescribed by a gastroenterology specialist

Patient meets all the following criteria:

- Patients with actively draining perianal or enterocutaneous fistulae that are refractory to a course of appropriate antibiotic therapy (e.g. ciprofloxacin with or without metronidazole for a minimum of 3 weeks);

PLUS

Patient has failed a trial of one (1) immunosuppressive agent:

- Azathioprine 2 to 2.5 mg/kg/day for a minimum of 3 months or treatment discontinued at < 3 months due to severe adverse: reactions.

OR

- 6-mercaptopurine 50-70 mg/day for a minimum of 3 months or treatment discontinued at <3 months due to severe adverse reactions.

7. For the treatment of adult patients with moderately to severely active ULCERATIVE COLITIS who meet the following:

- Prescribed by expert in gastroenterology
- Partial Mayo score > 4
- Inadequate response to conventional therapies:
 - 5-ASA 4grams/day for 6 weeks; PLUS
 - Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication.

100MG POWDER FOR SOLUTION

02419475 INFLECTRA

HOS

02470373 RENFLEXIS

UNK

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**INFLIXIMAB (REMICADE)**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for an initial three doses of 3 mg/kg, administered at 0, 2 and 6 weeks.

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX; AND
- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond the initial three doses will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- >20% reduction in number of tender and swollen joints; PLUS
- >20% improvement in Physician Global Assessment scale; PLUS either
- >20% improvement in Patient Global Assessment scale; OR
- >20% reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of moderately to severely active CROHN'S DISEASE

Coverage is provided for an initial three doses of 5 mg/kg, administered at 0, 2 and 6 weeks.

- Prescribed by a gastroenterology specialist

Patient meets the following criteria:

- Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication; PLUS
- Azathioprine 2 mg/kg/day for a minimum of 12 weeks; OR
- 6-mercaptopurine 1 mg/day for a minimum of 12 weeks; OR
- MTX (oral or parenteral) 15 mg per week for a minimum of 12 weeks.

Coverage beyond the initial three doses will be based on improvement in the CDAI or HBI scores.

- At least a 100-point reduction in the Crohn's Disease Activity Index (CDAI) OR at least a 3-point reduction in the Harvey Bradshaw Index (HBI).

3. For the treatment of FISTULIZING CROHN'S DISEASE

Coverage is provided for an initial three doses of 5 mg/kg, administered at 0, 2 and 6 weeks.

- Prescribed by a gastroenterology specialist

Patient meets all the following criteria:

- Patients with actively draining perianal or enterocutaneous fistulae that are refractory to a course of appropriate antibiotic therapy (e.g. ciprofloxacin with or without metronidazole for a minimum of 3 weeks);

PLUS

Patient has failed a trial of one (1) immunosuppressive agent:

- Azathioprine 2 to 2.5 mg/kg/day for a minimum of 3 months or treatment discontinued at < 3 months due to severe adverse reactions. OR
- 6-mercaptopurine 50-70 mg/day for a minimum of 3 months or treatment discontinued at <3 months due to severe adverse reactions.

Coverage beyond the initial three doses will be based on improvement or closure of actively draining fistulae

- Closure of individual fistulae as evidenced by no, or minimal, fistulae drainage and bleeding.

100MG/VIAL POWDER FOR SOLUTION

02244016 REMICADE

JSO

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**TOCILIZUMAB (IV)**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for 16 weeks at an initial dose of 4 mg/kg/dose every 4 weeks.

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

- MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX;

AND

- MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX;
- A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond 16 weeks, at a dose of up to 8 mg/kg/dose (maximum dose of 800 mg per infusion) every 4 weeks, is based on a 20% improvement from baseline in swollen and tender joint counts, plus a 20% improvement in 2 of 5 baseline clinical parameters.

- $>20\%$ reduction in number of tender and swollen joints; PLUS
- $>20\%$ improvement in Physician Global Assessment scale; PLUS either
- $>20\%$ improvement in Patient Global Assessment scale; OR
- $>20\%$ reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of active SYSTEMIC JUVENILE IDIOPATHIC ARTHRITIS

Initial 16-week coverage is provided at a dose of 12 mg/kg once every two weeks for children weighing < 30 kg and 8 mg/kg for children weighing ≥ 30 kg.

- Prescribed by a rheumatologist

In patients two to seventeen years of age and older who meet the following criteria:

- Have responded inadequately to non-steroidal anti-inflammatory drugs (NSAIDs) and systemic corticosteroids (with or without methotrexate), due to intolerance or lack of efficacy.

Coverage beyond 16 weeks is based on a $>30\%$ improvement in 3 of 6 baseline clinical parameters

Patient has experienced 3 of 6 of the following variables:

- $>30\%$ reduction in the number of active joints
- $>30\%$ reduction in the number of joints with loss of range of motion
- $>30\%$ improvement in the Physician Global Assessment scale
- $>30\%$ improvement in the Patient or Parent Global Assessment scale
- $>30\%$ improvement in the Child Health Assessment Questionnaire (CHAQ)
- $>30\%$ reduction in ESR

AND

- No more than one of these variables has worsened by greater than 30%

3. For the treatment of severely active polyarticular JUVENILE IDIOPATHIC ARTHRITIS

Initial 16-week coverage is provided at a dose of 10 mg/kg once every four weeks for children weighing < 30 kg and 8 mg/kg for children weighing ≥ 30 kg.

- Prescribed by a rheumatologist

In patients two years of age and older who meet the following criteria:

- ≥ 5 swollen joints; AND
- ≥ 3 joints with limited range of motion and/or pain/tenderness; AND
- Condition is refractory to an adequate trial of a therapeutic dose of methotrexate.

Coverage beyond 16 weeks is based on a $>30\%$ improvement in 3 of 6 baseline clinical parameters

Patient has experienced 3 of 6 of the following variables:

- $>30\%$ reduction in the number of active joints
- $>30\%$ reduction in the number of joints with loss of range of motion
- $>30\%$ improvement in the Physician Global Assessment scale
- $>30\%$ improvement in the Patient or Parent Global Assessment scale
- $>30\%$ improvement in the Child Health Assessment Questionnaire (CHAQ)
- $>30\%$ reduction in ESR; AND
- No more than one of these variables has worsened by greater than 30%

80MG/4ML SOLUTION

02350092 ACTEMRA

HLR

200MG/10ML SOLUTION

02350106 ACTEMRA

HLR

400MG/20ML SOLUTION

02350114 ACTEMRA

HLR

92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**TOCILIZUMAB (SC)**

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage is provided for an initial period of one year. Initial approvals for patients < 100 kg will be for a dose of 162 mg every other week up to a maximum dose of 162 mg every week (Maximum 51 doses). For patients weighing 100 kg or more, coverage is provided at a dose of 162 mg weekly (Maximum 52 doses).
 • Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

• MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX;

AND

• MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:

• A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond the initial three doses will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- >20% reduction in number of tender and swollen joints; PLUS
- >20% improvement in Physician Global Assessment scale; PLUS either
- >20% improvement in Patient Global Assessment scale; OR
- >20% reduction in the acute phase as measured by ESR or CRP.

2. For the treatment of GIANT CELL ARTERITIS in adults

Coverage is limited to 52 weeks per treatment course at a dose of 162 mg s/c weekly. Treatment can be repeated if relapse occurs.

- Patient has been diagnosed with new-onset or relapsing active giant cell arteritis; AND
- Patient is receiving moderate- to high-dose oral corticosteroids (equivalent to prednisone 20 mg to 60 mg daily).

162MG SOLUTION

02424770 ACTEMRA

HLR

TOFACITINIB CITRATE

Limited use benefit (prior approval required).

1. For the treatment of severely active RHEUMATOID ARTHRITIS

Coverage of tofacitinib in adult patients ≥ 18 years is provided at a MAXIMUM dose of 10mg daily for an initial period of one year.

- Prescribed by a rheumatologist

Coverage is provided, in combination with methotrexate (MTX) or other disease modifying anti-rheumatic drugs (DMARDs), for the reduction in signs and symptoms of severely active RA in adult patients ≥ 18 years who have failed:

• MTX (oral or parenteral) at a dose ≥ 20 mg weekly (≥ 15 mg weekly if patient is ≥ 65 years) for a minimum of 12 weeks of continuous treatment. Note: Patients who do not exhibit a clinical response to oral MTX or who experience gastrointestinal intolerance may consider a trial of parenteral MTX;

AND

• MTX in combination with at least two other DMARDs, such as sulfasalazine and hydroxychloroquine, for a minimum of 12 weeks of continuous treatment; OR, if the patient has a contraindication, failure, or intolerance to MTX:

• A combination of at least two DMARDs, such as sulfasalazine, hydroxychloroquine, azathioprine, leflunomide, cyclosporine or gold, for a minimum of 12 weeks of continuous treatment.

Coverage beyond the initial three doses will be based on a 20% improvement in 3 of 5 baseline clinical parameters.

- >20% reduction in number of tender and swollen joints; PLUS
- >20% improvement in Physician Global Assessment scale; PLUS either
- >20% improvement in Patient Global Assessment scale; OR
- >20% reduction in the acute phase as measured by ESR or CRP.

5MG TABLET

02423898 XELJANZ

PFI

92:44.00 IMMUNOSUPPRESSIVE AGENTS**ALEMTUZUMAB**

Limited use benefit (prior approval required).

Coverage is provided for two years (i.e. two treatment courses/eight doses) for adult patients who meet ALL of the following criteria:

- For the treatment of relapsing-remitting multiple sclerosis (RRMS) diagnosed according to the 2017 McDonald clinical criteria and magnetic resonance imaging (MRI) evidence; AND
- Prescribed by a specialist with experience in the treatment of multiple sclerosis; AND
- Active disease defined by clinical and imaging features (i.e. significant increase in T2 lesion load compared with that from a previous MRI scan OR at least one gadolinium-enhancing lesion); AND
- Failure to respond to full and adequate courses of at least ONE initial trial of at least 6 months of interferon, glatiramer, dimethyl fumarate or teriflunomide therapy OR documented intolerance to at least 2 therapies; AND
- At least one relapse while on at least six months of a disease modifying therapy (an interferon, glatiramer, dimethyl fumarate or teriflunomide) within the last 10 years; AND
- At least two attacks (first episode or relapse) in the previous two years, with at least one attack in the previous year; AND
- An Expanded Disability Status Scale (EDSS) score of five (5) or less.

12MG SOLUTION

02418320 LEMTRADA

GEE

CYCLOSPORINE

Limited use benefit (prior approval required).

For transplant therapy.

ST 10MG CAPSULE

02237671 NEORAL

NVR

ST 25MG CAPSULE

02150689 NEORAL

NVR

02247073 SANDOZ CYCLOSPORINE

SDZ

ST 50MG CAPSULE

02150662 NEORAL

NVR

02247074 SANDOZ CYCLOSPORINE

SDZ

ST 100MG CAPSULE

02150670 NEORAL

NVR

02242821 SANDOZ CYCLOSPORINE

SDZ

ST 100MG/ML SOLUTION

02244324 APO-CYCLOSPORINE

APX

02150697 NEORAL

NVR

FINGOLIMOD (FINGOLIMOD HYDROCHLORIDE)

Limited use benefit (prior approval required).

Initial Coverage (one year):

For the treatment of patients with Relapsing Remitting Multiple Sclerosis (RRMS) who meet ALL of the following criteria:

- Failure to respond to full and adequate courses of at least ONE initial disease-modifying therapy (an interferon, glatiramer acetate, dimethyl fumarate or teriflunomide) OR documented intolerance to at least 2 therapies; AND
- One or more clinically disabling relapses in the previous year; AND
- Significant increase in T2 lesion load compared with that from a previous MRI scan or at least one gadolinium-enhancing lesion; AND
- Requested and followed by a neurologist experienced in the management of RRMS; AND
- Recent Expanded Disability Status Scale (EDSS) score.

Renewal Coverage (two years):

- EDSS scores must be provided (exam must have occurred within that last 90 days).
- Patients must be stable or have experienced no more than 1 disabling attack/relapse in the past year.

0.5MG CAPSULE

02365480 GILENYA

NVR

92:44.00 IMMUNOSUPPRESSIVE AGENTS**MEPOLIZUMAB**

Limited use benefit (prior approval required).

For initial 12-month coverage:

For the adjunctive treatment of severe eosinophilic asthma in adults who are inadequately controlled with high-dose inhaled corticosteroids plus one or more additional asthma controller(s) (e.g. long-acting beta-agonist);

AND

• Have had a blood eosinophil count of $\geq 0.15 \times 10^9/L$ before initiation of Nucala (levels must have been drawn within 3 months of the start of treatment); OR

• Have had a blood eosinophil count of $\geq 0.3 \times 10^9/L$ within the 12-month period prior to starting Nucala

AND

• Show reversibility on spirometry (a rise in FEV₁ of at least 12% AND at least 200 mL);

AND

• Have experienced two or more clinically significant asthma exacerbations* in the past 12 months period prior to starting Nucala; or

• Have received maintenance therapy with daily oral corticosteroids for at least 3 months prior to starting Nucala.

For 12-month renewal coverage:

For the adjunctive treatment of severe eosinophilic asthma in adult patients who have experienced a decrease in clinically significant exacerbations with mepolizumab treatment as demonstrated by:

• Patient has experienced a decrease in clinically significant asthma exacerbations* with Nucala treatment; OR

• Patient's oral corticosteroid maintenance dose decreased by at least 25 % from the pre-treatment dose.

Coverage for Nucala is provided for a maximum dose of 100 mg every four weeks.

* A clinically significant asthma exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least three days or the patient visited an emergency department or was hospitalized.

100MG POWDER FOR SOLUTION

02449781 NUCALA

GSK

MYCOPHENOLATE MOFETIL

Limited use benefit (prior approval required).

For transplant therapy.

ST 250MG CAPSULE

02383780 ACH-MYCOPHENOLATE

ACC

02352559 APO-MYCOPHENOLATE

APX

02192748 CELLCEPT

HLR

02386399 JAMP-MYCOPHENOLATE

JMP

02457369 MYCOPHENOLATE MOFETIL

SAN

02371154 MYLAN-MYCOPHENOLATE

MYL

02320630 SANDOZ MYCOPHENOLATE

SDZ

02364883 TEVA-MYCOPHENOLATE

TEV

02433680 VAN-MYCOPHENOLATE

VAN

ST 200MG POWDER FOR SUSPENSION

02242145 CELLCEPT

HLR

ST 500MG TABLET

02352567 APO-MYCOPHENOLATE

APX

02237484 CELLCEPT

HLR

02380382 JAMP-MYCOPHENOLATE

JMP

02378574 MYCOPHENOLATE

ACC

02457377 MYCOPHENOLATE MOFETIL

SAN

02370549 MYLAN-MYCOPHENOLATE

MYL

02313855 SANDOZ MYCOPHENOLATE

SDZ

02348675 TEVA-MYCOPHENOLATE

TEV

02432625 VAN-MYCOPHENOLATE

VAN

MYCOPHENOLATE SODIUM

Limited use benefit (prior approval required).

For transplant therapy.

ST 180MG TABLET (ENTERIC COATED)

02372738 APO-MYCOPHENOLIC ACID

APX

02264560 MYFORTIC

NVR

92:44.00 IMMUNOSUPPRESSIVE AGENTS**MYCOPHENOLATE SODIUM**

Limited use benefit (prior approval required).

For transplant therapy.

ST **360MG TABLET (ENTERIC COATED)**

02372746 APO-MYCOPHENOLIC ACID

APX

02264579 MYFORTIC

NVR

SIROLIMUS

Limited use benefit (prior approval required).

Coverage will be provided as a second line therapy for patients failing mycophenolate mofetil.

ST **1MG/ML SOLUTION**

02243237 RAPAMUNE

PFI

ST **1MG TABLET**

02247111 RAPAMUNE

PFI

TACROLIMUS MONOHYDRATE

Limited use benefit (prior approval required).

For transplant therapy.

ST **0.5MG CAPSULE**

02243144 PROGRAF

AST

02416816 SANDOZ TACROLIMUS

SDZ

ST **1MG CAPSULE**

02175991 PROGRAF

AST

02416824 SANDOZ TACROLIMUS

SDZ

ST **5MG CAPSULE**

02175983 PROGRAF

AST

ST **0.5MG CAPSULE (EXTENDED RELEASE)**

02296462 ADVAGRAF

AST

ST **1MG CAPSULE (EXTENDED RELEASE)**

02296470 ADVAGRAF

AST

ST **3MG CAPSULE (EXTENDED RELEASE)**

02331667 ADVAGRAF

AST

ST **5MG CAPSULE (EXTENDED RELEASE)**

02296489 ADVAGRAF

AST

ST **5MG CAPSULE (IMMEDIATE RELEASE)**

02416832 SANDOZ TACROLIMUS

SDZ

5MG/ML SOLUTION

02176009 PROGRAF

AST

92:44.00 IMMUNOSUPPRESSIVE AGENTS**VEDOLIZUMAB**

Limited use benefit (prior approval required).

1. For the treatment of moderately to severely active CROHN'S DISEASE

Coverage is provided for an initial period of 14 weeks at a dose of 300 mg weeks zero, two and six and then every eight weeks. Maintenance therapy is provided at a dose not exceeding 300 mg every eight weeks.

- Prescribed by a gastroenterology specialist

Patient meets the following criteria:

- Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication; PLUS
- Azathioprine 2 mg/kg/day for a minimum of 12 weeks; OR
- 6-mercaptopurine 1 mg/day for a minimum of 12 weeks; OR
- MTX (oral or parenteral) 15 mg per week for a minimum of 12 weeks.

Coverage beyond the initial 14 week period will be based on improvement in the CDAI or HBI scores.

- At least a 100-point reduction in the Crohn's Disease Activity Index (CDAI) OR at least a 3-point reduction in the Harvey Bradshaw Index (HBI).

2. For the treatment of adult patients with moderately to severely active ULCERATIVE COLITIS who meet the following:

Coverage is provided for an initial period of 14 weeks at a dose of 300 mg at weeks zero, two and six and then every eight weeks. Maintenance therapy is provided at a dose not exceeding 300 mg every eight weeks.

- Prescribed by expert in gastroenterology
- Partial Mayo score > 4; AND
- Inadequate response to conventional therapies:
 - 5-ASA 4grams/day for 6 weeks; PLUS
 - Glucocorticoids equivalent to prednisone 40 mg/day for a minimum of 2 weeks OR treatment discontinued due to intolerance or contraindication.

Coverage beyond the initial 14 week period will be based on improvement in the partial Mayo score of ≥ 2 points.

300MG POWDER FOR SOLUTION

02436841 ENTIVIO

TAK

92:92.00 OTHER MISCELLANEOUS THERAPEUTIC AGENTS**ABOBOTULINUMTOXINA**

Limited use benefit (prior approval required).

Treatment of cervical dystonia (spasmodic torticollis) in adults; OR
Symptomatic treatment of focal spasticity affecting upper limbs in adults; OR
Lower limb spasticity in patients 2 years of age and older.

300U POWDER FOR SOLUTION

02460203 DYSPORT THERAPEUTIC

IPS

500U POWDER FOR SOLUTION

02456117 DYSPORT THERAPEUTIC

IPS

INCOBOTULINUMTOXINA

Limited use benefit (prior approval required).

For the treatment of:

- strabismus and blepharospasm associated with dystonia, including benign essential blepharospasm or VII nerve disorder in patients 12 years of age or older; OR
- cervical dystonia (spasmodic torticollis).

50UNIT/VIAL POWDER FOR SOLUTION

02371081 XEOMIN

MEZ

100U/VIAL POWDER FOR SOLUTION

02324032 XEOMIN

MEZ

ONABOTULINUMTOXINA

Limited use benefit (prior approval required).

For the treatment of:

- strabismus and blepharospasm associated with dystonia, including benign essential blepharospasm or VII nerve disorder in patients 12 years of age or older; OR
- cervical dystonia (spasmodic torticollis); OR
- urinary incontinence due to neurogenic detrusor over activity resulting from neurogenic bladder associated with MS or subcervical spinal cord injury; OR
- overactive bladder.

50IU INJECTION

09857386 BOTOX

ALL

92:92.00 OTHER MISCELLANEOUS THERAPEUTIC AGENTS**ONABOTULINUMTOXINA**

Limited use benefit (prior approval required).

For the treatment of:

- strabismus and blepharospasm associated with dystonia, including benign essential blepharospasm or VII nerve disorder in patients 12 years of age or older; OR
- cervical dystonia (spasmodic torticollis); OR
- urinary incontinence due to neurogenic detrusor over activity resulting from neurogenic bladder associated with MS or subcervical spinal cord injury; OR
- overactive bladder.

200IU INJECTION

09857387 BOTOX

ALL

100IU POWDER FOR SOLUTION

01981501 BOTOX

ALL

94:00 DEVICES**94:00.00 DEVICES****SPACER DEVICE**

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 2 spacer devices every 12 months.

DEVICE

96899962 AEROCHAMBER AC BOYZ	TRU
96899963 AEROCHAMBER AC GIRLZ	TRU
96899969 AEROCHAMBER PLUS FLOWVU LARGE	TRU
96899970 AEROCHAMBER PLUS FLOWVU MEDIUM	TRU
96899968 AEROCHAMBER PLUS FLOWVU MOUTH	TRU
96899971 AEROCHAMBER PLUS FLOWVU SMALL	TRU
96899977 AEROTRACH PLUS	UNK
96899956 COMPACT SPACE PLUS LARGE MASK	MIN
96899955 COMPACT SPACE PLUS MEDIUM MASK	MIN
96899953 COMPACT SPACE PLUS NO MASK	MIN
96899954 COMPACT SPACE PLUS SMALL MASK	MIN
99400507 E-Z SPACER	WEP
99400511 E-Z SPACER (MASK ONLY)	WEP
99400508 E-Z SPACER WITH SMALL MASK	WEP
00901012 INSPIRA CHAMBER W LARGE MASK	LUP
00900003 INSPIRA CHAMBER W MEDIUM MASK	LUP
00900001 INSPIRA CHAMBER W MOUTHPIECE	LUP
00900002 INSPIRA CHAMBER W SMALL MASK	LUP
99400501 OPTICHAMBER	AUC
96899961 OPTICHAMBER DIAMOND (CHAMBER)	AUC
96899958 OPTICHAMBER DIAMOND LARGE MASK	AUC
96899959 OPTICHAMBER DIAMOND MEDIUM MASK	AUC
96899960 OPTICHAMBER DIAMOND SMALL MASK	AUC
99400504 OPTICHAMBER LARGE MASK	AUC
99400503 OPTICHAMBER MEDIUM MASK	AUC
99400502 OPTICHAMBER SMALL MASK	AUC
99400505 OPTIHALER	AUC
99400787 POCKET CHAMBER	MCA
99400791 POCKET CHAMBER WITH ADULT MASK	MCA
99400788 POCKET CHAMBER WITH INFANT MASK	MCA
99400790 POCKET CHAMBER WITH MEDIUM MASK	MCA
99400789 POCKET CHAMBER WITH SMALL MASK	MCA
96899974 RESPICHAMBER SILICONE MEDIUM MASK	TRU
96899973 RESPICHAMBER SILICONE SMALL MASK	TRU

94:00.00 DEVICES**SPACER DEVICE**

Limited use benefit with quantity and frequency limits (prior approval is not required).

Coverage is granted for 2 spacer devices every 12 months.

DEVICE

96899972 RESPICHAMBER VHC W MOUTHPIECE TRU

94:01.00 DEVICES (DIABETIC)**INSULIN PUMP SUPPLIES**

Limited use benefit (prior approval required).

- Insulin pump supplies are approved for NIHB clients following the approval of an insulin pump by NIHB; OR
- Insulin pump supplies are approved for NIHB clients with Type 1 diabetes if an insulin pump was partially or totally covered by another insurance.

DEVICE

97799674 CARTRIDGE FOR IR200 UNK

97799342 INSET 30 INFUSION SETS UNK

99401038 INSULIN PUMP BATTERY AUC

09991458 IV3000 SMW

COMFORT ANGLED DEVICE

97799682 COMFORT ANGLED INFSET 17MM UNK

97799683 COMFORT ANGLED INFSET 17MM UNK

COMFORT SHORT ANGLED DEVICE

97799678 COMFORT SRT ANGLED INFSET 13 UNK

97799679 COMFORT SRT ANGLED INFSET 13 UNK

CONTACT DETACH DEVICE

97799672 CONTACT DETACH 90 DEGREE 6MMX60CM UNK

97799610 CONTACT DETACH 90 DEGREE 8MMX60CM UNK

INSET II DEVICE

97799685 INSET II 90 DEGREE 6MMX110CM UNK

97799687 INSET II 90 DEGREE 6MMX60CM UNK

97799684 INSET II 90 DEGREE 9MMX110CM UNK

97799686 INSET II 90 DEGREE 9MMX60CM UNK

MIO DEVICE

97799491 MIO BLUE 6MMX18 MDT

97799438 MIO BLUE 6MMX23 MDT

97799490 MIO CLEAR 6MMX32 MDT

97799489 MIO CLEAR 9MMX32 MDT

97799492 MIO PINK 6MMX18 MDT

97799437 MIO PINK 6MMX23 MDT

OMNIPOD DEVICE

09991327 PODS UNK

PARADIGM SILHOUETTE DEVICE

97799715 PARADIGM SILHOUETTE 13MMX 43 MDT

97799485 PARADIGM SILHOUETTE 13MMX18" MDT

97799716 PARADIGM SILHOUETTE 13MMX23 MDT

97799484 PARADIGM SILHOUETTE 13MMX32" MDT

97799718 PARADIGM SILHOUETTE 17MMX23 MDT

97799483 PARADIGM SILHOUETTE 17MMX32" MDT

97799719 PARADIGM SILHOUETTE 17MMX43 MDT

97799529 PARADIGM SILHOUETTE CANNULA 13MM MDT

97799528 PARADIGM SILHOUETTE CANNULA 17MM MDT

QUICK-SET DEVICE

97799486 QUICK-SET 6MMX18 MDT

94:01.00 DEVICES (DIABETIC)**INSULIN PUMP SUPPLIES**

Limited use benefit (prior approval required).

- Insulin pump supplies are approved for NIHB clients following the approval of an insulin pump by NIHB; OR
- Insulin pump supplies are approved for NIHB clients with Type 1 diabetes if an insulin pump was partially or totally covered by another insurance.

QUICK-SET DEVICE

97799744	QUICK-SET 6MMX23 TUBING	MDT
97799487	QUICK-SET 6MMX32	MDT
97799743	QUICK-SET 6MMX43 TUBING	MDT
97799742	QUICK-SET 9MMX23 TUBING	MDT
97799488	QUICK-SET 9MMX32	MDT
97799741	QUICK-SET 9MMX43 TUBING	MDT

RAPID-D DEVICE

97799650	RAPID-D 10MM/110CM	ROD
97799652	RAPID-D 10MM/60CM	ROD
97799651	RAPID-D 10MM/80CM	ROD
97799656	RAPID-D 6MM/110CM	ROD
97799658	RAPID-D 6MM/60CM	ROD
97799657	RAPID-D 6MM/80CM	ROD
97799653	RAPID-D 8MM/110CM	ROD
97799655	RAPID-D 8MM/60CM	ROD
97799654	RAPID-D 8MM/80CM	ROD

SURE-T DEVICE

97799521	PARADIGM SURE-T 29G 6MMX18	MDT
97799520	PARADIGM SURE-T 29G 6MMX23	MDT
97799519	PARADIGM SURE-T 29G 8MMX23	MDT

TENDER DEVICE

97799644	TENDER-1 17MM/110CM	ROD
97799646	TENDER-1 17MM/60CM	ROD
97799645	TENDER-1 17MM/80CM	ROD
97799638	TENDER-2 17MM/110CM	ROD
97799640	TENDER-2 17MM/60CM	ROD
97799639	TENDER-2 17MM/80CM	ROD

TENDER "MINI" DEVICE

97799647	TENDER-1 MINI INF SET 13MM/110CM	ROD
97799649	TENDER-1 MINI INFSET 13MM/60CM	ROD
97799648	TENDER-1 MINI INFSET 13MM/80CM	ROD
97799641	TENDER-2 MINI INF SET 13MM/110CM	ROD
97799643	TENDER-2 MINI INFSET 13MM/60CM	ROD
97799642	TENDER-2 MINI INFSET 13MM/80CM	ROD

ULTRAFLEX DEVICE

97799665	ULTRAFLEX 1 10MM/110CM	ROD
97799667	ULTRAFLEX 1 10MM/60CM	ROD
97799666	ULTRAFLEX 1 10MM/80CM	ROD
97799668	ULTRAFLEX 1 8MM/110CM	ROD
97799670	ULTRAFLEX 1 8MM/60CM	ROD
97799669	ULTRAFLEX 1 8MM/80CM	ROD

643MMX" DEVICE

09991616	INSET 6MMX43"	UNK
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DRESS

09991615	IV3000 STANDARD	SMW
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94:01.00 DEVICES (DIABETIC)**INSULIN PUMP SUPPLIES**

Limited use benefit (prior approval required).

- Insulin pump supplies are approved for NIHB clients following the approval of an insulin pump by NIHB; OR
- Insulin pump supplies are approved for NIHB clients with Type 1 diabetes if an insulin pump was partially or totally covered by another insurance.

3ML NEEDLE

00951417 T : SLIM X2 CARTRIDGE (SK) UNK

PATCH

09991614 MMT-174 ADHESIVE UNK

SYRINGE

97799707 RESERVOIR PARADIGM 5X1.8ML MDT

97799706 RESERVOIR PARADIGM 7X3.0ML MDT

LANCET

Limited use benefit (prior approval not required).

The number of lancets that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 600 lancets per 100 days.
- Clients managing diabetes with high risk of causing hypoglycemia will be allowed 400 lancets per 365 days.
- Clients managing diabetes medication with low risk of causing hypoglycemia will be allowed 200 lancets per 365 days.
- Clients managing diabetes through diet/lifestyle therapy only (no insulin or anti-diabetes medications) will be allowed 200 lancets per 365 days.

Please note that the test strip limit is 500/100 days. Due to lancet pack sizes, 600 per 100 days will be reimbursed.

LANCET

97799494 ACCU-CHEK FASTCLIK LANCET ROD

97799495 ACCU-CHEK FASTCLIK LANCET ROD

97799817 ACCU-CHEK MULTICLIX LANCET ROD

97799946 ACCU-CHEK MULTICLIX LANCET ROD

97799945 ACCU-CHEK SOFTCLIX LANCET ROD

97799466 BG STAR LANCET SAC

97799541 EZ HEALTH ORACLE LANCET TRE

97799825 FINGERSTIX LANCET BAY

97799292 FIRST CANADIAN HEALTH LANCETS ARA

97799826 FREESTYLE LANCET BAY

97799918 MICROLET LANCET BAY

97799810 MPD THIN LANCET (NS) MPD

97799811 MPD THIN LANCET (NS) MPD

97799807 MPD ULTRA THIN LANCET (100) MPD

97799808 MPD ULTRA THIN LANCET (200) MPD

97799970 ONETOUCH ULTRASOFT LANCET JAJ

97799348 ULTILET CLASSIC LANCET UNK

21G LANCET

97799804 MONOLET 21G LANCET TYC

28G LANCET

97799232 DROPLET PERSONAL LANCET 28G SFA

97799253 FIRST CANHEALTH 28G LANCET ARA

97799766 ITEST SAFETY 28G LANCET AUC

97799801 MONOLET THIN (MONOJECT) 28G TYC

30G LANCET

97799254 FIRST CANHEALTH 30G LANCET ARA

97799388 MEDI+SURE SOFT 30G TWIST MEC

97799389 MEDI+SURE SOFT 33G TWIST MEC

97799431 ONE TOUCH DELICA 30G LANCET JAJ

33G LANCET

97799690 BD ULTRAFINE 33G LANCET BTD

97799234 DROPLET PERSONAL LANCET 33G SFA

94:01.00 DEVICES (DIABETIC)**LANCET**

Limited use benefit (prior approval not required).

The number of lancets that will be covered by the NIHB Program will depend on the client's medical treatment:

- Clients managing diabetes with insulin will be allowed 600 lancets per 100 days.
- Clients managing diabetes with high risk of causing hypoglycemia will be allowed 400 lancets per 365 days.
- Clients managing diabetes medication with low risk of causing hypoglycemia will be allowed 200 lancets per 365 days.
- Clients managing diabetes through diet/lifestyle therapy only (no insulin or anti-diabetes medications) will be allowed 200 lancets per 365 days.

Please note that the test strip limit is 500/100 days. Due to lancet pack sizes, 600 per 100 days will be reimbursed.

33G LANCET

97799255	FIRST CANHEALTH 33G LANCET	ARA
97799767	ITEST ULTRA-THIN 33G LANCET	AUC
97799501	ONETOUCH DELICA 33G LANCET	JAJ

96:00 PHARMACEUTICAL AIDS**96:00.00 PHARMACEUTICAL AIDS****DEVICE (METHADONE)**

Limited use benefit (prior approval is not required).

Coverage is granted for 1 device.

MISCELLANEOUS

91500016	METHADONE LOCK BOX	UNK
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INFANT FORMULA

Limited use benefit (prior approval required).

Infant formula coverage for children < 1 YEAR OF AGE (Corrected Gestational Age for Prematurity).

ORAL LIQUID

95900000	ALIMENTUM	ABB
95900001	ALIMENTUM	UNK
95900003	ENFAMIL A+	MJO
95900007	ENFAMIL A+ READY TO FEED	MJO
95900152	ENFAMIL ENFACARE A+	MJO
95900026	NUTRAMIGEN A+	MJO

POWDER

95900047	ALIMENTUM PDR (400G)	ABB
95900164	ENFAMIL A+ 663G PDR	MJO
95900009	ENFAMIL ENFACARE A+ 363G PDR	MJO
95900025	NEOCATE W/ DHA & ARA 400G PDR	UNK
95900027	NUTRAMIGEN A+ LGG 566G PDR	MJO
95900035	PURAMINO A+ PDR	MJO
95900036	SIMILAC ADVANCE NEOSURE 363G	UNK

Appendix A - Limited Use Benefits and Criteria

Non-Insured Health Benefits

AA-TRIMEBUTINE	26	AEROCHAMBER PLUS FLOWVU LARGE	114	APO-METHYLPHENIDATE ER	55
ABATACEPT	96	AEROCHAMBER PLUS FLOWVU MEDIUM	114	APO-METHYLPHENIDATE SR	56
ABENOL	45	AEROCHAMBER PLUS FLOWVU MOUTH	114	APO-MONTELUKAST	70
ABIRATERONE ACETATE	9	AEROCHAMBER PLUS FLOWVU SMALL	114	APO-MOXIFLOXACIN	3
ABOBOTULINUMTOXINA	113	AEROTRACH PLUS	114	APO-MYCOPHENOLATE	111
ACAMPROSATE CALCIUM	64	AFATINIB DIMALEATE	10	APO-MYCOPHENOLIC ACID	111
ACCEL-DONEPEZIL	23	AFINITOR	13	APO-OMEPRAZOLE	79
ACCEL-SEVELAMER	69	AFINITOR DISPERZ	13	APO-OXYCODONE/ACET	38
ACCU-CHEK ADVANTAGE	66	AFLIBERCEPT	74	APO-PANTOPRAZOLE	80
ACCU-CHEK AVIVA	66	AG-ZOLMITRIPTAN ODT	63	APO-PREGABALIN	50
ACCU-CHEK COMPACT	66	ALDARA P	87	APO-RABEPRAZOLE	81
ACCU-CHEK FASTCLIX LANCET	117	ALECENSARO	10	APO-RALOXIFENE	84
ACCU-CHEK GUIDE (ON)	66	ALECTINIB	10	APO-RIVASTIGMINE	25
ACCU-CHEK GUIDE (SK)	66	ALEMTUZUMAB	110	APO-RIZATRIPTAN	61
ACCU-CHEK MOBILE BG	66	ALIMENTUM	118	APO-RIZATRIPTAN RPD	61
ACCU-CHEK MOBILE CASSETT	67	ALIMENTUM PDR (400G)	118	APO-SILDENAFIL R	35
ACCU-CHEK MULTICLIX LANCET	117	ALIROCUMAB	34	APO-SUMATRIPTAN	62
ACCU-CHEK SOFTCLIX LANCET	117	ALMOTRIPTAN	60	APO-TADALAFIL PAH	35
ACCU-TREND	67	ALMOTRIPTAN MALATE	60	APO-TENOFOVIR	5
ACET 325	45	ALPRAZOLAM	56	APO-VARENICLINE	30
ACET 650	45	ALPRAZOLAM	56	APO-VORICONAZOLE	5
ACETAMINOPHEN	45	AMBRISENTAN	36	APO-ZOLMITRIPTAN	62
ACETAMINOPHEN	45	AMERGE	60	APO-ZOLMITRIPTAN RAPID	63
ACETAMINOPHEN, CAFFEINE CITRATE, CODEINE PHOSPHATE	38	AMIKACIN SULFATE	1	APREPITANT	76
ACETAMINOPHEN, CODEINE PHOSPHATE	38	AMIKACIN SULFATE	1	APTIOM	48
ACETAMINOPHEN, OXYCODONE HYDROCHLORIDE	38	AMPHETAMINE, DEXTROAMPHETAMINE	54	AQUA-E	91
ACÉTAMINOPHÈNE	46	ANDRODERM	83	AQUA-E/ML	91
ACÉTAMINOPHÈNE BLASON SHIELD	46	ANDROGEL	83	AQUASOL E	91
ACETYLSALICYLIC ACID	37	ANTI-NAUSEANT	76	AQUASOL E VITAMIN E	91
ACETYLSALICYLIC ACID	37	APIXABAN	31	ARICEPT	23
ACETYLSALICYLIC ACID, OXYCODONE HYDROCHLORIDE	38	APO ACETAMINOPHEN	46	ASA EC	37
ACH-MYCOPHENOLATE	111	APO DIMENHYDRINATE	76	ASAPHEN	37
ACLASTA	96	APO OXAZEPAM	59	ASATAB	37
ACT AMPHETAMINE XR	54	APO-ACETAMINOPHEN	46	ASCENCIA CONTOUR	67
ACT BUPRENORPHINE/NALOXONE	45	APO-ADEFOVIR	6	ASCENCIA BREEZE 2	67
ACT CABERGOLINE	63	APO-ALMOTRIPTAN	60	ASENAPINE MALEATE	53
ACT DEXTROAMPHETAMINE SR	54	APO-ALPRAZ	56	ATASOL 15	38
ACT LEVOFLOXACIN	3	APO-AMBRISENTAN	71	ATASOL 30	38
ACT METHYLPHENIDATE ER	55	APO-ATOMOXETINE	64	ATASOL FORTE	46
ACT NABILONE	77	APO-BENZYDAMINE	73	ATIVAN	58
ACT PREGABALIN	50	APO-BOSENTAN	36	ATIVAN SUBLINGUAL	58
ACT RALOXIFENE	84	APO-BROMAZEPAM	57	ATOMOXETINE	64
ACT RIZATRIPTAN	61	APO-CABERGOLINE	63	ATOMOXETINE HYDROCHLORIDE	64
ACT RIZATRIPTAN ODT	61	APO-CLONAZEPAM	47	AUBAGIO	95
ACT SUMATRIPTAN	62	APO-CYCLOBENZAPRINE	28	AURO-CYCLOBENZAPRINE	28
ACTEMRA	108	APO-CYCLOSPORINE	110	AURO-DONEPEZIL	23
ADALIMUMAB	97	APO-DICLOFENAC	37	AURO-ENTECAVIR	7
ADCIRCA	35	APO-DONEPEZIL	23	AURO-GABAPENTIN	48
ADDERALL XR	54	APO-ENTECAVIR	7	AURO-GALANTAMINE ER	24
ADEFOVIR DIPIVOXIL	6	APO-ERLOTINIB	12	AURO-LACOSAMIDE	50
ADEMPAS	72	APO-GABAPENTIN	48	AURO-MONTELUKAST	70
ADVAGRAF	112	APO-GEFITINIB	13	AURO-MOXIFLOXACIN	3
ADVAIR 100 DISKUS	28	APO-HYDROMORPHONE	40	AURO-PANTOPRAZOLE	80
ADVAIR 125	28	APO-IMATINIB	15	AURO-PREGABALIN	50
ADVAIR 250	28	APO-IMIQUIMOD	87	AURO-RIZATRIPTAN	61
ADVAIR 250 DISKUS	28	APO-LANSOPRAZOLE	77	AURO-TENOFOVIR	5
ADVAIR 500 DISKUS	28	APO-LEVOFLOXACIN	3	AVELOX	3
AEROCHAMBER AC BOYZ	114	APO-LINEZOLID	4	AVONEX	93
AEROCHAMBER AC GIRLZ	114	APO-LORAZEPAM	58	AVONEX PEN	93
		APO-METHYLPHENIDATE	55	AXERT	60
				AXITINIB	10
				AZTREONAM	2
				BANZEL	53
				BARACLUDE	7

Appendix A - Limited Use Benefits and Criteria

Non-Insured Health Benefits

BD ULTRAFINE 33G LANCET	117	CODEINE PHOSPHATE	39	DOM-LANSOPRAZOLE	77
BENZDAMINE HYDROCHLORIDE	73	COLISTIMETHATE FOR U.S.P	4	DOM-MONTELUKAST	70
BETASERON	94	COLISTIN	4	DOM-PANTOPRAZOLE	80
BETASERON INITIATION KIT	94	COLY-MYCIN M PARENTERAL	4	DOM-PREGABALIN	50
BG STAR	67	COMFORT ANGLED INFSET 17MM	115	DOM-RABEPRAZOLE EC	82
BG STAR (ON)	67	COMFORT SRT ANGLED INFSET 13	115	DOM-RIZATRIPTAN RDT	61
BG STAR LANCET	117	COMPACT SPACE PLUS LARGE MASK	114	DOM-SUMATRIPTAN	62
BIO-DONEPEZIL	23	COMPACT SPACE PLUS MEDIUM MASK	114	DOM-ZOLMITRIPTAN	62
BIO-MONTELUKAST	70	COMPACT SPACE PLUS NO MASK	114	DONEPEZIL	23
BIO-MOXIFLOXACIN	3	COMPACT SPACE PLUS SMALL MASK	114	DONEPEZIL HYDROCHLORIDE	23
BIO-OMEPRAZOLE	79	CONCERTA	56	DOSTINEX	63
BIO-PANTOPRAZOLE	80	CONTACT DETACH 90 DEGREE 6MMX60CM	115	DROPLET PERSONAL LANCET 28G	117
BISMUTH	76	CONTACT DETACH 90 DEGREE 8MMX60CM	115	DROPLET PERSONAL LANCET 33G	117
BISMUTH SUBSALICYLATE	76	CONTOUR BG (ON)	67	DYSPORT THERAPEUTIC	113
BISMUTH SUBSALICYLATE	76	CONTOUR NEXT	67	EDOXABAN	32
BOSENTAN MONOHYDRATE	36	CONTOUR NEXT (ON)	67	ELBASVIR, GRAZOPREVR	7
BOSULIF	11	COPAXONE	93	ELIDEL	88
BOSUTINIB	11	COSENTYX	89	ELIQUIS	31
BOTOX	113	COSENTYX (STYLO)	89	EMEND	76
BREEZE 2 BG (ON)	67	COSENTYX PEN (ON)	89	EMEND TRI-PACK	76
BRENZYS	101	COTELLIC	11	EMPAGLIFLOZIN	86
BREO ELLIPTA	26	CRIZOTINIB	11	ENABLEX	90
BRIVARACETAM	47	CYCLOBENZAPRINE	28	ENBREL	101
BRIVLERA	47	CYCLOBENZAPRINE HYDROCHLORIDE	28	ENBREL SURECLICK	101
BRODALUMAB	87	CYCLOSPORINE	110	ENFAMIL A+	118
BROMAZEPAM	57	DABIGATRAN ETEXILATE MESILATE	31	ENFAMIL A+ 663G PDR	118
BROMAZEPAM	57	DABRAFENIB	12	ENFAMIL A+ READY TO FEED	118
BUPRENORPHINE HYDROCHLORIDE, NALOXONE HYDROCHLORIDE	45	DACLATASVIR	7	ENFAMIL ENFACARE A+	118
BUPROPION HYDROCHLORIDE (ZYBAN)	53	DAKLINZA	7	ENFAMIL ENFACARE A+ 363G PDR	118
CABERGOLINE	63	DAPAGLIFLOZIN PROPANEDIOL MONOHYDRATE	85	ENFAMIL POLYVISOL	91
CAFFEINE CITRATE	56	DARIFENACIN HYDROBROMIDE	90	ENFAMIL TRIVISOL	91
CAFFEINE CITRATE	56	DENOSUMAB (PROLIA)	95	ENTECAVIR MONOHYDRATE	7
CAMPRAL	64	DENOSUMAB (XGEVA)	95	ENTRESTO	37
CANAGLIFLOZIN	85	DEVICE (METHADONE)	118	ENTYVIO	113
CAPRELSA	22	DEXEDRINE	54	ENZALUTAMIDE	12
CARNITOR	69	DEXEDRINE SPANSULE	54	EPCLUSA	8
CARTRIDGE FOR IR200	115	DEXTROAMPHETAMINE	55	EPLERENONE	36
CAYSTON	2	DEXTROAMPHETAMINE SULFATE	54	ERELZI	102
CEFTAZIDIME	1	DIASTAT	58	ERLOTINIB HYDROCHLORIDE	12
CEFTAZIDIME	1	DIASTAT 2X10MG RECTAL PACK	58	ERTAPENEM	2
CELLCEPT	111	DIASTAT 2X15MG RECTAL PACK	58	ESBRIET	70
CENTRUM	91	DIAZEPAM	57	ESLICARBAZEPINE ACETATE	48
CENTRUM DHA	92	DIAZEPAM	58	ETANERCEPT	100
CENTRUM FOR WOMEN	91	DIAZEPAM (DIASTAT)	58	ETANERCEPT (BRENZYS)	101
CENTRUM JUNIOR COMPLETE	91	DICETEL	83	ETANERCEPT (ERELZI)	102
CENTRUM PRENATAL	92	DICLOFENAC SODIUM	37	EURO-ASA	37
CERITINIB	11	DICLOFENAC SODIUM (TOPICAL)	37	EVEROLIMUS	13
CERTOLIZUMAB PEGOL	99	DICLOFENAC TOPICAL	37	EVISTA	84
CESAMET	76	DIENOGEST	86	EVOLOCUMAB	35
CHAMPIX	30	DIFICID	2	EXELON	25
CHAMPIX STARTER PACK	30	DILAUDID	41	EXTAVIA	94
CHU NICOTINE ANTI SMOKING AID	29	DIMENHYDRINATE	76	EXTEMPORANEOUS MIXTURE (GENDER AFFIRMING)	92
CIMZIA	99	DIMETHYL FUMARATE	66	EXTEMPORANEOUS MIXTURE (LU)	92
CLONAPAM	47	DOLORAL 1	42	EYLEA	74
CLONAZEPAM	47	DOLORAL 5	42	EZ HEALTH ORACLE	67
CLONAZEPAM	47	DOM-ATOMOXETINE	64	EZ HEALTH ORACLE LANCET	117
COBIMETINIB	11	DOM-CYCLOBENZAPRINE	28	E-Z SPACER	114
CODEINE	39	DOM-GABAPENTIN	48	E-Z SPACER (MASK ONLY)	114
CODEINE CONTIN CR	39			E-Z SPACER WITH SMALL MASK	114
CODEINE MONOHYDRATE, CODEINE SULFATE TRIHYDRATE	39			FEBUXOSTAT	93
CODEINE PHOSPHATE	39			FENTANYL	39
				FENTANYL	39

Appendix A - Limited Use Benefits and Criteria

Non-Insured Health Benefits

LYRICA	51	M-MOXIFLOXACIN	3	NICORETTE INHALER	29
MAR-DONEPEZIL	23	MMT-174 ADHESIVE	117	NICORETTE LOZENGE	29
MAR-GABAPENTIN	48	MOGADON	59	NICORETTE QUICKMIST	30
MAR-GALANTAMINE ER	24	MONOLET 21G LANCET	117	NICOTINE (GUM)	29
MAR-MONTELUKAST	70	MONOLET THIN (MONOJECT) 28G	117	NICOTINE (INHALER)	29
MAR-MOXIFLOXACIN	3	MONTELUKAST	70	NICOTINE (LOZENGE)	29
MAR-PANTOPRAZOLE	80	MONTELUKAST SODIUM	70	NICOTINE (PATCH)	29
MAR-PREGABALIN	51	MONTELUKAST SODIUM	70	NICOTINE (SPRAY)	30
MAR-RIZATRIPTAN	61	MONUROL	9	NICOTINE GUM	29
MAR-RIZATRIPTAN ODT	61	MORPHINE HYDROCHLORIDE	42	NICOTINE TRANSDERMAL	30
MAR-ZOLMITRIPTAN	63	MORPHINE SR	43	NICOTINE TRANSDERMAL SYSTEM	30
M-ASA	37	MORPHINE SULFATE	42	NILOTINIB	17
MATERNA	92	MORPHINE SULFATE (KADIAN)	44	NINTEDANIB ESILATE	69
MAVIRET	7	MOTION SICKNESS	76	NITRAZEPAM	59
MAXALT	61	MOXIFLOXACIN	3	NOVA MAX	67
MAXALT RPD	61	MOXIFLOXACIN HYDROCHLORIDE	3	NOVA-T	66
M-DONEPEZIL	23	MOZOBIL	33	NOVO-GESIC	46
MEDI+SURE	67	M-PANTOPRAZOLE	80	NOVO-GESIC FORTE	46
MEDI+SURE (ON)	67	MPD THIN LANCET (NS)	117	NRA-PREGABALIN	51
MEDI+SURE SOFT 30G TWIST	117	MPD ULTRA THIN LANCET (100)	117	NUCALA	111
MEDI+SURE SOFT 33G TWIST	117	MPD ULTRA THIN LANCET (200)	117	NUTRAMIGEN A+	118
MED-MOXIFLOXACIN	3	M-PREGABALIN	51	NUTRAMIGEN A+ LGG 566G PDR	118
MED-RIVASTIGMINE	25	MS CONTIN SR	43	OBETICHOLIC ACID	82
MEKINIST	22	MS IR	43	OCALIVA	82
MEPOLIZUMAB	111	MULTIVITAMINS (CHILDREN AND YOUTH)	91	OCRELIZUMAB	95
MEROPENEM	2	MULTIVITAMINS (PRENATAL)	92	OCREVUS	95
MEROPENEM	2	MYCOPHENOLATE	111	OFEV	69
M-ESLON	42	MYCOPHENOLATE MOFETIL	111	OLAPARIB	18
METADOL	42	MYCOPHENOLATE MOFETIL	111	OMALIZUMAB	73
METFORMIN HYDROCHLORIDE, DAPAGLIFLOZIN	86	MYCOPHENOLATE SODIUM	111	OMEPRAZOLE	79
METFORMIN HYDROCHLORIDE, EMPAGLIFLOZIN	86	MYFORTIC	111	OMEPRAZOLE MAGNESIUM	79
METHADONE HYDROCHLORIDE (METADOL)	42	MYLAN-ALMOTRIPTAN	60	OMEPRAZOLE-20	79
METHADONE LOCK BOX	118	MYLAN-	45	ONABOTULINUMTOXINA	113
METHYLPHENIDATE	55	BUPRENORPHINE/NALOXONE		ONBREZ BREEZHALER	27
METHYLPHENIDATE HYDROCHLORIDE	55	MYLAN-GALANTAMINE ER	24	ONE A DAY WOMEN	91
MICROLET LANCET	117	MYLAN-LANSOPRAZOLE	77	ONE TOUCH DELICA 30G LANCET	117
MIDOSTAURIN	17	MYLAN-MYCOPHENOLATE	111	ONE TOUCH ULTRA	68
MINT-EPLERENONE	36	MYLAN-PANTOPRAZOLE T	80	ONETOUCH DELICA 33G LANCET	118
MINT-MONTELUKAST	70	MYLAN-RIZATRIPTAN ODT	61	ONETOUCH ULTRASOFT LANCET	117
MINT-PANTOPRAZOLE	80	MYLAN-SUMATRIPTAN	62	ONETOUCH VERIO	68
MINT-PREGABALIN	51	MYLAN-TENOFOVIR DISOPROXIL	6	ONETOUCH VERIO (ON)	68
MINT-ZOLMITRIPTAN	63	MYLAN-VANCOMYCIN	4	ONGLYZA	84
MIO BLUE 6MMX18	115	MYRBETRIQ	90	OPIOID COMPOUNDED	92
MIO BLUE 6MMX23	115	NABILONE	76	OPTICHAMBER	114
MIO CLEAR 6MMX32	115	NARATRIPTAN HYDROCHLORIDE	60	OPTICHAMBER DIAMOND (CHAMBER)	114
MIO CLEAR 9MMX32	115	NAT-DONEPEZIL	23	OPTICHAMBER DIAMOND LARGE MASK	114
MIO PINK 6MMX18	115	NAT-ERLOTINIB	12	OPTICHAMBER DIAMOND MEDIUM MASK	114
MIO PINK 6MMX23	115	NAT-IMATINIB	15	OPTICHAMBER DIAMOND SMALL MASK	114
MIRABEGRON	90	NAT-OMEPRAZOLE DR	79	OPTICHAMBER LARGE MASK	114
MIRENA	83	NAT-RIZATRIPTAN ODT	61	OPTICHAMBER MEDIUM MASK	114
MISC LIMITED USE COMPOUND INTERNAL	92	NAT-TENOFOVIR	6	OPTICHAMBER SMALL MASK	114
MISC LIMITED USE EXTERNAL COMPOUND MIXTURE	92	NATURES BOUNTY PRENATAL VITAMINS	92	OPTIHALER	114
MISCELLANEOUS COMPOUNDED EXTERNAL POWDER	92	NAT-ZOLMITRIPTAN	63	ORENCIA	96
MISCELLANEOUS COMPOUNDED EYE/EAR DROP	92	NEOCATE W/ DHA & ARA 400G PDR	118	OSIMERTINIB	18
MISCELLANEOUS COMPOUNDED INJECTION/INFUSION	92	NEORAL	110	OXAZEPAM	59
MISCELLANEOUS COMPOUNDED SUPPOSITORY	92	NESTL MATERNA	92	OXAZEPAM	59
		NEULASTA	32	OXCARBAZEPINE (SUSPENSION)	50
		NEUPRO	63	OXEZE TURBUHALER	27
		NEURONTIN	48	OXPAM	59
		NICHIT	29	OXYCODONE	44
		NICODERM	30	OXYCODONE HYDROCHLORIDE	44
		NICORETTE GUM	29		

Appendix A - Limited Use Benefits and Criteria

Non-Insured Health Benefits

OXYCODONE/ACET	38	PMS-ENTECAVIR	7	PROTOPIC	90
OXYCODONE-ACET	38	PMS-ERLOTINIB	12	PURAMINO A+ PDR	118
OXY-IR	44	PMS-FENTANYL MTX	39	QUICK-SET 6MMX18	115
PALBOCICLIB	18	PMS-GABAPENTIN	48	QUICK-SET 6MMX23 TUBING	116
PAL-TIZANIDINE	28	PMS-GALANTAMINE ER	24	QUICK-SET 6MMX32	116
PANTOLOC	80	PMS-HYDROMORPHONE	41	QUICK-SET 6MMX43 TUBING	116
PANTOPRAZOLE	80	PMS-IMATINIB	15	QUICK-SET 9MMX23 TUBING	116
PANTOPRAZOLE MAGNESIUM	80	PMS-LANSOPRAZOLE	77	QUICK-SET 9MMX32	116
PANTOPRAZOLE MAGNESIUM	80	PMS-LEVOFLOXACIN	3	QUICK-SET 9MMX43 TUBING	116
PANTOPRAZOLE SODIUM	80	PMS-LORAZEPAM	58	QUINSAIR	3
PANTOPRAZOLE T	80	PMS-METHYLPHENIDATE	55	RABEPRAZOLE	81
PANTOPRAZOLE-40	81	PMS-METHYLPHENIDATE ER	56	RABEPRAZOLE EC	81
PARADIGM SILHOUETTE 13MMX 43	115	PMS-MONTELUKAST	70	RABEPRAZOLE SODIUM	81
PARADIGM SILHOUETTE 13MMX18"	115	PMS-NABILONE	77	RALOXIFENE HYDROCHLORIDE	84
PARADIGM SILHOUETTE 13MMX23	115	PMS-OMEPRAZOLE	79	RAN-DONEPEZIL	23
PARADIGM SILHOUETTE 13MMX32"	115	PMS-OXYCODONE	44	RAN-FENTANYL MATRIX	39
PARADIGM SILHOUETTE 17MMX23	115	PMS-PANTOPRAZOLE	81	RAN-GABAPENTIN	48
PARADIGM SILHOUETTE 17MMX32"	115	PMS-PREGABALIN	51	RANIBIZUMAB	75
PARADIGM SILHOUETTE 17MMX43	115	PMS-PROGESTERONE	87	RAN-LANSOPRAZOLE	77
PARADIGM SILHOUETTE CANNULA 13MM	115	PMS-RABEPRAZOLE	81	RAN-MONTELUKAST	70
PARADIGM SILHOUETTE CANNULA 17MM	115	PMS-RALOXIFENE	84	RAN-NABILONE	76
PARADIGM SURE-T 29G 6MMX18	116	PMS-RIVASTIGMINE	25	RAN-OMEPRAZOLE	79
PARADIGM SURE-T 29G 6MMX23	116	PMS-RIZATRIPTAN RDT	61	RAN-PANTOPRAZOLE	81
PARADIGM SURE-T 29G 8MMX23	116	PMS-SILDENAFIL R	35	RAN-PREGABALIN	51
PARIET	81	PMS-SUMATRIPTAN	62	RAN-RABEPRAZOLE	81
PAT-GALANTAMINE ER	24	PMS-TENOFOVIR	6	RAPAMUNE	112
PAZOPANIB	19	PMS-VANCOMYCIN 1 G	5	RAPID-D 10MM/110CM	116
PDP-ACETAMINOPHEN	45	PMS-ZOLMITRIPTAN	63	RAPID-D 10MM/60CM	116
PEDIAPHEN	45	PMS-ZOLMITRIPTAN ODT	63	RAPID-D 10MM/80CM	116
PEDIATRIX	45	POCKET CHAMBER	114	RAPID-D 6MM/110CM	116
PEDIAVIT	91	POCKET CHAMBER WITH ADULT MASK	114	RAPID-D 6MM/60CM	116
PEGASYS	6	POCKET CHAMBER WITH INFANT MASK	114	RAPID-D 6MM/80CM	116
PEGETRON KIT	6	POCKET CHAMBER WITH MEDIUM MASK	114	RAPID-D 8MM/110CM	116
PEGFILGRASTIM	32	POCKET CHAMBER WITH SMALL MASK	114	RAPID-D 8MM/60CM	116
PEGINTERFERON ALFA-2A	6	PODS	115	RAPID-D 8MM/80CM	116
PEGINTERFERON ALFA-2B, RIBAVIRIN	6	POLYSACCHARIDE IRON COMPLEX	31	RATIO-LENOLTEC NO 2	38
PEGINTERFERON BETA-1A	6	POMALIDOMIDE	19	RATIO-LENOLTEC NO 3	38
PEPTO BISMOL	76	POMALYST	19	REBIF	94
PHARIXIA	73	PONATINIB HYDROCHLORIDE	19	REDDY-PROGESTERONE	87
PHARMA-LACOSAMIDE	50	PRADAXA	31	REGORAFENIB	20
PIMECROLIMUS	88	PRALUENT	34	REMICADE	107
PINAVERIUM BROMIDE	83	PRECISION XTRA	68	RENAGEL	69
PIPERACILLIN	2	PREGABALIN	50	RENFLEXIS	106
SODIUM/TAZOBACTAM SODIUM	2	PREGABALIN	51	RENVELA	69
PIPERACILLIN, TAZOBACTAM	2	PRENATAL AND POSTPARTUM VITAMINS AND MINERALS	92	REPATHA	35
PIRFENIDONE	70	PREVACID	77	RESERVOIR PARADIGM 5X1.8ML	117
PLEGRIDY	6	PREVACID FASTAB	78	RESERVOIR PARADIGM 7X3.0ML	117
PLERIXAFOR	33	PRIVA-PANTOPRAZOLE	81	RESPICHAMBER SILICONE MEDIUM MASK	114
PMS HYDROMORPHONE	41	PRO-AAS	37	RESPICHAMBER SILICONE SMALL MASK	114
PMS-ACETAMINOPHEN	38	PROCET-30	38	RESPICHAMBER VHC W MOUTHPIECE	115
PMS-AMPHETAMINES XR	54	PRO-CLONAZEPAM	47	RESTORIL	60
PMS-ATOMOXETINE	64	PRO-GABAPENTIN	48	REVATIO	35
PMS-BENZYDAMINE	73	PROGESTERONE	87	REVLIMID	15
PMS-BOSENTAN	36	PROGRAF	112	RIBAVIRIN	8
PMS-BUPRENORPHINE-NALOXONE	45	PROLIA	95	RIFAXIMIN	4
PMS-CLONAZEPAM	47	PRO-LORAZEPAM	58	RIOCIGUAT	72
PMS-CLONAZEPAM-R	47	PROMETRIUM	87	RITUXAN	20
PMS-CYCLOBENZAPRINE	28	PROPRANOLOL (HEMANGIOL)	36	RITUXIMAB	20
PMS-DIAZEPAM	57	PRO-RABEPRAZOLE	81	RIVA OXAZEPAM	59
PMS-DICLOFENAC	37			RIVA-ATOMOXETINE	64
PMS-DIMENHYDRINATE	76			RIVA-CLONAZEPAM	47
PMS-DONEPEZIL	23			RIVACOCET	38

Appendix A - Limited Use Benefits and Criteria

Non-Insured Health Benefits

RIVA-CYCLOBENZAPRINE	28	SECUKINUMAB	89	TEMPRA INFANT	45
RIVA-DONEPEZIL	23	SELEXIPAG	72	TENDER-1 17MM/110CM	116
RIVA-GABAPENTIN	48	SEPTA DONEPEZIL	23	TENDER-1 17MM/60CM	116
RIVA-LANSOPRAZOLE	77	SEPTA-ZOLMITRIPTAN-ODT	63	TENDER-1 17MM/80CM	116
RIVA-MONTELUKAST	70	SEREVENT DISKUS	28	TENDER-1 MINI INF SET 13MM/110CM	116
RIVA-MOXIFLOXACIN	3	SEVELAMER CARBONATE	69	TENDER-1 MINI INFSET 13MM/60CM	116
RIVA-OMEPRAZOLE DR	79	SEVELAMER HYDROCHLORIDE	69	TENDER-1 MINI INFSET 13MM/80CM	116
RIVA-PANTOPRAZOLE	81	SIDEKICK	68	TENDER-2 17MM/110CM	116
RIVA-PREGABALIN	51	SILDENAFIL CITRATE	35	TENDER-2 17MM/60CM	116
RIVA-RABEPRAZOLE	82	SILIQ	87	TENDER-2 17MM/80CM	116
RIVA-RABEPRAZOLE EC	81	SIMILAC ADVANCE NEOSURE 363G	118	TENDER-2 MINI INF SET 13MM/110CM	116
RIVAROXABAN	32	SIMPONI	104	TENDER-2 MINI INFSET 13MM/60CM	116
RIVAROXABAN (10)	32	SINGULAIR	70	TENDER-2 MINI INFSET 13MM/80CM	116
RIVASA	37	SIROLIMUS	112	TENOFOVIR DISOPROXIL FUMARATE	5
RIVASTIGMINE	25	SITAGLIPTIN PHOSPHATE MONOHYDRATE	85	TERIFLUNOMIDE	95
RIVASTIGMINE HYDROGEN TARTRATE	25	SITAGLIPTIN PHOSPHATE MONOHYDRATE, METFORMIN HYDROCHLORIDE	85	TESTIM	83
RIVA-ZOLMITRIPTAN	63	SOFOSBUVIR	8	TESTOSTERONE (TOPICAL)	83
RIVOTRIL	47	SOFOSBUVIR, LEDIPASVIR	8	TEVA-ALMOTRIPTAN	60
RIZATRIPTAN BENZOATE	61	SOFOSBUVIR, VELPATASVIR	8	TEVA-ALPRAZOLAM	57
RIZATRIPTAN ODT	61	SOFOSBUVIR, VELPATASVIR, VOXILAPREVIR	9	TEVA-ATOMOXETINE	64
RIZATRIPTAN RDT	61	SOVALDI	8	TEVA-BOSENTAN	36
ROTIGOTINE	63	SPACER DEVICE	114	TEVA-BROMAZEPAM	57
RUFINAMIDE	53	SPIRIT TEST STRIP (ON)	68	TEVA-CLONAZEPAM	47
RUGBY NICOTINE POLACRILEX GUM	29	STATEX	42	TEVA-CYCLOBENZAPRINE	28
RUXOLITINIB	21	STELARA	93	TEVA-DONEPEZIL	23
RYDAPT	17	STIVARGA	20	TEVA-EMTEC-30	38
SALMETEROL XINAFOATE	28	STRATTERA	64	TEVA-ERLOTINIB	12
SALMETEROL XINAFOATE, FLUTICASON PROPRIONATE	28	STRESSTABS FOR WOMEN	91	TEVA-FENTANYL	39
SANDOZ ALMOTRIPTAN	60	SUBOXONE	45	TEVA-GABAPENTIN	48
SANDOZ AMPHETAMINE XR	54	SUMATRIPTAN	62	TEVA-HYDROMORPHONE	41
SANDOZ ATOMOXETINE	64	SUMATRIPTAN DF	62	TEVA-IMATINIB	15
SANDOZ BOSENTAN	36	SUMATRIPTAN SUCCINATE	62	TEVA-LACOSAMIDE	50
SANDOZ CYCLOSPORINE	110	SUNITINIB MALATE	21	TEVA-LANSOPRAZOLE	77
SANDOZ DONEPEZIL	23	SUPEUDOL	44	TEVA-LEVOFLOXACIN	3
SANDOZ FENTANYL	39	SURE STEP	68	TEVA-LORAZEPAM	58
SANDOZ LACOSAMIDE	50	SURETEST (ON)	68	TEVA-METHYLPHENIDATE	56
SANDOZ LANSOPRAZOLE	77	SUTENT	21	TEVA-MONTELUKAST	70
SANDOZ LEVOFLOXACIN	3	SYMBICORT 100 TURBUHALER	27	TEVA-MORPHINE SR	43
SANDOZ LINEZOLID	4	SYMBICORT 200 TURBUHALER	27	TEVA-MOXIFLOXACIN	3
SANDOZ METHYLPHENIDATE SR	56	SYNJARDY	86	TEVA-MYCOPHENOLATE	111
SANDOZ MONTELUKAST	70	T : SLIM X2 CARTRIDGE (SK)	117	TEVA-NABILONE	77
SANDOZ MORPHINE SR	43	TACROLIMUS (PROTOPIC)	90	TEVA-NARATRIPTAN	60
SANDOZ MOXIFLOXACIN	3	TACROLIMUS MONOHYDRATE	112	TEVA-OMEPRAZOLE	79
SANDOZ MYCOPHENOLATE	111	TADALAFIL	35	TEVA-OXYCOCET	38
SANDOZ NARATRIPTAN	60	TAFINLAR	12	TEVA-OXYCODAN	38
SANDOZ OMEPRAZOLE	79	TAGRISSO	18	TEVA-PANTOPRAZOLE	81
SANDOZ	38	TALTZ	88	TEVA-PANTOPRAZOLE MAGNESIUM	80
OXYCODONE/ACETAMINOPHEN		TARCEVA	12	TEVA-PREGABALIN	51
SANDOZ PANTOPRAZOLE	81	TARO-DICLOFENAC	37	TEVA-PROGESTERONE	87
SANDOZ PREGABALIN	51	TARO-IMIQUIMOD PUMP	87	TEVA-RABEPRAZOLE	82
SANDOZ RABEPRAZOLE	81	TARO-SUMATRIPTAN	62	TEVA-RIZATRIPTAN ODT	61
SANDOZ RIVASTIGMINE	25	TARO-TESTOSTERONE	83	TEVA-SILDENAFIL R	35
SANDOZ RIZATRIPTAN ODT	61	TARO-ZOLEDRONIC ACID	96	TEVA-SUMATRIPTAN	62
SANDOZ SUMATRIPTAN	62	TASIGNA	17	TEVA-SUMATRIPTAN DF	62
SANDOZ TACROLIMUS	112	TECFIDERA	66	TEVA-TEMAZEPAM	60
SANDOZ VORICONAZOLE	5	TECTA	80	TEVA-TENOFOVIR	6
SANDOZ ZOLMITRIPTAN	63	TEMAZEPAM	60	TEVA-TOBRAMYCIN	1
SANDOZ ZOLMITRIPTAN ODT	63	TEMAZEPAM	60	TEVA-VORICONAZOLE	5
SAPHRIS	53	TEMPRA CHILDREN'S	45	TEVA-ZOLMITRIPTAN	63
SAXAGLIPTIN HYDROCHLORIDE	84	TEMPRA CHILDREN'S DOUBLE STRENGTH	45	TEVA-ZOLMITRIPTAN OD	63
SAXAGLIPTIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE	84				

Appendix A - Limited Use Benefits and Criteria

Non-Insured Health Benefits

THRIVE GUM (NS)	29	VENCLEXTA	22
THRIVE NICOTINE LOZENGES	29	VENETOCLAX	22
THRIVE NICOTINELL GUM	29	VERTEPORFIN	75
TIZANIDINE	28	VFEND	5
TIZANIDINE HYDROCHLORIDE	28	VIMPAT	50
TOBI PODHALER	1	VIREAD	5
TOBRAMYCIN	1	VISANNE	86
TOBRAMYCIN	1	VISUDYNE	75
TOBRAMYCIN INHALATION	1	VITAMIN E	91
TOBRAMYCINE	1	VITAMIN E	91
TOCILIZUMAB (IV)	108	VOLIBRIS	36
TOCILIZUMAB (SC)	109	VORICONAZOLE	5
TOFACITINIB CITRATE	109	VOSEVI	9
TOVIAZ	90	VOTRIENT	19
TRACLEER	36	VYVANSE	55
TRAJENTA	84	WAMPOLE COMPLETE MULT-PRE AND POST NATAL WITH FOLIC ACID	92
TRAMETINIB	22	XALKORI	11
TRANSDERMAL NICOTINE	30	XANAX	57
TRANSDERMAL NICOTINE PATCHDAY	30	XANAX TS	57
TRAVEL	76	XARELTO	32
TRAVEL ON	76	XELJANZ	109
TRIA TEC-30	38	XEOMIN	113
TRIAZOLAM	60	XGEVA	95
TRIAZOLAM	60	XIGDUO	86
TRILEPTAL	50	XOLAIR	73
TRIMEBUTINE	26	XTANDI	12
TRIMEBUTINE MALEATE	26	ZAXINE	4
TROSEC	90	ZELBORAF	22
TROSPIMUM CHLORIDE	90	ZENHALE	27
TRUE TRACK	68	ZEPATIER	7
TRUETEST	68	ZOLEDRONIC ACID	96
TYLENOL	45	ZOLEDRONIC ACID MONOHYDRATE	96
TYLENOL EXTRA STRENGTH	46	ZOLMITRIPTAN	62
TYLENOL JR STRENGTH FASTMELTS	46	ZOLMITRIPTAN	63
TYLENOL JUNIOR STRENGTH	46	ZOLMITRIPTAN ODT	63
TYLENOL WITH CODEINE NO.2	38	ZOMIG	63
TYLENOL WITH CODEINE NO.3	38	ZOMIG RAPIMELT	63
ULIPRISTAL ACETATE	83	ZYBAN	53
ULORIC	93	ZYDELIG	14
ULTILET CLASSIC LANCET	117	ZYKADIA	11
ULTRAFLEX 1 10MM/110CM	116	ZYTIGA	9
ULTRAFLEX 1 10MM/60CM	116	ZYVOXAM	4
ULTRAFLEX 1 10MM/80CM	116		
ULTRAFLEX 1 8MM/110CM	116		
ULTRAFLEX 1 8MM/60CM	116		
ULTRAFLEX 1 8MM/80CM	116		
UPTRAVI	72		
USTEKINUMAB	93		
VALIUM	58		
VALSARTAN, SACUBITRIL	37		
VANCOMYCIN	4		
VANCOMYCIN HYDROCHLORIDE	4		
VANCOMYCIN HYDROCHLORIDE (INJECTION)	4		
VANDETANIB	22		
VAN-MYCOPHENOLATE	111		
VAN-OMEPRAZOLE	79		
VAN-RIZATRIPTAN	61		
VAN-RIZATRIPTAN ODT	62		
VAN-ZOLMITRIPTAN ODT	63		
VARENICLINE TARTRATE	30		
VEDOLIZUMAB	113		
VEMURAFENIB	22		

**APPENDIX B
SPECIAL FORMULARY FOR
CHRONIC RENAL FAILURE PATIENTS**

The Special Formulary for Chronic Renal Failure Patients defines selected drugs (for example: darbepoetin alfa, calcium products, water-soluble multivitamin products and selected nutritional supplements formulated for renal patients) that are covered for identified eligible NIHB clients in chronic renal failure.

These drugs are covered in addition to the drugs and products listed in the NIHB Drug Benefit List.

08:00 ANTI-INFECTIVE AGENTS**08:12.02 AMINOGLYCOSIDES****GENTAMICIN SULFATE****10MG/ML INJECTION**

02225123 CIDOMYCIN UNK

10MG SOLUTION

02470462 GENTAMICIN TEL

40MG SOLUTION

02457008 GENTAMICIN TEL

08:12.06 CEPHALOSPORINS**CEFAZOLIN SODIUM****500MG POWDER FOR SOLUTION**

02437104 CEFAZOLIN RAX

1G POWDER FOR SOLUTION

02465469 CEFAZOLIN UNK

10G POWDER FOR SOLUTION

02452162 CEFAZOLIN FKD

02465477 CEFAZOLIN UNK

20G POWDER FOR SOLUTION

02237141 CEFAZOLIN FKD

100G POWDER FOR SOLUTION

02401029 CEFAZOLIN FKD

**20:00 BLOOD FORMATION
COAGULATION AND
THROMBOSIS****20:16.00 HEMATOPOIETIC AGENTS****DARBEPOETIN ALFA****25MCG/ML SOLUTION**

02392313 ARANESP AMG

40MCG/ML SOLUTION

02392321 ARANESP AMG

60MCG/ML SOLUTION

02246348 ARANESP AMG

100MCG/ML SOLUTION

02391740 ARANESP AMG

02391759 ARANESP AMG

02392348 ARANESP AMG

99004917 ARANESP AMG

99004925 ARANESP AMG

200MCG/ML SOLUTION

02391767 ARANESP AMG

02391775 ARANESP AMG

02391783 ARANESP AMG

02392356 ARANESP AMG

99004909 ARANESP AMG

99004933 ARANESP AMG

500MCG/ML SOLUTION

02391791 ARANESP AMG

20:16.00 HEMATOPOIETIC AGENTS**DARBEPOETIN ALFA****500MCG/ML SOLUTION**

02391805 ARANESP AMG

02391821 ARANESP AMG

02392364 ARANESP AMG

09857185 ARANESP AMG

EPOETIN ALFA**1,000U/0.5ML SOLUTION**

02231583 EPREX JSO

2,000U/0.5ML SOLUTION

02231584 EPREX JSO

3,000U/0.3ML SOLUTION

02231585 EPREX JSO

4,000U/0.4ML SOLUTION

02231586 EPREX JSO

5000U/0.5ML SOLUTION

02243400 EPREX JSO

6000U/0.6ML SOLUTION

02243401 EPREX JSO

8000U/0.8ML SOLUTION

02243403 EPREX JSO

10,000/ML SOLUTION

02231587 EPREX JSO

20,000U/0.5ML SOLUTION

02243239 EPREX JSO

30,000U/0.75ML SOLUTION

02288680 EPREX JSO

40,000U/ML SOLUTION

02240722 EPREX JSO

**40:00 ELECTROLYTIC, CALORIC,
AND WATER BALANCE****40:12.00 REPLACEMENT PREPARATIONS****CALCIUM****250MG TABLET**

00645958 CALCIUM NOP

625MG TABLET (COATED)

00682047 APOCAL APX

CALCIUM CARB-GLUCONOLACTATE**500MG TABLET**

02232482 CALCIUMSANDOZ FORTE GSK

1,000MG TABLET

02232483 GRAMCAL GSK

SODIUM PHOSPHATE**123MG POWDER FOR SOLUTION**

80027202 PHOSPHATE NOVARTIS NVR

500MG TABLET

00225819 PHOSPHATE-NOVARTIS NVC

The Special Formulary for Chronic Renal Failure Patients defines selected drugs (for example: darbepoetin alfa, calcium products, water-soluble multivitamin products and selected nutritional supplements formulated for renal patients) that are covered for identified eligible NIHB clients in chronic renal failure.

These drugs are covered in addition to the drugs and products listed in the NIHB Drug Benefit List.

40:12.00 REPLACEMENT PREPARATIONS**ZINC GLUCONATE****50MG TABLET**

00503169 ZINC	VTH
00505463 ZINC	JAM

56:00 GASTROINTESTINAL DRUGS**56:04.00 ANTACIDS AND ADSORBENTS****ALUMINUM HYDROXIDE****500MG CAPSULE**

02135620 BASALJEL	AUP
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320MG/ML SUSPENSION

00572527 ALUGEL	ATL
-----------------	-----

325MG/5ML SUSPENSION

02125862 AMPHOJEL	AUP
-------------------	-----

600MG TABLET

02124971 AMPHOJEL	AUP
-------------------	-----

CALCIUM**500MG TABLET**

01970240 TUMS	GSK
---------------	-----

750MG TABLET

01967932 TUMS EXTRA STRENGTH	GSK
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1,000MG TABLET

02151138 TUMS ULTRA STRENGTH	GSK
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SODIUM BICARBONATE**500MG TABLET**

80030520 JAMP-SODIUM BICARBONATE	JMP
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80022194 SANDOZ SODIUM BICARBONATE	SDZ
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84:00 SKIN AND MUCOUS MEMBRANE AGENTS (SMMA)**84:04.04 SMMA - ANTIBIOTICS****GENTAMICIN SULFATE****1MG OINTMENT**

00872881 PMS-GENTAMICIN	PMS
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88:00 VITAMINS**88:28.00 MULTIVITAMIN PREPARATIONS****MULTIVITAMINS****TABLET/CAPLET**

00123803 B COMPLEX PLUS C	JAM
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80007498 BC VITAMINS	WNP
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02245391 DIAMINE	EUR
------------------	-----

80063438 M-PLAVITE	MAN
--------------------	-----

80001432 RENAVIDE	MAC
-------------------	-----

00558796 STRESS PLEX	JAM
----------------------	-----

96:00 PHARMACEUTICAL AIDS**96:00.00 PHARMACEUTICAL AIDS****NUTRITIONAL SUPPLEMENT****ORAL LIQUID**

00999483 BOOST DIABETIC O/L	NVC
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09853154 BOOST FRUIT BEVERAGE	NES
-------------------------------	-----

95999970 BOOST HIPROTEIN	NES
--------------------------	-----

95999963 BOOST ORIGINAL	NES
-------------------------	-----

95999975 BOOST PLUS	NES
---------------------	-----

97904341 ENSURE	ABB
-----------------	-----

00801054 ENSURE HIGH PROTEIN	ABB
------------------------------	-----

97904333 ENSURE PLUS	ABB
----------------------	-----

97904317 ENSURE WITH FIBRE	ABB
----------------------------	-----

00920347 GLUCERNA	ABB
-------------------	-----

09854392 GLUCERNA	ABB
-------------------	-----

99004267 GLUCERNA	ABB
-------------------	-----

09854393 GLUCERNA TUBE FEEDING	ABB
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09853723 NEPRO	ABB
----------------	-----

99002639 NEPRO	ABB
----------------	-----

99100702 NEPRO CARB STEADY	ABB
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00907995 NOVASOURCE	NVC
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09854258 NOVASOURCE	NES
---------------------	-----

99101180 PEDIASURE COMPLETE	ABB
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GROW&GAIN	
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09853731 SUPLENA	ABB
------------------	-----

99002647 SUPLENA	ABB
------------------	-----

POWDER

09991056 RESOURCE BENEPROTEIN	NVC
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99003783 RESOURCE BENEPROTEIN	NES
-------------------------------	-----

ALUGEL	2
ALUMINUM HYDROXIDE	2
AMPHOJEL	2
APOCAL	1
ARANESP	1
B COMPLEX PLUS C	2
BASALJEL	2
BC VITAMINS	2
BOOST DIABETIC O/L	2
BOOST FRUIT BEVERAGE	2
BOOST HIPROTEIN	2
BOOST ORIGINAL	2
BOOST PLUS	2
CALCIUM	1
CALCIUM	1
CALCIUM CARB- GLUCONOLACTATE	1
CALCIUMSANDOZ FORTE	1
CEFAZOLIN	1
CEFAZOLIN SODIUM	1
CIDOMYCIN	1
DARBEPOETIN ALFA	1
DIAMINE	2
ENSURE	2
ENSURE HIGH PROTEIN	2
ENSURE PLUS	2
ENSURE WITH FIBRE	2
EPOETIN ALFA	1
EPREX	1
GENTAMICIN	1
GENTAMICIN SULFATE	1
GLUCERNA	2
GLUCERNA TUBE FEEDING	2
GRAMCAL	1
JAMP-SODIUM BICARBONATE	2
M-PLAVITE	2
MULTIVITAMINS	2
NEPRO	2
NEPRO CARB STEADY	2
NOVASOURCE	2
NUTRITIONAL SUPPLEMENT	2
PEDIASURE COMPLETE GROW&GAIN	2
PHOSPHATE NOVARTIS	1
PHOSPHATE-NOVARTIS	1
PMS-GENTAMICIN	2
RENAVITE	2
RESOURCE BENEPROTEIN	2
SANDOZ SODIUM BICARBONATE	2
SODIUM BICARBONATE	2
SODIUM PHOSPHATE	1
STRESS PLEX	2
SUPLENA	2
TUMS	2
TUMS EXTRA STRENGTH	2
TUMS ULTRA STRENGTH	2
ZINC	2
ZINC GLUCONATE	2

APPENDIX C
PALLIATIVE CARE FORMULARY

Effective April 1, 2009, recipients diagnosed with a terminal illness and are near the end of life will be eligible to receive a list of supplemental benefits that are not included in the NIHB Drug Benefit List. The Palliative Care Formulary includes medications and nutritional supplements used to provide comfort to those near the end of life.

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Please note: During the six month coverage period, a maximum 30 day supply will be reimbursed at any one time.

12:00 AUTONOMIC DRUGS

12:08.08 ANTIMUSCARINICS / ANTISPASMODICS

ATROPINE SULFATE

0.4MG/ML SOLUTION

02094681 ATROPINE	ALV
00960624 ATROPINE SULFATE	UNK

0.6MG/ML SOLUTION

00012076 ATROPINE SULFATE	GSK
00392693 ATROPINE SULFATE	SDZ
00392782 ATROPINE SULFATE	SDZ

GLYCOPYRROLATE

0.2MG/ML LIQUID

02382857 GLYCOPYRROLATE	OMG
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0.2MG SOLUTION

02382849 GLYCOPYRROLATE MULTIDOSE	OMG
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0.2MG/ML SOLUTION

02039508 GLYCOPYRROLATE	SDZ
-------------------------	-----

1MG SOLUTION

02469332 CUVPOSA	PEI
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HYOSCINE BUTYLBROMIDE

20MG/ML SOLUTION

00363839 BUSCOPAN	SAC
02229868 HYOSCINE BUTYLBROMIDE	SDZ

SCOPOLAMINE HYDROBROMIDE

0.4MG/ML SOLUTION

00541869 SCOPOLAMINE	PFI
02242810 SCOPOLAMINE	OMG

0.6MG/ML SOLUTION

00541877 SCOPOLAMINE	PFI
02242811 SCOPOLAMINE	OMG

28:00 CENTRAL NERVOUS SYSTEM AGENTS

28:04.92 GENERAL ANESTHETICS, MISC.

KETAMINE HYDROCHLORIDE

10MG/ML SOLUTION

00224391 KETALAR	ERF
02246795 KETAMINE	SDZ
02387301 KETAMINE	SDZ

50MG/ML SOLUTION

00224405 KETALAR	ERF
02246796 KETAMINE	SDZ
02387328 KETAMINE	SDZ
02387336 KETAMINE	SDZ

28:08.08 OPIATE AGONISTS

EXTEMPORANEOUS MIXTURE

INJECTION

99506019 FENTANYL STERILE INFUSION	UNK
99506017 HYDROMORPHONE HP STERILE INFUSION	UNK
99506018 MORPHINE HP STERILE INFUSION	UNK

FENTANYL

12MCG/HR PATCH

02454440 APO-FENTANYL MATRIX	APX
02334186 DURAGESIC	JSO
99100480 FENTANYL	JNO
02376768 PAT-FENTANYL MATRIX	KLA

25MCG/HR PATCH

02304120 FENTANYL TRANSDERMAL SYSTEM	ACG
02376776 PAT-FENTANYL MATRIX	KLA
02325403 RAN-FENTANYL MATRIX	RBY

37MCG/HR PATCH

02386860 CO FENTANYL	OBT
02327139 SANDOZ FENTANYL	SDZ

50MCG/HR PATCH

02304139 FENTANYL TRANSDERMAL SYSTEM	ACG
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28:08.08 OPIATE AGONISTS

FENTANYL

50MCG/HR PATCH

02376784 PAT-FENTANYL MATRIX	KLA
02325411 RAN-FENTANYL MATRIX	RBY

75MCG/HR PATCH

02304147 FENTANYL TRANSDERMAL SYSTEM	ACG
02376792 PAT-FENTANYL MATRIX	KLA
02325438 RAN-FENTANYL MATRIX	RBY

100MCG/HR PATCH

02304155 FENTANYL TRANSDERMAL SYSTEM	ACG
02376806 PAT-FENTANYL MATRIX	KLA
02325446 RAN-FENTANYL MATRIX	RBY

FENTANYL CITRATE

50MCG LIQUID

02384124 FENTANYL CITRATE SDZ	SDZ
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50MCG/ML SOLUTION

00888346 FENTANYL CITRATE	PFI
02240434 FENTANYL CITRATE	SDZ

HYDROMORPHONE HYDROCHLORIDE

2MG/ML SOLUTION

02145901 HYDROMORPHONE	SDZ
------------------------	-----

10MG SOLUTION

02460610 HYDROMORPHONE HYDROCHLORIDE HP 10	RAX
--	-----

10MG/ML SOLUTION

02145928 HYDROMORPHONE HP	SDZ
---------------------------	-----

20MG/ML SOLUTION

02145936 HYDROMORPHONE HP	SDZ
---------------------------	-----

50MG/ML SOLUTION

02146126 HYDROMORPHONE HP	SDZ
99003163 HYDROMORPHONE HP	UNK

100MG/ML SOLUTION

02244797 HYDROMORPHONE HP FORTE	SDZ
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28:08.08 OPIATE AGONISTS

METHADONE HYDROCHLORIDE (BC ONLY)

POWDER

09991180 METHADONE PDR (PAIN)	UNK
09991552 METHADONE PDR (PALLIATIVE)	UNK

METHADONE HYDROCHLORIDE (METADOL)

1MG/ML SOLUTION

02247694 METADOL	PAL
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1MG TABLET

02247698 METADOL	PAL
------------------	-----

5MG TABLET

02247699 METADOL	PAL
------------------	-----

10MG TABLET

02247700 METADOL	PAL
------------------	-----

25MG TABLET

02247701 METADOL	PAL
------------------	-----

MORPHINE SULFATE

2MG/ML LIQUID

02242484 MORPHINE SULFATE	SDZ
---------------------------	-----

10MG LIQUID

00392588 MORPHINE SULFATE	SDZ
---------------------------	-----

15MG LIQUID

00392561 MORPHINE SULFATE	SDZ
---------------------------	-----

50MG/ML LIQUID

02137267 MORPHINE SULPHATE	HOS
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0.5MG/ML SOLUTION

02021056 MORPHINE LP EPIDURAL	SDZ
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01949047 MORPHINE-EPD	PFI
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1MG/ML SOLUTION

02021048 MORPHINE LP	SDZ
----------------------	-----

01980696 MORPHINE SULFATE	SDZ
---------------------------	-----

01949055 MORPHINE-EPD	PFI
-----------------------	-----

2MG/ML SOLUTION

00850314 MORPHINE SULFATE	PFI
---------------------------	-----

01964437 MORPHINE SULFATE	SDZ
---------------------------	-----

5MG/ML SOLUTION

01964429 MORPHINE SULFATE	SDZ
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28:08.08 OPIATE AGONISTS

MORPHINE SULFATE

10MG/ML SOLUTION

00850322 MORPHINE SULFATE PFI

25MG/ML SOLUTION

00676411 MORPHINE HP SDZ

50MG/ML SOLUTION

00617288 MORPHINE HP SDZ

28:12.04 ANTICONVULSANTS - BARBITURATES

PHENOBARBITAL

30MG SOLUTION

02304082 PHENOBARBITAL SODIUM SDZ

120MG SOLUTION

02304090 PHENOBARBITAL SODIUM SDZ

28:12.12 ANTICONVULSANTS - HYDANTOINS

PHENYTOIN

50MG LIQUID

00780626 PHENYTOIN SODIUM SDZ

28:16.08 ANTIPSYCHOTIC AGENTS

METHOTRIMEPRAZINE HYDROCHLORIDE

25MG/ML SOLUTION

01927698 NOZINAN SAC

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

DIAZEPAM

5MG/ML SOLUTION

00399728 DIAZEPAM SDZ

02386143 DIAZEPAM SDZ

DIAZEPAM (DIASTAT)

5MG/ML GEL

02238162 DIASTAT VAE

09853340 DIASTAT 2X10MG RECTAL PACK ELN

09853430 DIASTAT 2X15MG RECTAL PACK ELN

28:24.08 ANXIOLYTICS, SEDATIVES AND HYPNOTICS - BENZODIAZEPINES

LORAZEPAM

4MG/ML LIQUID

02243278 LORAZEPAM SDZ

2MG/ML SOLUTION

02438704 LORAZEPAM SDZ

MIDAZOLAM

1MG/ML SOLUTION

02240285 MIDAZOLAM SDZ

02242904 MIDAZOLAM FKD

02243934 MIDAZOLAM NOP

5MG SOLUTION

02423766 MIDAZOLAM PFI

5MG/ML SOLUTION

02240286 MIDAZOLAM SDZ

02242905 MIDAZOLAM FKD

02243935 MIDAZOLAM NOP

02382903 MIDAZOLAM SDZ

40:00 ELECTROLYTIC, CALORIC, AND WATER BALANCE

40:28.08 LOOP DIURETICS

FUROSEMIDE

10MG LIQUID

00527033 FUROSEMIDE SDZ

10MG/ML SOLUTION

02382539 FUROSEMIDE SDZ

02384094 FUROSEMIDE ALV

52:00 EYE, EAR, NOSE AND THROAT (EENT)

52:92.00 MISCELLANEOUS EENT DRUGS

ARTIFICIAL SALIVA

0.05MG SPRAY

02238696 MOISTIR PMS

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56:00 GASTROINTESTINAL DRUGS

56:08.00 ANTIDIARRHEA AGENTS

DIPHENOXYLATE HYDROCHLORIDE, ATROPINE SULFATE

2.5MG & 0.025MG TABLET

00036323 LOMOTIL PFI

56:22.20 5-HT3 RECEPTOR ANTAGONISTS

GRANISETRON HYDROCHLORIDE

1MG LIQUID

02322765 GRANISETRON HYDROCHLORIDE OMG

1MG/ML SOLUTION

02385414 GRANISETRON SDZ

ONDANSETRON HYDROCHLORIDE

2MG/ML INJECTION

09857323 ONDANSETRON (ON) NOP

02291703 ONDANSETRON W/P APX

09857324 ZOFRAN (ON) GSK

09857325 ZOFRAN (ON) GSK

2MG LIQUID

02271761 ONDANSETRON OMEGA - (PRESERVATIVE FREE SINGLE DOSE VIALS) OMG

02271788 ONDANSETRON OMEGA -(WITH PRESERVATIVE MULTIDOSE VIAL) OMG

2MG SOLUTION

02420414 JAMP-ONDANSETRON JMP

02420422 JAMP-ONDANSETRON JMP

02462257 ONDANSETRON RAX

02464578 ONDANSETRON RAX

02279436 ONDANSETRON -(WITH PRESERVATIVE) SDZ

02461420 ONDANSETRON BP AUR

02213745 ZOFRAN NVR

2MG/ML SOLUTION

02265524 ONDANSETRON TEV

02274418 ONDANSETRON SDZ

02279428 ONDANSETRON SDZ

56:22.20 5-HT3 RECEPTOR ANTAGONISTS

ONDANSETRON HYDROCHLORIDE

2MG/ML SOLUTION

02390019 ONDANSETRON MYL

02390051 ONDANSETRON MYL

56:22.92 MISCELLANEOUS ANTIEMETICS

NABILONE

0.25MG CAPSULE

02441497 APO-NABILONE APX

02345897 APP-NABILONE UNK

02380897 PMS-NABILONE PMS

0.5MG CAPSULE

02441500 APO-NABILONE APX

02345927 APP-NABILONE UNK

1MG CAPSULE

02441519 APO-NABILONE APX

02345935 APP-NABILONE UNK

SCOPOLAMINE

1.5MG PATCH

00550094 TRANSDERM-V NVC

80024336 TRANSDERM-V NVR

56:28.12 HISTAMINE H2-ANTAGONISTS

RANITIDINE HYDROCHLORIDE

25MG/ML SOLUTION

02256711 RANITIDINE SDZ

56:32.00 PROKINETIC AGENTS

METOCLOPRAMIDE HYDROCHLORIDE

5MG/ML LIQUID

02185431 METOCLOPRAMIDE SDZ

02243563 METOCLOPRAMIDE OMEGA OMG

56:92.00 MISCELLANEOUS GI DRUGS

METHYLNALTREXONE BROMIDE

20MG SOLUTION

02308215 RELISTOR SLX

02356481 RELISTOR SLX

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56:92.00 MISCELLANEOUS GI DRUGS

METHYLNALTREXONE BROMIDE

20MG SOLUTION

02356503 RELISTOR SLX

96:00 PHARMACEUTICAL AIDS

96:00.00 PHARMACEUTICAL AIDS

ADMINISTRATION DIN

MISCELLANEOUS

91500004 STERILE PREPERATION FEE UNK

NUTRITIONAL SUPPLEMENT

ORAL LIQUID

00999483	BOOST DIABETIC O/L	NVC
09853154	BOOST FRUIT BEVERAGE	NES
95999970	BOOST HIPROTEIN	NES
95999963	BOOST ORIGINAL	NES
95999975	BOOST PLUS	NES
97904341	ENSURE	ABB
00801054	ENSURE HIGH PROTEIN	ABB
97904333	ENSURE PLUS	ABB
97904317	ENSURE WITH FIBRE	ABB
09854392	GLUCERNA	ABB
99101180	PEDIASURE COMPLETE GROW&GAIN	ABB
99003554	RESOURCE 2.0	NES
09853170	RESOURCE 2.0 O/L	NES

Appendix C - Palliative Care Formulary

Non-Insured Health Benefits

ADMINISTRATION DIN	5	LORAZEPAM	3
APO-FENTANYL MATRIX	1	METADOL	2
APO-NABILONE	4	METHADONE HYDROCHLORIDE (BC ONLY)	2
APP-NABILONE	4	METHADONE HYDROCHLORIDE (METADOL)	2
ARTIFICIAL SALIVA	3	METHADONE PDR (PAIN)	2
ATROPINE	1	METHADONE PDR (PALLIATIVE)	2
ATROPINE SULFATE	1	METHOTRIMEPRAZINE HYDROCHLORIDE	3
ATROPINE SULFATE	1	METHYLNALTREXONE BROMIDE	4
BOOST DIABETIC O/L	5	METOCLOPRAMIDE	4
BOOST FRUIT BEVERAGE	5	METOCLOPRAMIDE HYDROCHLORIDE	4
BOOST HIPROTEIN	5	METOCLOPRAMIDE OMEGA	4
BOOST ORIGINAL	5	MIDAZOLAM	3
BOOST PLUS	5	MIDAZOLAM	3
BUSCOPAN	1	MOISTIR	3
CO FENTANYL	1	MORPHINE HP	3
CUVPOSA	1	MORPHINE HP STERILE INFUSION	1
DIASTAT	3	MORPHINE LP	2
DIASTAT 2X10MG RECTAL PACK	3	MORPHINE LP EPIDURAL	2
DIASTAT 2X15MG RECTAL PACK	3	MORPHINE SULFATE	2
DIAZEPAM	3	MORPHINE SULFATE	2
DIAZEPAM	3	MORPHINE SULPHATE	2
DIAZEPAM (DIASTAT)	3	MORPHINE-EPD	2
DIPHENOXYLATE HYDROCHLORIDE, ATROPINE SULFATE	4	NABILONE	4
DURAGESIC	1	NOZINAN	3
ENSURE	5	NUTRITIONAL SUPPLEMENT	5
ENSURE HIGH PROTEIN	5	ONDANSETRON	4
ENSURE PLUS	5	ONDANSETRON (ON)	4
ENSURE WITH FIBRE	5	ONDANSETRON -(WITH PRESERVATIVE)	4
EXTEMPORANEOUS MIXTURE	1	ONDANSETRON BP	4
FENTANYL	1	ONDANSETRON HYDROCHLORIDE	4
FENTANYL	1	ONDANSETRON OMEGA - (PRESERVATIVE FREE SINGLE DOSE VIALS)	4
FENTANYL CITRATE	2	ONDANSETRON OMEGA -(WITH PRESERVATIVE MULTIDOSE VIAL)	4
FENTANYL CITRATE	2	ONDANSETRON W/P	4
FENTANYL CITRATE SDZ	2	PAT-FENTANYL MATRIX	1
FENTANYL STERILE INFUSION	1	PEDIASURE COMPLETE GROW&GAIN	5
FENTANYL TRANSDERMAL SYSTEM	1	PHENOBARBITAL	3
FUROSEMIDE	3	PHENOBARBITAL SODIUM	3
FUROSEMIDE	3	PHENYTOIN	3
GLUCERNA	5	PHENYTOIN SODIUM	3
GLYCOPYRROLATE	1	PMS-NABILONE	4
GLYCOPYRROLATE	1	RAN-FENTANYL MATRIX	1
GLYCOPYRROLATE MULTIDOSE	1	RANITIDINE	4
GRANISETRON	4	RANITIDINE HYDROCHLORIDE	4
GRANISETRON HYDROCHLORIDE	4	RELISTOR	4
GRANISETRON HYDROCHLORIDE	4	RESOURCE 2.0	5
HYDROMORPHONE	2	RESOURCE 2.0 O/L	5
HYDROMORPHONE HP	2	SANDOZ FENTANYL	1
HYDROMORPHONE HP FORTE	2	SCOPOLAMINE	1
HYDROMORPHONE HP STERILE INFUSION	1	SCOPOLAMINE	4
HYDROMORPHONE HYDROCHLORIDE	2	SCOPOLAMINE HYDROBROMIDE	1
HYDROMORPHONE HYDROCHLORIDE HP 10	2	STERILE PREPERATION FEE	5
HYOSCINE BUTYLBROMIDE	1	TRANSDERM-V	4
HYOSCINE BUTYLBROMIDE	1	ZOFAN	4
JAMP-ONDANSETRON	4	ZOFAN (ON)	4
KETALAR	1		
KETAMINE	1		
KETAMINE HYDROCHLORIDE	1		
LOMOTIL	4		
LORAZEPAM	3		

APPENDIX D
FORMULARY FOR ADJUNCT MEDICATIONS
USED DURING ACTIVE CANCER TREATMENT

The NIHB Program has established a new formulary to streamline access to adjunctive (non-chemotherapy) medications frequently used by clients undergoing active cancer treatment.

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Clients are automatically enrolled for a period of six months. If cancer treatment is of a longer duration, access to the formulary will be granted to align with the treatment duration. In the event that treatment duration is not known and the treatment plan extends beyond six months, access to this formulary may be extended upon request.

08:00 ANTI-INFECTIVE AGENTS

08:12.24 TETRACYCLINES

MINOCYCLINE HYDROCHLORIDE

50MG CAPSULE

02084090 APO-MINOCYCLINE	APX
02153394 MINOCYCLINE	PDL
02230735 MYLAN-MINOCYCLINE	MYL
02294419 PMS-MINOCYCLINE	PMS
02237313 SANDOZ MINOCYCLINE	SDZ
02108143 TEVA-MINOCYCLINE	TEV

100MG CAPSULE

02084104 APO-MINOCYCLINE	APX
02154366 MINOCYCLINE	PDL
02230736 MYLAN-MINOCYCLINE	MYL
02294427 PMS-MINOCYCLINE	PMS
02237314 SANDOZ MINOCYCLINE	SDZ
02108151 TEVA-MINOCYCLINE	TEV

12:00 AUTONOMIC DRUGS

12:12.08 BETA ADRENERGIC AGONISTS

**SALMETEROL XINAFOATE, FLUTICASONE
PROPRIONATE**

25MCG & 125MCG AEROSOL

02245126 ADVAIR 125	GSK
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25MCG & 250MCG AEROSOL

02245127 ADVAIR 250	GSK
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50MCG & 100MCG POWDER

02240835 ADVAIR 100 DISKUS	GSK
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50MCG & 250MCG POWDER

02240836 ADVAIR 250 DISKUS	GSK
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50MCG & 500MCG POWDER

02240837 ADVAIR 500 DISKUS	GSK
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**20:00 BLOOD FORMATION
COAGULATION AND
THROMBOSIS**

20:16.00 HEMATOPOIETIC AGENTS

DARBEPOETIN ALFA

25MCG/ML SOLUTION

02392313 ARANESP	AMG
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40MCG/ML SOLUTION

02392321 ARANESP	AMG
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60MCG/ML SOLUTION

02246348 ARANESP	AMG
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20:16.00 HEMATOPOIETIC AGENTS

DARBEPOETIN ALFA

100MCG/ML SOLUTION

02391740 ARANESP	AMG
02391759 ARANESP	AMG
02392348 ARANESP	AMG
99004917 ARANESP	AMG
99004925 ARANESP	AMG

200MCG/ML SOLUTION

02391767 ARANESP	AMG
02391775 ARANESP	AMG
02391783 ARANESP	AMG
02392356 ARANESP	AMG
99004909 ARANESP	AMG
99004933 ARANESP	AMG

500MCG/ML SOLUTION

02391791 ARANESP	AMG
02391805 ARANESP	AMG
02391821 ARANESP	AMG
02392364 ARANESP	AMG
09857185 ARANESP	AMG

EPOETIN ALFA

1,000U/0.5ML SOLUTION

02231583 EPREX	JSO
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2,000U/0.5ML SOLUTION

02231584 EPREX	JSO
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3,000U/0.3ML SOLUTION

02231585 EPREX	JSO
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4,000U/0.4ML SOLUTION

02231586 EPREX	JSO
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5000U/0.5ML SOLUTION

02243400 EPREX	JSO
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6000U/0.6ML SOLUTION

02243401 EPREX	JSO
----------------	-----

8000U/0.8ML SOLUTION

02243403 EPREX	JSO
----------------	-----

10,000/ML SOLUTION

02231587 EPREX	JSO
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20,000U/0.5ML SOLUTION

02243239 EPREX	JSO
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30,000U/0.75ML SOLUTION

02288680 EPREX	JSO
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40,000U/ML SOLUTION

02240722 EPREX	JSO
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**Appendix D - Formulary for Adjunct Medications Used
During Active Cancer Treatment**

Non-Insured Health Benefits

The NIHB Program has established a new formulary to streamline access to adjunctive (non-chemotherapy) medications frequently used by clients undergoing active cancer treatment.

Clients who are approved for oral chemotherapy drugs are given access to all of the medications and nutritional supplements in the formulary. Additionally, clients who request, and are approved, for one of the medications on the formulary for a cancer-related indication are also granted access.

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20:16.00 HEMATOPOIETIC AGENTS		28:12.92 MISCELLANEOUS ANTICONSULSANTS	
PEGFILGRASTIM		PREGABALIN	
10MG/ML SOLUTION		50MG CAPSULE	
02249790 NEULASTA	AMG	02377047 RIVA-PREGABALIN	RIV
28:00 CENTRAL NERVOUS SYSTEM AGENTS		02390825 SANDOZ PREGABALIN	SDZ
28:12.92 MISCELLANEOUS ANTICONSULSANTS		02361175 TEVA-PREGABALIN	TEV
PREGABALIN		75MG CAPSULE	
25MG CAPSULE		02402939 ACT PREGABALIN	ACG
02402912 ACT PREGABALIN	ACG	02394251 APO-PREGABALIN	APX
02394235 APO-PREGABALIN	APX	02433885 AURO-PREGABALIN	AUR
02433869 AURO-PREGABALIN	AUR	02402572 DOM-PREGABALIN	DPC
02402556 DOM-PREGABALIN	DPC	02435993 JAMP-PREGABALIN	JMP
02435977 JAMP-PREGABALIN	JMP	02268434 LYRICA	PFI
02268418 LYRICA	PFI	02417545 MAR-PREGABALIN	MAR
02417529 MAR-PREGABALIN	MAR	02424185 MINT-PREGABALIN	MIN
02423804 MINT-PREGABALIN	MIN	02479133 NRA-PREGABALIN	UNK
02479117 NRA-PREGABALIN	UNK	02359626 PMS-PREGABALIN	PMS
02359596 PMS-PREGABALIN	PMS	02396513 PREGABALIN	PDL
02396483 PREGABALIN	PDL	02403714 PREGABALIN	SIV
02403692 PREGABALIN	SIV	02405555 PREGABALIN	SAN
02405539 PREGABALIN	SAN	02476320 PREGABALIN	RIV
02476304 PREGABALIN	RIV	02392836 RAN-PREGABALIN	RBY
02392801 RAN-PREGABALIN	RBY	02377055 RIVA-PREGABALIN	RIV
02377039 RIVA-PREGABALIN	RIV	02390833 SANDOZ PREGABALIN	SDZ
02390817 SANDOZ PREGABALIN	SDZ	02361183 TEVA-PREGABALIN	TEV
02361159 TEVA-PREGABALIN	TEV	150MG CAPSULE	
50MG CAPSULE		02402955 ACT PREGABALIN	ACG
02402920 ACT PREGABALIN	ACG	02394278 APO-PREGABALIN	APX
02394243 APO-PREGABALIN	APX	02433907 AURO-PREGABALIN	AUR
02433877 AURO-PREGABALIN	AUR	02402580 DOM-PREGABALIN	DPC
02402564 DOM-PREGABALIN	DPC	02436000 JAMP-PREGABALIN	JMP
02435985 JAMP-PREGABALIN	JMP	02268450 LYRICA	PFI
02268426 LYRICA	PFI	02417561 MAR-PREGABALIN	MAR
02417537 MAR-PREGABALIN	MAR	02424207 MINT-PREGABALIN	MIN
02423812 MINT-PREGABALIN	MIN	02479168 NRA-PREGABALIN	UNK
02479125 NRA-PREGABALIN	UNK	02359634 PMS-PREGABALIN	PMS
02359618 PMS-PREGABALIN	PMS	02396521 PREGABALIN	PDL
02396505 PREGABALIN	PDL	02403722 PREGABALIN	SIV
02403706 PREGABALIN	SIV	02405563 PREGABALIN	SAN
02405547 PREGABALIN	SAN	02476347 PREGABALIN	RIV
02476312 PREGABALIN	RIV	02392844 RAN-PREGABALIN	RBY
02392828 RAN-PREGABALIN	RBY	02377063 RIVA-PREGABALIN	RIV
		02390841 SANDOZ PREGABALIN	SDZ
		02361205 TEVA-PREGABALIN	TEV

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**28:12.92 MISCELLANEOUS
ANTICONVULSANTS**

PREGABALIN

300MG CAPSULE

02402998	ACT PREGABALIN	ACG
02394294	APO-PREGABALIN	APX
02436019	JAMP-PREGABALIN	JMP
02268485	LYRICA	PFI
02359642	PMS-PREGABALIN	PMS
02396548	PREGABALIN	PDL
02403730	PREGABALIN	SIV
02405598	PREGABALIN	SAN
02476371	PREGABALIN	RIV
02392860	RAN-PREGABALIN	RBV
02377071	RIVA-PREGABALIN	RIV
02390868	SANDOZ PREGABALIN	SDZ
02361248	TEVA-PREGABALIN	TEV

**48:00 RESPIRATORY TRACT
AGENTS**

48:10.24 LEUKOTRIENE MODIFIERS

MONTELUKAST SODIUM

4MG GRANULES

02358611	SANDOZ MONTELUKAST	SDZ
02247997	SINGULAIR	FRS

10MG TABLET

02374609	APO-MONTELUKAST	APX
02401274	AURO-MONTELUKAST	AUR
02445735	BIO-MONTELUKAST	UNK
02376695	DOM-MONTELUKAST	DPC
02391422	JAMP-MONTELUKAST	JMP
02399997	MAR-MONTELUKAST	MAR
02408643	MINT-MONTELUKAST	MIN
02379333	MONTELUKAST	SAN
02379856	MONTELUKAST	PDL
02382474	MONTELUKAST	SIV
02379236	MONTELUKAST SODIUM	ACC
02373947	PMS-MONTELUKAST	PMS
02389517	RAN-MONTELUKAST	RBV
02398826	RIVA-MONTELUKAST	RIV
02328593	SANDOZ MONTELUKAST	SDZ
02238217	SINGULAIR	FRS
02355523	TEVA-MONTELUKAST	TEV

4MG TABLET (CHEWABLE)

02377608	APO-MONTELUKAST	APX
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48:10.24 LEUKOTRIENE MODIFIERS

MONTELUKAST SODIUM

4MG TABLET (CHEWABLE)

02422867	AURO-MONTELUKAST	AUR
02442353	JAMP-MONTELUKAST	JMP
02399865	MAR-MONTELUKAST	MAR
02408627	MINT-MONTELUKAST	MIN
02379317	MONTELUKAST	SAN
02379821	MONTELUKAST	PDL
02382458	MONTELUKAST	SIV
02354977	PMS-MONTELUKAST	PMS
02402793	RAN-MONTELUKAST	RBV
02330385	SANDOZ MONTELUKAST	SDZ
02243602	SINGULAIR	FRS
02355507	TEVA-MONTELUKAST	TEV

5MG TABLET (CHEWABLE)

02377616	APO-MONTELUKAST	APX
02422875	AURO-MONTELUKAST	AUR
02442361	JAMP-MONTELUKAST	JMP
02399873	MAR-MONTELUKAST	MAR
02408635	MINT-MONTELUKAST	MIN
02379325	MONTELUKAST	SAN
02379848	MONTELUKAST	PDL
02382466	MONTELUKAST	SIV
02354985	PMS-MONTELUKAST	PMS
02402807	RAN-MONTELUKAST	RBV
02330393	SANDOZ MONTELUKAST	SDZ
02238216	SINGULAIR	FRS
02355515	TEVA-MONTELUKAST	TEV

**52:00 EYE, EAR, NOSE AND
THROAT (EENT)**

**52:28.00 EENT - MOUTHWASHES AND
GARGLES**

BENZYDAMINE HYDROCHLORIDE

0.15% MOUTHWASH

02239044	APO-BENZYDAMINE	APX
02229777	PHARIXIA	PED
02239537	PMS-BENZYDAMINE	PMS

52:92.00 MISCELLANEOUS EENT DRUGS

ARTIFICIAL SALIVA

0.05MG SPRAY

02238696	MOISTIR	PMS
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Appendix D - Formulary for Adjunct Medications Used During Active Cancer Treatment

Non-Insured Health Benefits

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56:00 GASTROINTESTINAL DRUGS

56:08.00 ANTIDIARRHEA AGENTS

DIPHENOXYLATE HYDROCHLORIDE, ATROPINE SULFATE

2.5MG & 0.025MG TABLET

00036323 LOMOTIL PFI

56:22.20 5-HT3 RECEPTOR ANTAGONISTS

ONDANSETRON HYDROCHLORIDE

2MG/ML INJECTION

09857323 ONDANSETRON (ON) NOP
 02291703 ONDANSETRON W/P APX
 09857324 ZOFRAN (ON) GSK
 09857325 ZOFRAN (ON) GSK

2MG LIQUID

02271761 ONDANSETRON OMEGA - (PRESERVATIVE FREE SINGLE DOSE VIALS) OMG
 02271788 ONDANSETRON OMEGA -(WITH PRESERVATIVE MULTIDOSE VIAL) OMG

2MG SOLUTION

02420414 JAMP-ONDANSETRON JMP
 02420422 JAMP-ONDANSETRON JMP
 02462257 ONDANSETRON RAX
 02464578 ONDANSETRON RAX
 02279436 ONDANSETRON -(WITH PRESERVATIVE) SDZ
 02461420 ONDANSETRON BP AUR
 02213745 ZOFRAN NVR

2MG/ML SOLUTION

02265524 ONDANSETRON TEV
 02274418 ONDANSETRON SDZ
 02279428 ONDANSETRON SDZ
 02390019 ONDANSETRON MYL
 02390051 ONDANSETRON MYL

56:22.32 MISCELLANEOUS ANTIEMETICS

APREPITANT

80MG CAPSULE

02298791 EMEND FRS

125MG CAPSULE

02298805 EMEND FRS

125MG & 80MG CAPSULE

02298813 EMEND TRI-PACK FRS

56:22.92 MISCELLANEOUS ANTIEMETICS

NABILONE

0.25MG CAPSULE

02441497 APO-NABILONE APX
 02312263 CESAMET VAE
 02380897 PMS-NABILONE PMS
 02358077 RAN-NABILONE RBY
 02392925 TEVA-NABILONE TEV

0.5MG CAPSULE

02393581 ACT NABILONE ACG
 02441500 APO-NABILONE APX
 02256193 CESAMET VAE
 02380900 PMS-NABILONE PMS
 02358085 RAN-NABILONE RBY
 02384884 TEVA-NABILONE TEV

1MG CAPSULE

02393603 ACT NABILONE ACG
 02441519 APO-NABILONE APX
 00548375 CESAMET VAE
 02380919 PMS-NABILONE PMS
 02358093 RAN-NABILONE RBY
 02384892 TEVA-NABILONE TEV

92:00 UNCLASSIFIED THERAPEUTIC AGENTS

92:24.00 BONE RESORPTION INHIBITORS

DENOSUMAB (XGEVA)

120MG/1.7ML SOLUTION

02368153 XGEVA AMG

96:00 PHARMACEUTICAL AIDS

96:00.00 PHARMACEUTICAL AIDS

NUTRITIONAL SUPPLEMENT

ORAL LIQUID

00999483 BOOST DIABETIC O/L NVC
 09853154 BOOST FRUIT BEVERAGE NES
 95999970 BOOST HIPROTEIN NES
 95999963 BOOST ORIGINAL NES
 95999975 BOOST PLUS NES
 97904341 ENSURE ABB
 00801054 ENSURE HIGH PROTEIN ABB
 97904333 ENSURE PLUS ABB
 97904317 ENSURE WITH FIBRE ABB
 09854392 GLUCERNA ABB

**Appendix D - Formulary for Adjunct Medications Used
During Active Cancer Treatment**

Non-Insured Health Benefits

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96:00.00 PHARMACEUTICAL AIDS

NUTRITIONAL SUPPLEMENT

ORAL LIQUID

99101180 PEDIASURE COMPLETE
GROW&GAIN

ABB

Appendix D - Formulary for Adjunct Medications Used During Active Cancer Treatment

Non-Insured Health Benefits

ACT NABILONE	4	ONDANSETRON -(WITH PRESERVATIVE)	4
ACT PREGABALIN	2	ONDANSETRON BP	4
ADVAIR 100 DISKUS	1	ONDANSETRON HYDROCHLORIDE	4
ADVAIR 125	1	ONDANSETRON OMEGA - (PRESERVATIVE FREE SINGLE DOSE VIALS)	4
ADVAIR 250	1	ONDANSETRON OMEGA -(WITH PRESERVATIVE MULTIDOSE VIAL)	4
ADVAIR 250 DISKUS	1	ONDANSETRON W/P	4
ADVAIR 500 DISKUS	1	PEDIASURE COMPLETE GROW&GAIN	5
APO-BENZYDAMINE	3	PEGFILGRASTIM	2
APO-MINOCYCLINE	1	PHARIXIA	3
APO-MONTELUKAST	3	PMS-BENZYDAMINE	3
APO-NABILONE	4	PMS-MINOCYCLINE	1
APO-PREGABALIN	2	PMS-MONTELUKAST	3
APREPITANT	4	PMS-NABILONE	4
ARANESP	1	PMS-PREGABALIN	2
ARTIFICIAL SALIVA	3	PREGABALIN	2
AURO-MONTELUKAST	3	PREGABALIN	2
AURO-PREGABALIN	2	RAN-MONTELUKAST	3
BENZYDAMINE HYDROCHLORIDE	3	RAN-NABILONE	4
BIO-MONTELUKAST	3	RAN-PREGABALIN	2
BOOST DIABETIC O/L	4	RIVA-MONTELUKAST	3
BOOST FRUIT BEVERAGE	4	RIVA-PREGABALIN	2
BOOST HIPROTEIN	4	SALMETEROL XINAFOATE, FLUTICASONE PROPIONATE	1
BOOST ORIGINAL	4	SANDOZ MINOCYCLINE	1
BOOST PLUS	4	SANDOZ MONTELUKAST	3
CESAMET	4	SANDOZ PREGABALIN	2
DARBEPOETIN ALFA	1	SINGULAIR	3
DENOSUMAB (XGEVA)	4	TEVA-MINOCYCLINE	1
DIPHENOXYLATE HYDROCHLORIDE, ATROPINE SULFATE	4	TEVA-MONTELUKAST	3
DOM-MONTELUKAST	3	TEVA-NABILONE	4
DOM-PREGABALIN	2	TEVA-PREGABALIN	2
EMEND	4	XGEVA	4
EMEND TRI-PACK	4	ZOFRAN	4
ENSURE	4	ZOFRAN (ON)	4
ENSURE HIGH PROTEIN	4		
ENSURE PLUS	4		
ENSURE WITH FIBRE	4		
EPOETIN ALFA	1		
EPREX	1		
GLUCERNA	4		
JAMP-MONTELUKAST	3		
JAMP-ONDANSETRON	4		
JAMP-PREGABALIN	2		
LOMOTIL	4		
LYRICA	2		
MAR-MONTELUKAST	3		
MAR-PREGABALIN	2		
MINOCYCLINE	1		
MINOCYCLINE HYDROCHLORIDE	1		
MINT-MONTELUKAST	3		
MINT-PREGABALIN	2		
MOISTIR	3		
MONTELUKAST	3		
MONTELUKAST SODIUM	3		
MONTELUKAST SODIUM	3		
MYLAN-MINOCYCLINE	1		
NABILONE	4		
NEULASTA	2		
NRA-PREGABALIN	2		
NUTRITIONAL SUPPLEMENT	4		
ONDANSETRON	4		
ONDANSETRON (ON)	4		

APPENDIX E
EXTEMPORANEOUS MIXTURES

To be eligible under the NIHB Program, extemporaneous mixtures (compounds) must have at least one ingredient listed on the DBL and must not duplicate the formulation of commercially manufactured drug products. Mixtures that contain exception or limited use drugs must receive prior approval by the DEC. Mixtures that contain ingredients excluded from the Program will not be eligible for coverage.

All extemporaneous mixtures must be submitted with the corresponding pseudo-DIN to be reimbursed appropriately. Pharmaceutical powders of eligible ingredients may be used in lieu of tablets/capsules. These powders must be billed at AAC and must not exceed the maximum allowable AAC which is based on the price of the DIN of the comparable listed tablet or capsule.

Back Order Items and Compounding:

Providers who are preparing a compound to replace a commercially available product which is on back-order do not require a PA. The claim must be submitted using the corresponding miscellaneous pseudo-DIN. Providers are required to maintain documentation demonstrating that the commercially available product was on back-order at the time of dispense.

Compounds with Diclofenac:

Compounds with diclofenac as an ingredient require a PA and will be reviewed on a case-by-case basis

COMPOUNDED EXTERNAL LOTION

99502001 MENTHOL & CAMPHOR IN CORTICOSTEROID
LOTION

99502002 MISCELLANEOUS COMPOUNDED EXTERNAL
LOTION

COMPOUNDED EXTERNAL POWDER

99504000 MISCELLANEOUS COMPOUNDED EXTERNAL
POWDER

COMPOUNDED EYE/EAR DROP

99507000 MISCELLANEOUS COMPOUNDED EYE/EAR
DROP

99507001 ANTIFUNGAL DROPS

99507002 ANTIBIOTIC DROPS

99507003 ANTIVIRAL DROPS

COMPOUNDED INJECTION OR INFUSION

99506000 CEFAZOLIN STERILE INFUSION

99506001 CEFTRIAZONE STERILE INFUSION

99506003 PENICILLIN G STERILE INFUSION

99506004 GENTAMYCIN STERILE INFUSION

99506005 AMPICILLIN STERILE INFUSION

99506008 CLINDAMYCIN STERILE INFUSION

99506015 IRON SUCROSE STERILE INFUSION

99506021 MISCELLANEOUS COMPOUNDED
INJECTION/INFUSION

COMPOUNDED INTERNAL POWDER

99505000 MISCELLANEOUS COMPOUNDED INTERNAL
POWDER

99505003 PHENAZOPYRIDINE COMPOUNDED

99505004 BACKORDER INTERNAL POWDER

COMPOUNDED INTERNAL USE LIQUID

99503000 HYDROCHLOROTHIAZIDE ORAL LIQUID

99503001 SPIRONOLACTONE ORAL LIQUID

99503002 OMEPRAZOLE ORAL LIQUID

99503003 AMLODIPINE ORAL LIQUID

99503004 NITRO-FURANTOIN ORAL LIQUID

99503005 DOMPERIDONE ORAL LIQUID

99503006 TRANEXAMIC DENTAL MOUTHWASH

99503007 DEXAMETHASONE ORAL LIQUID

99503008 PREDNISONE ORAL LIQUID

99503009 ALDACTAZIDE ORAL LIQUID

99503010 LANSOPRAZOLE ORAL LIQUID

99503011 BACLOFEN ORAL LIQUID

99503012 METRONIDAZOLE ORAL LIQUID

99503013 ENALAPRIL ORAL LIQUID

99503014 PROPRANOLOL ORAL LIQUID

99503015 METOPROLOL ORAL LIQUID

99503016 AMIODARONE ORAL LIQUID

99503017 TRIMETHOPRIM ORAL LIQUID

99503018 ALLOPURINOL ORAL LIQUID

99503019 AZATHIOPRINE ORAL LIQUID

99503020 BENZODIAZEPINE ORAL LIQUID

99503021 CLONIDINE ORAL LIQUID

99503022 RIFAMPIN ORAL LIQUID

99503023 SOTALOL ORAL LIQUID

99503024 UROSODIOL ORAL LIQUID

99503025 MISCELLANEOUS COMPOUNDED INTERNAL
LIQUID

99503026 LEVETIRACETAM ORAL LIQUID

99503027 TOPIRAMATE ORAL LIQUID

99503028 ANTACID AND LIDOCAINE ORAL LIQUID

99503029 MAGIC MOUTHWASH

99503031 ISONIAZID ORAL LIQUID

99503032 OPIOID COMPOUNDED

99503033 MISC LIMITED USE COMPOUND INTERNAL

To be eligible under the NIHB Program, extemporaneous mixtures (compounds) must have at least one ingredient listed on the DBL and must not duplicate the formulation of commercially manufactured drug products. Mixtures that contain exception or limited use drugs must receive prior approval by the DEC. Mixtures that contain ingredients excluded from the Program will not be eligible for coverage.

All extemporaneous mixtures must be submitted with the corresponding pseudo-DIN to be reimbursed appropriately. Pharmaceutical powders of eligible ingredients may be used in lieu of tablets/capsules. These powders must be billed at AAC and must not exceed the maximum allowable AAC which is based on the price of the DIN of the comparable listed tablet or capsule.

Back Order Items and Compounding:

Providers who are preparing a compound to replace a commercially available product which is on back-order do not require a PA. The claim must be submitted using the corresponding miscellaneous pseudo-DIN. Providers are required to maintain documentation demonstrating that the commercially available product was on back-order at the time of dispense.

Compounds with Diclofenac:

Compounds with diclofenac as an ingredient require a PA and will be reviewed on a case-by-case basis

COMPOUNDED SUPPOSITORY

99508000 MISCELLANEOUS COMPOUNDED
SUPPOSITORY

COMPOUNDED TOPICAL CREAM

99500000 HYDROCORTISONE POWDER AND
CLOTRIMAZOLE CREAM
99500001 STEROID AND ANTIFUNGAL CREAM
99500002 MENTHOL &/OR CAMPHOR IN STEROID
99500003 SALICYLIC ACID IN CORTICOSTEROID CREAM
99500004 MISCELLANEOUS COMPOUNDED TOPICAL
CREAM
99500006 SULFUR IN NON-MEDICATED CREAM
99500008 MOMETASONE CREAM
99500009 LCD IN NON-MEDICATED CREAM
99500010 LCD IN CORTICOSTEROID CREAM
99504001 MISC LIMITED USE EXTERNAL COMPOUND
MIXTURE

COMPOUNDED TOPICAL OINTMENT

99501000 LCD IN CORTICOSTEROID OINTMENT
99501001 SALICYLIC ACID IN NON-MEDICATED
OINTMENT
99501002 SULFUR IN NON-MEDICATED OINTMENT
99501003 CALCIUM CHANNEL BLOCKER IN OINTMENT
99501004 MISCELLANEOUS COMPOUNDED TOPICAL
OINTMENT
99501005 LCD IN NON-MEDICATED OINTMENT
99501006 ALL PURPOSE NIPPLE OINTMENT
99501008 DILTIAZEM IN OINTMENT
99502000 CLINDAMYCIN IN DILUSOL OR DUONALC

GENDER AFFIRMING THERAPY

00915311 GENDER AFFIRMING TOPICAL HORMONES
00915312 GENDER AFFIRMING HORMONES

STERILE EXTEMPORANEOUS MIXTURE

00915000 STERILE EXTEMPORANEOUS MIXTURE (QC)

APPENDIX F
LIST OF DRUG MANUFACTURERS

Appendix F - List of Drug Manufacturers
Non-Insured Health Benefits

MFR	Manufacturer Name	MFR	Manufacturer Name
AAP	AA PHARMA INCORPORATED	DPI	DOMREX PHARMA INCORPORATED
ABB	ABBOTT LABORATORIES LIMITED	DPT	DERMTEK PHARMA INCORPORATED
ABV	ABBVIE CORPORATION	DUI	DUCHESNAY INCORPORATED
ACC	ACCORD HEALTHCARE INCORPORATED	EIS	EISAI LIMITED
ACG	ACTAVIS GROUP PTC EHF	ELN	ELAN PHARMACEUTICALS INCORPORATED
ACP	ACCEL PHARMA INCORPORATED	ERF	ERFA CANADA INCORPORATED
ADA	ADAMS LABS LIMITED	ETH	ETHYPHARM INCORPORATED
ADD	AVEVA DRUG DELIVERY SYSTEMS INCORPORATED	EUR	EURO-PHARM INTERNATIONAL CANADA INCORPORATED
ALC	ALCON CANADA INCORPORATED	FEI	FERRING INCORPORATED
ALK	ALK ABELLO A/S	FKD	FRESENIUS KABI CANADA LIMITED
ALL	ALLERGAN INCORPORATED	FMC	FRESENIUS MEDICAL CARE NORTH AMERICA
ALV	ALVEDA PHARMACEUTICALS INCORPORATED	FRS	MERCK FROSST CANADA LIMITED
AMD	AMDIPHARM LIMITED	GAC	GALDERMA CANADA INCORPORATED
AMG	AMGEN CANADA INCORPORATED	GEE	GENZYME CANADA INCORPORATED
ANG	ANGITA PHARMA INCORPORATED	GIL	GILEAD SCIENCES INCORPORATED
APC	APTALIS PHARMA CANADA ULC	GLK	GLENMARK PHARMACEUTICALS CANADA INCORPORATED
APL	AUROBINDO PHARMA LIMITED	GMP	GENERIC MEDICAL PARTNERS INCORPORATED
APU	ATNAHS PHARMA UK LIMITED	GPB	G POHL-BOSKAMP GMBH & CO KG
APX	APOTEX INCORPORATED	GSK	GLAXOSMITHKLINE INCORPORATED
ARA	ARA PHARMACEUTICALS INCORPORATED	HIL	HILL DERMACEUTICALS INCORPORATED
ARI	ARIAD PHARMACEUTICALS INCORPORATED	HJS	H.J. SUTTON INDUSTRIES LIMITED
ASP	ASPEN PHARMA TRADING LIMITED	HLR	HOFFMAN-LAROCHE LIMITED
AST	ASTELLAS PHARMA CANADA INCORPORATED	HLS	HLS THERAPEUTICS INC
ATL	LABORATORIE ATLAS INCORPORATED	HOD	NIPRO DIAGNOSTICS CANADA LIMITED
ATO	ATON PHARMA INCORPORATED, A DIVISION OF VALEANT PHARMACEUTICALS NORTH AMERICA LLC	HOS	HOSPIRA HEALTHCARE CORPORATION
AUC	AUTO CONTROL	HYD	HYDRATION PHARMACEUTICALS CANADA INCORPORATED
AUP	AURIUM PHARMA INCORPORATED	ICN	ICN CANADA LIMITED
AUR	AURO PHARMA INCORPORATED	IDE	INTERNATIONAL DERMATOLOGICALS INCORPORATED
AZC	ASTRAZENECA CANADA INCORPORATED	IND	INDIVIOR UK LIMITED
BAX	BAXTER CORPORATION	INS	INSIGHT PHARMACEUTICALS LLC
BAY	BAYER INCORPORATED, HEALTHCARE/DIAGNOSTICS	IPS	IPSEN LIMITED
BEN	BENCARD ALLERGY LABORATORIES	JAC	JACOBUS PHARMACEUTICAL COMPANY INCORPORATED
BEX	BERLEX CANADA INCORPORATED	JAJ	JOHNSON & JOHNSON
BGP	BGP PHARMA ULC	JAM	C.E. JAMIESON COMPANY LIMITED
BIO	BIONICHE PHARMA (CANADA) LIMITED	JMP	JAMP PHARMA CORPORATION
BMI	BIOMED 2002 INCORPORATED	JNO	JANSSEN-ORTHO INCORPORATED
BMS	BRISTOL-MYERS SQUIBB CANADA	JSO	JANSSEN INCORPORATED
BOE	BOEHRINGER INGELHEIM (CANADA) LIMITED	JUB	JUBILANT HOLLISTERSTIER LLC
BSH	BAUSCH & LOMB CANADA INCORPORATED	KAL	KALEO INCORPORATED
BSY	BIOSYENT PHARMA INCORPORATED	KIM	MCNEIL CONSUMER HEALTHCARE, A DIVISION OF JOHNSON & JOHNSON INCORPORATED
BTD	WEB PACK INTERNATIONAL INCORPORATED	KLA	PATRIOT A DIVISION OF JANSSEN INCORPORATED
BTU	BRAINTREE LABORATORIES INCORPORATED	LAL	LABORATOIRE LALCO INCORPORATED
CHE	CHEPLAPHARM ARZNEIMITTEL GMBH GERMANY	LAP	LABORATOIRE HRA PHARMA
CHU	CHURCH & DWIGHT CANADA CORP	LEO	LEO PHARMA INCORPORATED
CIP	CIPHER PHARMACEUTICALS INCORPORATED	LIL	ELI LILLY CANADA INCORPORATED
CLC	COLUMBIA LABORATORIES CANADA INCORPORATED	LIP	LINEPHARMA INTERNATIONAL LIMITED
COV	COVIDIEN CANADA	LUD	LUNDBECK CANADA INCORPORATED
DCM	D & C MOBILITY	LUK	LUNDBECK LLC
DDP	THE D DROPS COMPANY INCORPORATED	LUP	LUPIN PHARMA CANADA LIMITED
DOR	DORMER LABORATORIES INCORPORATED	MAC	MACDONALD'S PRESCRIPTION LAB LIMITED
DPC	DOMINION PHARMACAL		

Appendix F - List of Drug Manufacturers
Non-Insured Health Benefits

MFR	Manufacturer Name	MFR	Manufacturer Name
MAK	3M CANADA COMPANY	REC	DR REDDYS LABORATORIES INCORPORATED
MAN	MANTRA PHARMA INCORPORATED	RGL	RECRO GAINESVILLE LLC
MAR	MARCAN PHARMACEUTICALS INCORPORATED	RIV	LABORATORIE RIVA INCORPORATED
MAT	MALLINCKRODT CANADA ULC	RLI	RED LEAF MEDICAL INCORPORATED
MAY	MAYNE PHARMA (CANADA) INCORPORATED	ROD	ROCHE DIAGNOSTICS
MCA	MCARTHUR MEDICAL SALES INCORPORATED	RPH	RATIOPHARM INCORPORATED
MCL	MCNEIL CONSUMER PRODUCTS COMPANY	SAC	SANOFI-AVENTIS CANADA
MDF	MEDICAL FUTURES INCORPORATED	SAN	SANIS HEALTH INCORPORATED
MDS	MEDISCA PHARMACEUTIQUE INCORPORATED	SDZ	SANDOZ CANADA INCORPORATED
MDT	MEDTRONIC OF CANADA LIMITED	SEA	SEARCHLIGHT PHARMA INCORPORATED
MEC	MEDI+SURE CANADA INCORPORATED	SEV	SERVIER CANADA INCORPORATED
MEZ	MERZ PHARMACEUTICALS GMBH	SFA	HTL STREFA
MIN	MINT PHARMACEUTICALS INCORPORATED	SHI	SHIRE CANADA INCORPORATED
MJO	MEAD JOHNSON CANADA INCORPORATED	SIV	SIVEM PHARMACEUTICALS ULC
MPD	MEDICAL PLASTIC DEVICES INCORPORATED	SKY	LIFESCAN INCORPORATED, PART OF THE JOHNSON & JOHNSON
MSF	MEDISAFE DISTRIBUTION INCORPORATED	SLX	SALIX PHARMACEUTICALS INCORPORATED
MTC	MEDTECH PRODUCTS INCORPORATED	SMW	SMITH & NEPHEW CANADA
MYL	MYLAN PHARMACEUTICALS ULC	SNE	SMITH & NEPHEW INCORPORATED
NCA	NOVA DIABETES CARE	SPC	SUNOVION PHARMACEUTICALS CANADA INCORPORATED
NEB	NEOBOURNE PHARMA LP	SPH	SOLVAY PHARMA INCORPORATED
NES	NESTLÉ CANADA INCORPORATED	SPT	SEPTA PHARMACEUTICALS INCORPORATED
NOO	NOVO NORDISK CANADA INCORPORATED	SRO	EMD SERONO A DIVISION OF EMD INCORPORATED CANADA
NOP	NOVOPHARM LIMITED	STE	STERIMAX INCORPORATED
NPH	NATCO PHARMA CANADA INCORPORATED	STG	LABORATOIRES STERIGEN INCORPORATED
NUR	NUTRICORP INTERNATIONAL	SUN	SUN PHARMA GLOBAL FZE
NVC	NOVARTIS CONSUMER HEALTH CANADA INCORPORATED	SUS	SUNSTAR AMERICAS INCORPORATED
NVR	NOVARTIS PHARMACEUTICALS CANADA INCORPORATED	SWS	SWISS HERBAL REMEDIES LIMITED
OBT	COBALT PHARMACEUTICALS COMPANY	TAK	TAKEDA PHARMACEUTICALS AMERICA INCORPORATED
ODN	ODAN LABORATORIES LIMITED	TAN	TANTA PHARMACEUTICALS INCORPORATED
OMG	OMEGA LABORATORIES LIMITED	TAR	TARO PHARMACEUTICALS INCORPORATED
OPU	OPUS PHARMA	TEL	TELIGHT OU
ORM	ORIMED PHARMA INCORPORATED	TEV	TEVA CANADA LIMITED
OTS	OTSUKA PHARMACEUTICAL CORPORATION LIMITED	TIL	TILLOTTS PHARMA GMBH
PAL	PALADIN LABS INCORPORATED	TIP	H & P INDUSTRIES / THE TRIAD-GROUP
PDI	PROFESSIONAL DISPOSABLES INTERNATIONAL LIMITED	TLI	LABORATOIRES TRIANON INCORPORATED
PDL	PRO DOC LIMITED	TPT	TAROPHARMA, A DIVISION OF TARO PHARMACEUTICALS INCORPORATED
PED	PENDOPHARM INCORPORATED	TRE	TREMBLAY HARRISON INCORPORATED
PEI	PEDIAPHARM INCORPORATED	TRI	TRIANON LABORATORIES INCORPORATED
PER	PERRIGO INTERNATIONAL	TRM	ACERUS PHARMACEUTICALS CORPORATION
PFD	PROFESSIONAL DISPOSABLES	TRU	TRUDELL MEDICAL INTERNATIONAL
PFI	PFIZER CANADA INCORPORATED	TSN	TRIMEDIC SUPPLY NETWORK LIMITED
PFR	PURDUE PHARMA	TYC	KENDALL HEALTHCARE
PGI	PROCTOR & GAMBLE INCORPORATED	UCB	UBC PHARMA INCORPORATED
PHA	PHARMAPAR INCORPORATED	UMI	ULTIMED, INCORPORATED
PMS	PHARMASCIENCE INCORPORATED	UNK	
PMT	PHARMETICS INCORPORATED	VAE	VALEANT CANADA LIMITED
PPH	PAR PHARMACEUTICAL COMPANIES	VAN	VANC PHARMACEUTICALS INCORPORATED
PPI	PRESTIGE PHARMA INCORPORATED	VII	VIIV HEALTHCARE ULC
RAX	STERIMAX INC	VTH	VITA HEALTH PRODUCTS INCORPORATED
RBP	RB PHARMACEUTICALS LIMITED	WAM	WAMPOLE INCORPORATED
RBW	R.W. PACKAGING LIMITED	WEP	WE PHARMACEUTICALS
RBX	RANBAXY PHARMACEUTICALS CANADA INCORPORATED	WNP	WN PHARMACEUTICALS LIMITED

Appendix F - List of Drug Manufacturers**Non-Insured Health Benefits**

MFR	Manufacturer Name	MFR	Manufacturer Name
WPC	WELLSPRING PHARMACEUTICAL CANADA CORPORATION		
XED	XEDITON PHARMACEUTICALS INCORPORATED		
XEN	XENEX LABS INCORPORATED		

APPENDIX G
LIST OF EXCLUSIONS

Appendix G - Exclusions

Non-Insured Health Benefits

Certain drug products are not within the scope of the program. These products will not be reimbursed as benefits under the NIHB Program:

Anti-obesity drugs;
Household products (regular soaps and shampoos);
Cosmetics;
Alternative therapies, including glucosamine and evening primrose oil;
Megavitamins;
Drugs with investigational/experimental status;
Vaccinations for travel indications;
Hair growth stimulants;
Fertility agents and impotence drugs;
Selected over-the-counter products;
Opioid containing cough preparations;
Dalmane®, Somnol® and generics (flurazepam);
Darvon® and 642® (propoxyphene);
Fiorinal®, Fiorinal® C ¼, Fiorinal® C ½ and generics (Butalbital containing analgesics with and without codeine);
Librium®, Solium®, Medilium® and generics (chlordiazepoxide);
Stadol TM NS and generics (butorphanol tartrate nasal spray);
Tranxene® and generics (clorazepate); and
Imovane® and generics (zopiclone).

The following drugs are excluded from the NIHB Program as recommended by the Common Drug Review (CDR) and the NIHB Drugs and Therapeutics Advisory Committee (DTAC) because published evidence does not support the clinical value or cost of the drug relative to existing therapies, or there is insufficient clinical evidence to support coverage.

Of Note: The Appeal Process and the Emergency Supply Policy will not apply for the following drug products.

DIN	MFR	Brand Name	Strength and Format
02248722	ALL	ACULAR LS	0.4% SOLUTION
02259052	AST	AMEVIVE	15MG/ML POWDER FOR SOLUTION
02247916	BAY	CIPRO XL	500MG TABLET (EXTENDED RELEASE)
02251787	BAY	CIPRO XL	1,000MG TABLET (EXTENDED RELEASE)
02248417	FEI	GYNAZOLE	2% CREAM
01926799	SAC	IMOVANE	7.5MG TABLET
02216167	SAC	IMOVANE	5MG TABLET
02244521	AZC	NEXIUM	20MG TABLET (DELAYED RELEASE)
02244522	AZC	NEXIUM	40MG TABLET (DELAYED RELEASE)
02241804	TAK	PANTOLOC	20MG TABLET (ENTERIC COATED)
02248503	GSK	PAXIL	12.5MG TABLET (EXTENDED RELEASE)
02248504	GSK	PAXIL	25MG TABLET (EXTENDED RELEASE)
02229437	FMC	PHOSLO	667MG TABLET
02256290	PFI	RELPAX	20MG TABLET
02256304	PFI	RELPAX	40MG TABLET

**APPENDIX H
NEW LISTINGS**

Appendix H - New Listings

Non-Insured Health Benefits

The following items have been added to the NIHB Drug Benefit List since it was last published. Some of these items may have specific criteria for use. Please consult Appendix A for criteria.

DIN	MFR	Brand Name	Strength and Dosage Form	Date Added
02369230	ANG	AG-AMLODIPINE	5MG TABLET	2019-03-22
02369249	ANG	AG-AMLODIPINE	10MG TABLET	2019-03-22
02374617	ANG	AG-CITALOPRAM	10MG TABLET	2019-03-22
02339390	ANG	AG-CITALOPRAM	20MG TABLET	2019-03-22
02339404	ANG	AG-CITALOPRAM	40MG TABLET	2019-03-22
02475308	ANG	AG-DULOXETINE	30MG CAPSULE (DELAYED RELEASE)	2019-03-22
02475316	ANG	AG-DULOXETINE	60MG CAPSULE (DELAYED RELEASE)	2019-03-22
02475898	ANG	AG-EZETIMIBE	10MG TABLET	2019-03-22
02474409	ANG	AG-IRBESARTAN	150MG TABLET	2019-03-22
02474395	ANG	AG-IRBESARTAN	75MG TABLET	2019-03-22
02474417	ANG	AG-IRBESARTAN	300MG TABLET	2019-03-22
02441217	ANG	AG-LOSARTAN	100MG TABLET	2019-03-22
02441209	ANG	AG-LOSARTAN	50MG TABLET	2019-03-22
02441195	ANG	AG-LOSARTAN	25MG TABLET	2019-03-22
02475537	ANG	AG-PAROXETINE	10MG TABLET	2019-03-22
02475545	ANG	AG-PAROXETINE	20MG TABLET	2019-03-22
02475553	ANG	AG-PAROXETINE	30MG TABLET	2019-03-22
02458136	HLR	ALECSARO	150MG CAPSULE	2019-02-19
95900000	ABB	ALIMENTUM	ORAL LIQUID	2019-03-06
95900001	UNK	ALIMENTUM	ORAL LIQUID	2019-03-06
95900047	ABB	ALIMENTUM PDR (400G)	POWDER	2019-03-06
02375990	APX	ALLERGY REMEDY	10MG TABLET	2019-02-01
02419556	ACC	AMLODIPINE BESYLATE	2.5MG TABLET	2019-02-28
02419572	ACC	AMLODIPINE BESYLATE	10MG TABLET	2019-02-28
02419564	ACC	AMLODIPINE BESYLATE	5MG TABLET	2019-02-28
02477130	APX	APO-ACYCLOVIR	5% OINTMENT	2019-04-18
02475375	APX	APO-AMBRISANTAN	5MG TABLET	2019-01-18
02475383	APX	APO-AMBRISANTAN	10MG TABLET	2019-01-01
02468050	APX	APO-GEFITINIB	250MG TABLET	2019-02-27
02415305	APX	APO-TRAVOPROST-TIMOP	0.5% & 0.004% SOLUTION	2019-04-30
09991656	UNK	AQUA-E/ML	20U/ML LIQUID	2019-01-14
02445816	AUR	AURO-CANDESARTAN	32MG TABLET	2019-04-18
02445808	AUR	AURO-CANDESARTAN	16MG TABLET	2019-04-18
02468689	AUR	AURO-CEFIXIME	100MG POWDER FOR SUSPENSION	2019-01-18
02473305	AUR	AURO-QUINAPRIL HCTZ	20MG & 12.5MG TABLET	2019-05-02
02473321	AUR	AURO-QUINAPRIL HCTZ	20MG & 25MG TABLET	2019-05-02
02473291	AUR	AURO-QUINAPRIL HCTZ	10MG & 12.5MG TABLET	2019-05-02
80088060	BIO	BIO-CAL DR FORTE	500-400MGU TABLET	2019-04-18
02452936	UCB	BRIVLERA	10MG TABLET	2019-02-11
02452960	UCB	BRIVLERA	75MG TABLET	2019-02-11
02452979	UCB	BRIVLERA	100MG TABLET	2019-02-11
02452944	UCB	BRIVLERA	25MG TABLET	2019-02-11
02452952	UCB	BRIVLERA	50MG TABLET	2019-02-11
02150859	BAY	CANESTEN COMFORTAB 1	500MG VAGINAL TABLET	2019-01-01
80028872	PAL	CANTHACUR 07	1% LIQUID	2019-03-13
80023975	DOR	CANTHARONE 07	1%(W/V) LIQUID	2019-03-13
02378590	SAC	CAPRELSA	300MG TABLET	2019-05-01
02378582	SAC	CAPRELSA	100MG TABLET	2019-05-01
02329840	GIL	CAYSTON	75MG POWDER FOR SOLUTION	2019-04-01
02246362	PFI	CENTRUM	0MG TABLET	2019-05-07
80021452	PFI	CENTRUM	0MG TABLET	2019-05-07

Appendix H - New Listings

Non-Insured Health Benefits

The following items have been added to the NIHB Drug Benefit List since it was last published. Some of these items may have specific criteria for use. Please consult Appendix A for criteria.

DIN	MFR	Brand Name	Strength and Dosage Form	Date Added
80024482	PFI	CENTRUM FOR WOMEN	0MG TABLET	2019-05-07
02462842	EUR	CHLORHEXIDINE	0.12% MOUTHWASH	2019-05-01
02244849	RAX	COLISTIMETHATE FOR U.S.P	150MG POWDER FOR SOLUTION	2019-04-01
00476420	ERF	COLY-MYCIN M PARENTERAL	150MG POWDER FOR SOLUTION	2019-04-01
99101215	NVC	COSENTYX (STYLO)	150MG/ML INJECTION	2019-05-21
09857548	NVC	COSENTYX PEN (ON)	150MG/ML INJECTION	2019-05-21
02476134	TEL	DICLOFENAC SODIUM	1.5% SOLUTION	2019-05-01
02387174	FRS	DIFICID	200MG TABLET	2019-04-17
00392731	SDZ	DIMENHYDRINATE	10MG LIQUID	2019-05-17
95900003	MJO	ENFAMIL A+	ORAL LIQUID	2019-03-06
95900164	MJO	ENFAMIL A+ 663G PDR	POWDER	2019-06-01
95900007	MJO	ENFAMIL A+ READY TO FEED	ORAL LIQUID	2019-06-01
95900152	MJO	ENFAMIL ENFACARE A+	ORAL LIQUID	2019-06-01
95900009	MJO	ENFAMIL ENFACARE A+ 363G PDR	POWDER	2019-03-06
02457393	LEO	ENSTILAR	50MCG & 0.5MG AEROSOL, FOAM	2018-10-15
02396971	CIP	EPURIS	10MG CAPSULE	2019-03-13
02396998	CIP	EPURIS	20MG CAPSULE	2019-03-13
02397005	CIP	EPURIS	30MG CAPSULE	2019-03-13
02397013	CIP	EPURIS	40MG CAPSULE	2019-03-13
02248676	AZC	IRESSA	250MG TABLET	2019-02-27
02474794	JMP	JAMP ENALAPRIL	5MG TABLET	2019-05-02
02474816	JMP	JAMP ENALAPRIL	20MG TABLET	2019-05-02
02474786	JMP	JAMP ENALAPRIL	2.5MG TABLET	2019-05-02
02474808	JMP	JAMP ENALAPRIL	10MG TABLET	2019-05-02
02477025	JMP	JAMP PERINDOPRIL	8MG TABLET	2019-05-14
02477017	JMP	JAMP PERINDOPRIL	4MG TABLET	2019-05-14
02412470	JMP	JAMP-PYRANTEL PAMOATE	50MG SUSPENSION	2019-03-01
02463717	JMP	JAMP-RANITIDINE	150MG TABLET	2019-02-28
02463725	JMP	JAMP-RANITIDINE	300MG TABLET	2019-02-28
02479087	JMP	JAMP-TENOFOVIR	300MG TABLET	2019-05-02
02472392	JMP	JAMP-URSODIOL	250MG TABLET	2019-01-18
02472406	JMP	JAMP-URSODIOL	500MG TABLET	2019-01-18
02475774	VII	JULUCA	50MG & 25MG TABLET	2019-04-16
02400871	RAX	KETOTIFEN	0.25MG SOLUTION	2019-03-28
02474565	APX	LAPELGA	6MG SOLUTION	2019-04-24
02458640	SEV	LIXIANA	15MG TABLET	2019-04-03
02458659	SEV	LIXIANA	30MG TABLET	2019-04-03
02458667	SEV	LIXIANA	60MG TABLET	2019-04-03
02481227	MAR	MAR-DAPSONE	100MG TABLET	2019-01-18
02444771	IDE	MAR-ENALAPRIL	10MG TABLET	2019-05-28
02459450	MAR	MAR-ENALAPRIL	2.5MG TABLET	2019-05-28
02444798	IDE	MAR-ENALAPRIL	20MG TABLET	2019-05-28
02459469	MAR	MAR-ENALAPRIL	5MG TABLET	2019-05-28
02443716	MAR	MAR-RANITIDINE	300MG TABLET	2019-02-28
02467550	ABV	MAVIRET	100MG & 40MG TABLET	2019-04-01
02473216	MAN	M-DULOXETINE	60MG CAPSULE (DELAYED RELEASE)	2019-01-18
02473208	MAN	M-DULOXETINE	30MG CAPSULE (DELAYED RELEASE)	2019-01-18
80013276	PGI	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE	3G POWDER	2019-02-01
80013287	PGI	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE SUGAR FREE	3G POWDER	2019-02-01

Appendix H - New Listings

Non-Insured Health Benefits

The following items have been added to the NIHB Drug Benefit List since it was last published. Some of these items may have specific criteria for use. Please consult Appendix A for criteria.

DIN	MFR	Brand Name	Strength and Dosage Form	Date Added
80015505	PGI	METAMUCIL SMOOTH TEXTURE UNFLAVOURED UNSWEETENED	3G POWDER	2019-02-01
02480956	MIN	MINT-ABACAVIR	300MG TABLET	2019-05-28
02465612	MIN	MINT-BISOPROLOL	5MG TABLET	2019-04-18
02465620	MIN	MINT-BISOPROLOL	10MG TABLET	2019-04-18
02476916	MIN	MINT-CANDESARTAN	8MG TABLET	2019-02-14
02476924	MIN	MINT-CANDESARTAN	16MG TABLET	2019-02-14
02476789	MIN	MINT-PERINDOPRIL	8MG TABLET	2019-04-10
02476770	MIN	MINT-PERINDOPRIL	4MG TABLET	2019-04-10
02476762	MIN	MINT-PERINDOPRIL	2MG TABLET	2019-05-28
02476282	MAN	M-PRAVASTATIN	20MG TABLET	2019-04-18
02476290	MAN	M-PRAVASTATIN	40MG TABLET	2019-04-18
02476274	MAN	M-PRAVASTATIN	10MG TABLET	2019-04-18
02473542	MAN	M-RANITIDINE	300MG TABLET	2019-04-18
02473534	MAN	M-RANITIDINE	150MG TABLET	2019-01-18
02483912	NPH	NAT-ERLOTINIB	25MG TABLET	2019-05-02
02472511	NPH	NAT-TENOFOVIR	300MG TABLET	2019-05-02
80004919	VTH	NATURES BOUNTY PRENATAL VITAMINS	2MG TABLET	2019-05-28
95900025	UNK	NEOCATE W/ DHA & ARA 400G PDR	POWDER	2019-06-01
02476452	UNK	NRA-AMLODIPINE	2.5MG TABLET	2019-02-05
02476460	UNK	NRA-AMLODIPINE	5MG TABLET	2019-02-05
02476479	UNK	NRA-AMLODIPINE	10MG TABLET	2019-02-05
02481669	UNK	NRA-EZETIMIBE	10MG TABLET	2019-02-05
02479753	UNK	NRA-PAROXETINE	10MG TABLET	2019-02-14
02479788	UNK	NRA-PAROXETINE	30MG TABLET	2019-02-14
02479761	UNK	NRA-PAROXETINE	20MG TABLET	2019-02-14
95900026	MJO	NUTRAMIGEN A+	ORAL LIQUID	2019-06-01
95900027	MJO	NUTRAMIGEN A+ LGG 566G PDR	POWDER	2019-06-01
02467224	HLR	OCREVUS	30MG SOLUTION	2019-06-03
80045908	BAY	ONE A DAY WOMEN	2MG TABLET	2019-05-23
02479893	SIV	PERINDOPRIL ERBUMINE	8MG TABLET	2019-04-18
02479877	SIV	PERINDOPRIL ERBUMINE	2MG TABLET	2019-04-18
02479885	SIV	PERINDOPRIL ERBUMINE	4MG TABLET	2019-04-18
02413469	PMS	PMS-DICLOFENAC-MISOPROSTOL	200MCG & 50MG TABLET (DELAYED RELEASE)	2019-04-18
02413477	PMS	PMS-DICLOFENAC-MISOPROSTOL	200MCG & 75MG TABLET (DELAYED RELEASE)	2019-04-18
02476576	PMS	PMS-PROGESTERONE	100MG CAPSULE	2019-03-12
02419610	UNK	POMALYST	4MG CAPSULE	2019-05-21
02419580	UNK	POMALYST	1MG CAPSULE	2019-05-21
02419602	UNK	POMALYST	3MG CAPSULE	2019-05-21
02419599	UNK	POMALYST	2MG CAPSULE	2019-05-21
95900035	MJO	PURAMINO A+ PDR	POWDER	2019-06-01
02442302	UNK	QUINSAIR	240MG SOLUTION	2019-03-13
00903725	UNK	REFUSAL TO FILL	MISCELLANEOUS	2019-04-10
02470373	UNK	RENFLEXIS	100MG POWDER FOR SOLUTION	2019-04-17
02461757	OTS	REXULTI	0.5MG TABLET	2019-02-11
02461765	OTS	REXULTI	1MG TABLET	2019-02-11
02461749	OTS	REXULTI	0.25MG TABLET	2019-02-11
02461773	OTS	REXULTI	2MG TABLET	2019-02-11

Appendix H - New Listings

Non-Insured Health Benefits

The following items have been added to the NIHB Drug Benefit List since it was last published. Some of these items may have specific criteria for use. Please consult Appendix A for criteria.

DIN	MFR	Brand Name	Strength and Dosage Form	Date Added
02461781	OTS	REXULTI	3MG TABLET	2019-02-11
02461803	OTS	REXULTI	4MG TABLET	2019-02-11
02479397	RIV	RIVA-ARIPIRAZOLE	30MG TABLET	2019-01-18
02479389	RIV	RIVA-ARIPIRAZOLE	20MG TABLET	2019-01-18
02479362	RIV	RIVA-ARIPIRAZOLE	10MG TABLET	2019-01-18
02479346	RIV	RIVA-ARIPIRAZOLE	2MG TABLET	2019-01-18
02479370	RIV	RIVA-ARIPIRAZOLE	15MG TABLET	2019-01-18
02479354	RIV	RIVA-ARIPIRAZOLE	5MG TABLET	2019-01-18
02405970	RIV	RIVA-ATOMOXETINE	18MG CAPSULE	2019-01-24
02468476	RIV	RIVA-CLINDAMYCIN	150MG CAPSULE	2019-05-28
02482576	SDZ	SANDOZ AMOXI-CLAV	500MG & 125MG TABLET	2019-05-02
02479486	SDZ	SANDOZ FLUOXETINE	10MG CAPSULE	2019-04-10
02479494	SDZ	SANDOZ FLUOXETINE	20MG CAPSULE	2019-04-10
02473623	VAE	SILIQ	210MG SOLUTION	2019-03-19
95900036	UNK	SIMILAC ADVANCE NEOSURE 363G	POWDER	2019-03-06
80039441	PFI	STRESSTABS FOR WOMEN	10MG TABLET	2019-05-07
00951417	UNK	T : SLIM X2 CARTRIDGE (SK)	3ML NEEDLE	2019-03-01
02456222	AZC	TAGRISSO	80MG TABLET	2019-01-23
02456214	AZC	TAGRISSO	40MG TABLET	2019-01-23
02471051	TAR	TARO-DIPYRIDAMOLE/ ASA	200MG & 25MG CAPSULE (IMMEDIATE AND EXTENDED RELEASE)	2019-01-18
02482983	TAR	TARO-IMIQUIMOD PUMP	5% CREAM	2019-04-10
02453738	TEV	TEVA-FLUTICASONE	50MCG PUMP	2019-05-01
02475863	TEV	TEVA-MOMETASONE	50MCG SPRAY	2019-01-18
02389622	TEV	TEVA-TOBRAMYCIN	60MG SOLUTION	2019-04-01
02365154	NVR	TOBI PODHALER	28MG CAPSULE	2019-04-01
02443368	SDZ	TOBRAMYCIN INHALATION	300MG SOLUTION	2019-04-01
02458039	ABV	VENCLEXTA	10MG TABLET	2019-06-03
02458047	ABV	VENCLEXTA	50MG TABLET	2019-06-03
02458055	ABV	VENCLEXTA	100MG TABLET	2019-06-03
02458063	ABV	VENCLEXTA	100MG TABLET	2019-06-03
02411849	PAL	VEREGEN	10% OINTMENT	2019-03-13

ALPHABETICAL INDEX OF DRUG PRODUCTS

Non-Insured Health Benefits

24 HOUR ALLERGY REMEDY	1	ACH-ANASTROZOLE	16	ACT PRAMIPEXOLE	96
3TC	11	ACH-BICALUTAMIDE	16	ACT PREGABALIN	75
AA-AMILZIDE	105	ACH-CANDESARTAN	56	ACT QUETIAPINE	86
AA-CLOZAPINE	83	ACH-CAPECITABINE	17	ACT RALOXIFENE	129
AA-TRIMEBUTINE	28	ACH-ESCITALOPRAM	79	ACT RAMIPRIL	55
ABACAVIR SUFLATE, LAMIVUDINE	10	ACH-EZETIMIBE	39	ACT RANITIDINE	118
ABACAVIR SULFATE	10	ACH-FINASTERIDE	152	ACT REPAGLINIDE	131
ABACAVIR SULFATE, LAMIVUDINE	10	ACH-FLUOXETINE	79	ACT RIZATRIPTAN	94
ABACAVIR SULFATE, LAMIVUDINE, DOLUTEGRAVIR SODIUM	10	ACH-LETROZOLE	20	ACT RIZATRIPTAN ODT	94
ABACAVIR SULFATE, LAMIVUDINE, ZIDOVUDINE	10	ACH-MYCOPHENOLATE	158	ACT ROPINIROLE	96
ABATACEPT	155	ACH-PIOGLITAZONE	133	ACT ROSUVASTATIN	42
ABENOL	70	ACH-TELMISARTAN HCTZ	59	ACT SUMATRIPTAN	94
ABILIFY	82	ACITRETIN	142	ACT TEMOZOLOMIDE	24
ABILIFY MAINTENA	83	ACLASTA	155	ACT TERBINAFINE	9
ABIRATERONE ACETATE	16	ACLIDINIUM BROMIDE	27	ACT VALSARTAN	59
ABOBOTULINUMTOXINA	158	ACLIDINIUM BROMIDE, FORMOTEROL FUMARATE DIHYDRATE	29	ACT VENLAFAXINE XR	82
ACAMPROSATE CALCIUM	97	ACT ALENDRONATE	154	ACTEMRA	156
ACARBOSE	129	ACT AMLODIPINE	49	ACTIKERALL	143
ACCEL PIOGLITAZONE	133	ACT AMPHETAMINE XR	89	ACTONEL	154
ACCEL-CANDESARTAN	56	ACT ANASTROZOLE	16	ACULAR	112
ACCEL-CANDESARTAN/HCTZ	57	ACT ATENOLOL	47	ACUVAIL	112
ACCEL-CITALOPRAM	77	ACT AZITHROMYCIN	4	ACYCLOVIR	13
ACCEL-DONEPEZIL	26	ACT BUPRENORPHINE/NALOXONE	69	ADALAT XL	51
ACCEL-SEVELAMER	104	ACT BUPROPION XL	77	ADALIMUMAB	155
ACCEL-TOPIRAMATE	76	ACT CABERGOLINE	96	ADAPALENE	142
ACCU-CHEK ADVANTAGE	100	ACT CANDESARTAN	56	ADCIRCA	46
ACCU-CHEK AVIVA	100	ACT CELECOXIB	62	ADDERALL XR	89
ACCU-CHEK COMPACT	100	ACT CIPROFLOXACIN	6	ADEFOVIR DIPIVOXIL	13
ACCU-CHEK FASTCLIX LANCET	162	ACT CITALOPRAM	78	ADEMPAS	108
ACCU-CHEK GUIDE (ON)	100	ACT CLARITHROMYCIN XL	4	ADHESHIVE WIPES	160
ACCU-CHEK GUIDE (SK)	100	ACT CLOPIDOGREL	36	ADMINISTRATION DIN	166
ACCU-CHEK MOBILE BG	100	ACT DEXTROAMPHETAMINE SR	89	ADRENALIN	30
ACCU-CHEK MOBILE CASSETT	100	ACT DICLO-MISO	64	ADVAGRAF	158
ACCU-CHEK MULTICLIX LANCET	162	ACT DILTIAZEM CD	51	ADVAIR 100 DISKUS	30
ACCU-CHEK SOFTCLIX LANCET	162	ACT DILTIAZEM T	51	ADVAIR 125	30
ACCU-PRIL	54	ACT DORZOTIMOLOL	113	ADVAIR 250	30
ACCURETIC	55	ACT DUTASTERIDE	152	ADVAIR 250 DISKUS	30
ACCUTANE ROCHE	143	ACT ENALAPRIL	53	ADVAIR 500 DISKUS	30
ACCUTREND	100	ACT ESCITALOPRAM ODT	79	ADVIL	63
ACEBUTOLOL	47	ACT ETIDROCAL	154	ADVIL PEDIATRIC DROPS	63
ACEBUTOLOL HYDROCHLORIDE	47	ACT ETIDRONATE	154	AERIUS	1
ACENOCOUMAROL	34	ACT EXEMESTANE	19	AERIUS KIDS	1
ACET	70	ACT FAMCICLOVIR	13	AEROCHAMBER AC BOYZ	160
ACET 120	70	ACT FINASTERIDE	152	AEROCHAMBER AC GIRLZ	160
ACET 325	70	ACT FLUCONAZOLE	9	AEROCHAMBER PLUS FLOWVU LARGE	160
ACET 650	70	ACT FLUOXETINE	79	AEROCHAMBER PLUS FLOWVU MEDIUM	160
ACETAMINOPHEN	69	ACT FLUVOXAMINE	80	AEROCHAMBER PLUS FLOWVU MOUTH	160
ACETAMINOPHEN	69	ACT LATANOPROST/TIMOLOL	113	AEROCHAMBER PLUS FLOWVU	160
ACETAMINOPHEN, CAFFEINE CITRATE, CODEINE PHOSPHATE	65	ACT LEVETIRACETAM	74	AEROCHAMBER PLUS FLOWVU	160
ACETAMINOPHEN, CODEINE PHOSPHATE	65	ACT LEVOFLOXACIN	6	AEROTRACH PLUS	160
ACETAMINOPHEN, OXYCODONE HYDROCHLORIDE	65	ACT LOSARTAN	58	AFATINIB DIMALEATE	16
ACÉTAMINOPHÈNE	70	ACT LOVASTATIN	41	AFINITOR	19
ACÉTAMINOPHÈNE BLASON SHIELD	70	ACT MELOXICAM	64	AFINITOR DISPERZ	19
ACETAZOLAMIDE	113	ACT METFORMIN	129	AFLIBERCEPT	114
ACETAZOLAMIDE	113	ACT METHYLPHENIDATE ER	90	AG-AMLODIPINE	50
ACETYLSALICYLIC ACID	62	ACT MOXIFLOXACIN	110	AG-CELECOXIB	62
ACETYLSALICYLIC ACID	62	ACT NABILONE	118	AG-CITALOPRAM	77
ACETYLSALICYLIC ACID, OXYCODONE HYDROCHLORIDE	66	ACT OLANZAPINE ODT	85	AG-DULOXETINE	78
ACH-ALENDRONATE	153	ACT OLMESARTAN	59	AG-EZETIMIBE	39
		ACT OLMESARTAN HCT	59	AGGRENOLX	46
		ACT OLOPATADINE	110	AG-IRBESARTAN	57
		ACT ONDANSETRON	117	AG-LOSARTAN	58
		ACT PAROXETINE	80	AG-PAROXETINE	80
		ACT PIOGLITAZONE	133		

Non-Insured Health Benefits

AGRYLIN	36	ALYSENA 28	127	APO AMPI	5
AG-ZOLMITRIPTAN ODT	95	AMANTADINE HYDROCHLORIDE	10	APO ASA	62
AIROMIR	30	AMBRISENTAN	46	APO CARBAMAZEPINE	72
ALBALON	112	AMCINONIDE	139	APO CHLOROTHALIDONE	105
ALCOHOL PREP	161	AMERGE	93	APO DIMENHYDRINATE	117
ALCOHOL SWABS	161	AMI-HYDRO	105	APO FUROSEMIDE	104
ALCOHOL SWABS 6893 BUTTERFLY	161	AMIKACIN SULFATE	2	APO GLYBURIDE	132
ALCOHOL SWABS 6896 (150)	161	AMIKACIN SULFATE	2	APO HALOPERIDOL	84
ALCOHOL SWABS ANTISEPTIC SKIN CLEANERS	161	AMILORIDE	105	APO HYDRO	105
ALCOHOL SWABS BD REGULAR	161	AMILORIDE, HYDROCHLOROTHIAZIDE	105	APO IBUPROFEN	63
ALDACTAZIDE	105	AMIODARONE	39	APO INDOMETHACIN	64
ALDACTAZIDE ORAL LIQUID	61	AMIODARONE HYDROCHLORIDE	39	APO METHAZIDE	45
ALDACTONE	61	AMIODARONE ORAL LIQUID	39	APO METOPROLOL	48
ALDARA P	143	AMITRIPTYLINE	77	APO METOPROLOL (TYPE L)	48
ALECENSARO	16	AMITRIPTYLINE HYDROCHLORIDE	77	APO NAPROXEN	64
ALECTINIB	16	AMLODIPINE	49	APO OXAZEPAM	92
ALEMTUZUMAB	157	AMLODIPINE BESYLATE	49	APO PEN VK	5
ALENDRONATE	154	AMLODIPINE BESYLATE	49	APO PIROXICAM	65
ALENDRONATE SODIUM	153	AMLODIPINE BESYLATE, ATORVASTATIN CALCIUM	50	APO PREDNISONE	126
ALENDRONATE SODIUM, CHOLECALCIFEROL	154	AMLODIPINE BESYLATE, TELMISARTAN	50	APO PROPRANLOL	49
ALENDRONATE-70	154	AMLODIPINE ORAL LIQUID	50	APO TRIAZIDE	105
ALERTEC	90	AMOXICILLIN	5	APO-ABACAVIR	10
ALESSE 21	127	AMOXICILLIN	5	APO-ABACAVIR-LAMIVUDINE	10
ALESSE 28	127	AMOXICILLIN (SUGAR REDUCED)	5	APO-ABACAVIR-LAMIVUDINE-ZIDOVUDINE	10
ALFACALCIDOL	148	AMOXICILLIN, CLARITHROMYCIN, LANSOPRAZOLE	119	APO-ACEBUTOLOL	47
ALFUZOSIN	30	AMOXICILLIN, CLAVULANIC ACID	5	APO-ACETAMINOPHEN	70
ALFUZOSIN HYDROCHLORIDE	30	AMPHETAMINE, DEXTROAMPHETAMINE	89	APO-ACYCLOVIR	13
ALIMENTUM	166	AMPICILLIN	5	APO-ADEFOVIR	13
ALIMENTUM PDR (400G)	166	AMPICILLIN	5	APO-ALENDRONATE	153
ALIROCUMAB	44	AMPICILLIN SODIUM	5	APO-ALENDRONATE/VITAMIN D3	154
ALKERAN	21	AMPICILLIN SODIUM FOR BP	5	APO-ALFUZOSIN	30
ALL PURPOSE NIPPLE OINTMENT	150	AMPICILLIN STERILE INFUSION	5	APO-ALLOPURINOL	152
ALLEGRA 12 HOUR	1	ANAFRANIL	78	APO-ALMOTRIPTAN	93
ALLEGRA 24 HOUR	1	ANAGRELIDE HYDROCHLORIDE	36	APO-ALPRAZ	90
ALLER-AIDE	1	ANANDRON	21	APO-AMBRISENTAN	108
ALLERGENIC EXTRACT NON POLLENS	151	ANAPROX	64	APO-AMIODARONE	39
ALLERGENIC EXTRACT POLLENS	151	ANAPROX DS	65	APO-AMITRIPTYLINE	77
ALLERGY	1	ANASTROZOLE	16	APO-AMLODIPINE	50
ALLERGY ELIXIR	1	ANASTROZOLE	16	APO-AMLODIPINE-ATORVASTATIN	50
ALLERGY EXTRA STRENGTH	1	ANDROCUR	158	APO-AMOXI	5
ALLERGY RELIEF	1	ANDRODERM	126	APO-AMOXI CLAV	5
ALLERGY REMEDY	1	ANDROGEL	126	APO-AMOXI SUGAR FREE	5
ALLERJECT	30	ANETHOLE TRITHIONE	114	APO-ANASTROZOLE	16
ALLERNIX	1	ANODAN-HC	140	APO-ARIPIPIRAZOLE	82
ALLERNIX ELIXIR	1	ANORO ELLIPTA	28	APO-ASA LD	62
ALLERNIX EXTRA STRENGTH	1	ANTACID AND LIDOCAINE ORAL LIQUID	150	APO-ATENIDONE	47
ALLERNIX MULTI SYMPTOM	1	ANTIBIOTIC DROPS	150	APO-ATENOL	47
ALLERTIN	1	ANTIBIOTIC OINT	137	APO-ATOMOXETINE	97
ALLOPURINOL	152	ANTIFUNGAL DROPS	150	APO-ATORVASTATIN	40
ALLOPURINOL	152	ANTI-NAUSEANT	117	APO-AZATHIOPRINE	157
ALLOPURINOL ORAL LIQUID	152	ANTIVIRAL DROPS	150	APO-AZITHROMYCIN	4
ALMOTRIPTAN	93	ANUGESIC HC	141	APO-BACLOFEN	31
ALMOTRIPTAN MALATE	93	ANUSOL HC	140	APO-BECLOMETHASONE	111
ALOMIDE	110	APIDRA CARTRIDGE	131	APO-BENZYLAMINE	112
ALPHAGAN	112	APIDRA SOLOSTAR	131	APO-BICALUTAMIDE	17
ALPHAGAN P	112	APIDRA VIAL	131	APO-BISACODYL	115
ALPRAZOLAM	90	APIS MELLIFERA VENOM PROTEIN EXTRACT	151	APO-BISOPROLOL	47
ALPRAZOLAM	90	APIXABAN	35	APO-BOSENTAN	46
ALTACE	55	APO ACETAMINOPHEN	70	APO-BRIMONIDINE	112
ALTACE HCT	55			APO-BROMAZEPAM	91
ALVESCO	125			APO-BUSPIRONE	93
ALYSENA 21	127			APO-CABERGOLINE	96
				APOCAL	102
				APO-CANDESARTAN	56

Non-Insured Health Benefits

APO-CANDESARTAN/HCTZ	57	APO-GEFITINIB	19	APO-MOXIFLOXACIN	7
APO-CAPTO	52	APO-GEMFIBROZIL	40	APO-MYCOPHENOLATE	158
APO-CARVEDILOL	48	APO-GLICLAZIDE	132	APO-MYCOPHENOLIC ACID	158
APO-CEFACTOR	2	APO-GLICLAZIDE MR	132	APO-NALTREXONE	71
APO-CEFADROXIL	2	APO-GRANISETRON	117	APO-NAPRO-NA	64
APO-CEFPROZIL	2	APO-HALOPIRIDOL	84	APO-NAPRO-NA DS	65
APO-CEFUROXIME	3	APO-HYDRALAZINE	45	APO-NAPROXEN	64
APO-CELECOXIB	62	APO-HYDRO	105	APO-NAPROXEN EC	65
APO-CEPHALEX	3	APO-HYDROMORPHONE	67	APO-NEVIRAPINE XR	11
APO-CETIRIZINE	1	APO-HYDROXYQUINE	15	APO-NITROGLYCERIN	45
APO-CILAZAPRIL	52	APO-HYDROXYUREA	19	APO-NORTRIPTYLINE	80
APO-CILAZAPRIL/HCTZ	52	APO-IBUPROFEN	63	APO-OFLOXACIN	110
APO-CIPROFLOX	6	APO-IMATINIB	19	APO-OLANZAPINE	84
APO-CITALOPRAM	78	APO-IMIQUIMOD	143	APO-OLANZAPINE ODT	85
APO-CLARITHROMYCIN	4	APO-INDAPAMIDE	105	APO-OLMESARTAN	59
APO-CLARITHROMYCIN XL	4	APO-IPRAVENT	28	APO-OLMESARTAN/HCTZ	59
APO-CLINDAMYCIN	7	APO-IRBESARTAN	57	APO-OLOPATADINE	110
APO-CLOBAZAM	71	APO-IRBESARTAN/HCTZ	57	APO-OMEPRAZOLE	120
APO-CLONAZEPAM	71	APO-ISMN	45	APO-ONDANSETRON	117
APO-CLONIDINE	45	APO-K	103	APO-OXCARBAZEPINE	74
APO-CLOPIDOGREL	36	APO-KETOCONAZOLE	9	APO-OXYBUTYNIN	145
APO-CROMOLYN	108	APO-KETOROLAC	112	APO-OXYCODONE/ACET	65
APO-CYCLOBENZAPRINE	31	APO-LACTULOSE	102	APO-PANTOPRAZOLE	120
APO-CYCLOSPORINE	157	APO-LAMIVUDINE	11	APO-PAROXETINE	80
APO-DEXAMETHASONE	125	APO-LAMIVUDINE HBV	11	APO-PERINDOPRIL	54
APO-DICLO	63	APO-LAMIVUDINE-ZIDOVUDINE	11	APO-PERINDOPRIL-INDAPAMIDE	54
APO-DICLO SR	63	APO-LAMOTRIGINE	73	APO-PHENYTOIN SODIUM	71
APO-DICLOFENAC	63	APO-LANSOPRAZOLE	119	APO-PINAVERIUM	121
APO-DILTIAZ	52	APO-LANSOPRAZOLE-AMOXICILLIN- CLARITHROMYCIN	119	APO-PINDOL	49
APO-DILTIAZ CD	51	APO-LATANOPROST	113	APO-PIOGLITAZONE	133
APO-DIPIVEFRIN	112	APO-LATANOPROST-TIMOP	113	APO-PRAMIPEXOLE	96
APO-DIPYRIDAMOLE	46	APO-LEFLUNOMIDE	156	APO-PRAVASTATIN	41
APO-DIVALPROEX	76	APO-LETOZOLE	20	APO-PRAZO	46
APO-DOMPERIDONE	120	APO-LEVETIRACETAM	74	APO-PREGABALIN	75
APO-DONEPEZIL	26	APO-LEVOBUNOLOL	112	APO-PROCAINAMIDE	39
APO-DORZO-TIMOP	113	APO-LEVOCARB	95	APO-PROPAFENONE	39
APO-DOXAZOSIN	46	APO-LEVOFLOXACIN	6	APO-QUETIAPINE	86
APO-DOXY	7	APO-LINEZOLID	8	APO-QUETIAPINE XR	87
APO-DULOXETINE	78	APO-LISINOPRIL	53	APO-QUINAPRIL	54
APO-DUTASTERIDE	152	APO-LITHIUM CARBONATE	93	APO-QUINAPRIL/HCTZ	55
APO-EFAVIRENZ-EMTRICITABINE- TENOFVIR	11	APO-LOPERAMIDE	115	APO-RABEPRAZOLE	120
APO-EMTRICITABINE-TENOFOVIR	12	APO-LORATADINE	1	APO-RALOXIFENE	129
APO-ENALAPRIL	53	APO-LORAZEPAM	92	APO-RAMIPRIL	55
APO-ENTECAVIR	13	APO-LOSARTAN	58	APO-RAMIPRIL/HCTZ	55
APO-ERLOTINIB	18	APO-LOSARTAN/HCTZ	58	APO-RANITIDINE	118
APO-ESCITALOPRAM	79	APO-LOVASTATIN	41	APO-REPAGLINIDE	131
APO-EXEMESTANE	19	APO-MEDROXY	133	APO-RISEDRONATE	155
APO-EZETIMIBE	40	APO-MELOXICAM	64	APO-RISPERIDONE	87
APO-FAMCICLOVIR	13	APO-METFORMIN	129	APO-RIVASTIGMINE	27
APO-FAMOTIDINE	118	APO-METHOTREXATE	21	APO-RIZATRIPTAN	94
APO-FELODIPINE	51	APO-METHYLPHENIDATE	90	APO-RIZATRIPTAN RPD	94
APO-FENOFIBRATE	40	APO-METHYLPHENIDATE ER	90	APO-ROPINIROLE	96
APO-FENO-MICRO	40	APO-METHYLPHENIDATE SR	90	APO-ROSUVASTATIN	42
APO-FENO-SUPER	40	APO-METOCLOP	121	APO-SALVENT CFC FREE	30
APO-FERROUS GLUCONATE	34	APO-METOPROLOL	48	APO-SELEGILINE	97
APO-FINASTERIDE	152	APO-METOPROLOL (TYPE L)	48	APO-SERTRALINE	81
APO-FLECAINIDE	39	APO-METOPROLOL SR	48	APO-SILDENAFIL R	45
APO-FLUCONAZOLE	9	APO-METRONIDAZOLE	15	APO-SIMVASTATIN	42
APO-FLUOXETINE	79	APO-MIDODRINE	28	APO-SOLIFENACIN	145
APO-FLURBIPROFEN	63	APO-MINOCYCLINE	7	APO-SOTALOL	49
APO-FLUTICASONE	111	APO-MIRTAZAPINE	80	APO-SUCRALFATE	119
APO-FLUVOXAMINE	80	APO-MODAFINIL	90	APO-SUMATRIPTAN	94
APO-FOSINOPRIL	53	APO-MOMETASONE	111	APO-TADALAFIL PAH	46
APO-GABAPENTIN	72	APO-MONTELUKAST	107	APO-TAMOX	24

Non-Insured Health Benefits

APO-TAMSULOSIN	31	ASCORBIC ACID	147	AURO-IRBESARTAN	57
APO-TELMISARTAN	59	ASENAPINE MALEATE	83	AURO-IRBESARTAN HCT	57
APO-TELMISARTAN/HCTZ	59	ASMANEX TWISTHALER	126	AURO-LACOSAMIDE	73
APO-TEMOZOLOMIDE	24	ASPIRIN	62	AURO-LAMIVUDINE/ZIDOVUDINE	11
APO-TENOFOVIR	12	ATACAND	56	AURO-LAMOTRIGINE	73
APO-TERAZOSIN	46	ATACAND PLUS	57	AURO-LEVETIRACETAM	74
APO-TERBINAFINE	9	ATARAX	93	AURO-LISINOPRIL	53
APO-TETRABENAZINE	98	ATASOL 15	65	AURO-LOSARTAN	58
APO-THEO-LA	146	ATASOL 30	65	AURO-LOSARTAN HCT	58
APO-TIMOP	112	ATASOL FORTE	70	AURO-MELOXICAM	64
APO-TOLTERODINE	145	ATAZANAVIR SULFATE	10	AURO-METFORMIN	129
APO-TRAPIRAMATE	76	ATENOLOL	47	AURO-METRONIDAZOLE	15
APO-TRAVOPROST Z	113	ATENOLOL	47	AURO-MIRTAZAPINE	80
APO-TRAVOPROST-TIMOP	113	ATENOLOL, CHLORTHALIDONE	47	AURO-MIRTAZAPINE OD	80
APO-TRAZODONE	82	ATIVAN	92	AURO-MODAFINIL	90
APO-TRAZODONE D	82	ATIVAN SUBLINGUAL	92	AURO-MONTELUKAST	107
APO-TRIAMCINOLONE AQ	111	ATOMOXETINE	97	AURO-MOXIFLOXACIN	7
APO-VALACYCLOVIR	13	ATOMOXETINE HYDROCHLORIDE	97	AURO-NEVIRAPINE	11
APO-VALGANCICLOVIR	13	ATORVASTATIN	40	AURO-OLANZAPINE ODT	85
APO-VALPROIC	77	ATORVASTATIN CALCIUM	40	AURO-OLMESARTAN	59
APO-VALSARTAN	59	ATORVASTATIN-10	40	AURO-PANTOPRAZOLE	120
APO-VALSARTAN/HCTZ	60	ATORVASTATIN-20	40	AURO-PAROXETINE	80
APO-VARENICLINE	32	ATORVASTATIN-40	41	AURO-PERINDOPRIL	54
APO-VENLAFAXINE XR	82	ATORVASTATIN-80	41	AURO-PRAMIPEXOLE	96
APO-VERAP	52	ATOVAQUONE	15	AURO-PRAVASTATIN	41
APO-VERAP SR	52	ATRIPLA	11	AURO-PREGABALIN	75
APO-VORICONAZOLE	9	ATROPINE	112	AURO-QUETIAPINE	86
APO-WARFARIN	36	ATROPINE SULFATE	112	AURO-QUINAPRIL HCTZ	55
APO-ZIDOVUDINE	12	ATROVENT HFA	28	AURO-RAMIPRIL	55
APO-ZOLMITRIPTAN	95	AUBAGIO	153	AURO-REPAGLINIDE	131
APO-ZOLMITRIPTAN RAPID	95	AURANOFIN	123	AURO-RISEDRONATE	155
APRACLONIDINE HYDROCHLORIDE	114	AURO-ABACAVIR/LAMIVUDINE	10	AURO-RIZATRIPTAN	94
APREPITANT	118	AURO-ALENDRONATE	154	AURO-ROSUVASTATIN	42
APRI 21	127	AURO-ALFUZOSIN	31	AURO-SERTRALINE	81
APRI 28	127	AURO-AMLODIPINE	50	AURO-SIMVASTATIN	42
APTIOM	72	AURO-AMOXICILLIN	5	AURO-SOLIFENACIN	145
APTIVUS	12	AURO-ARIPIRAZOLE	82	AURO-TELMISARTAN	59
APX-OXCARBAZEPINE	74	AURO-ATORVASTATIN	40	AURO-TELMISARTAN HCTZ	59
AQUA-E	149	AURO-BETAHISTINE	98	AURO-TENOFOVIR	12
AQUA-E/ML	149	AURO-CANDESARTAN	56	AURO-TERBINAFINE	9
AQUASOL E	149	AURO-CANDESARTAN HCT	57	AURO-TOPIRAMATE	76
AQUASOL E VITAMIN E	149	AURO-CARVEDILOL	48	AURO-TRANDOLAPRIL	56
ARAVA	156	AURO-CEFIXIME	2	AURO-VALACYCLOVIR	13
ARICEPT	26	AURO-CEFPROZIL	2	AURO-VALGANCICLOVIR	13
ARIMIDEX	16	AURO-CEFUROXIME	3	AURO-VALSARTAN	59
ARIPIRAZOLE	82	AURO-CELECOXIB	62	AURO-VALSARTAN HCT	60
ARIPIRAZOLE (MAINTENA)	83	AURO-CEPHALEXIN	3	AURO-VENLAFAXINE XR	82
ARISTOCORT C	141	AURO-CIPROFLOXACIN	6	AURO-ZIPRASIDONE	88
ARISTOCORT R	141	AURO-CITALOPRAM	78	AVALIDE	57
ARNUITY ELLIPTA	111	AURO-CLINDAMYCIN	7	AVAPRO	57
AROMASIN	19	AURO-CLOPIDOGREL	36	AVELOX	7
ARTHROTEC	64	AURO-CYCLOBENZAPRINE	31	AVENTYL	80
ARTIFICIAL TEARS	114	AURO-DONEPEZIL	26	AVIANE 21	127
ASA	62	AURO-DULOXETINE	78	AVIANE 28	127
ASA DAILY LOW DOSE	62	AURO-DUTASTERIDE	152	AVODART	152
ASA EC	62	AURO-EFAVIRENZ	11	AVONEX	153
ASACOL	121	AURO-ENTECAVIR	13	AVONEX PEN	153
ASAPHEN	62	AURO-ESCITALOPRAM	79	AXERT	93
ASAPHEN EC	62	AURO-EZETIMIBE	40	AXID	118
ASATAB	62	AURO-FINASTERIDE	152	AXITINIB	16
ASATAB EC	62	AURO-FLECAINIDE	39	AZARGA	113
ASCENCIA CONTOUR	100	AURO-FLUOXETINE	79	AZATHIOPRINE	157
ASCENCIA BREEZE 2	100	AURO-GABAPENTIN	72	AZATHIOPRINE ORAL LIQUID	157
ASCORBIC ACID	147	AURO-GALANTAMINE ER	26	AZATHIOPRINE-50	157

Non-Insured Health Benefits

AZELAIC ACID	142	BD SLIP TIP 5ML SYRINGE	163	BIO K-20 POTASSIUM	103
AZILSARTAN MEDOXOMIL	56	BD SLIP TIP 60ML SYRINGE	164	BIO-AMLODIPINE	49
AZITHROMYCIN	4	BD SLIP TIP SUB Q 26G SYRINGE	164	BIO-ANASTROZOLE	16
AZITHROMYCIN	4	BD SYRINGE + NEEDLE	164	BIO-ATENOLOL	47
AZOPT	113	BD SYRINGE WITH ULTRA-FINE NEEDLE	164	BIO-CAL DR FORTE	149
AZTREONAM	3	BD TUBERCULIN 21GX1 SYRINGE	164	BIOCALCIUM	102
B-12	147	BD TUBERCULIN 25GX5/8 SYRINGE	164	BIOCALCIUMD	103
B6	147	BD TUBERCULIN 26GX3/8 SYRINGE	164	BIOCALD FORTE	102
BABY DDROPS	148	BD TUBERCULIN 27GX1/2 SYRINGE	164	BIO-CELECOXIB	62
BACIMYXIN ONGUENT	137	BD ULTRA 29G.1/2CC SYRINGE	164	BIO-CIPROFLOXACIN	6
BACITIN	137	BD ULTRA 29G.1CC SYRINGE	164	BIO-CITALOPRAM	77
BACITRACIN ZINC	137	BD ULTRAFINE 31G 5MM PEN NEEDLE	162	BIODERM	137
BACKORDER INTERNAL POWDER	150	BD ULTRAFINE 31G 8MM PEN NEEDLE	162	BIO-DOMPERIDONE	121
BACKUP PLAN ONESTEP	127	BD ULTRAFINE 33G LANCET	162	BIO-DONEPEZIL	26
BACLOFEN	31	BD ULTRA-FINE II 30GX0.5CC SYRINGE	164	BIO-FLUOXETINE	79
BACLOFEN	31	BD ULTRA-FINE III PEN NEEDLE	162	BIO-FUROSEMIDE	104
BACLOFEN ORAL LIQUID	31	BD ULTRA-FINE NANO PEN NEEDLE	163	BIO-HYDROCHLOROTHIAZIDE	105
BACTERIOSTATIC SODIUM CHLORIDE	103	BD ULTRA-FINE PEN NEEDLE 29G	163	BIO-IRBESARTAN	57
BACTERIOSTATIC WATER	105	BECLOMETHASONE DIPROPIONATE	111	BIO-LETROZOLE	20
BACTROBAN	137	BEDUZIL	147	BIO-LEVETIRACETAM	74
BANZEL	76	BENADRYL	1	BIO-LOSARTAN	58
BARACLUDE	13	BENADRYL CHILDRENS	1	BIO-MODAFINIL	90
BARRIERE	141	BENAZEPRIL	52	BIO-MONTELUKAST	107
BASAGLAR	131	BENAZEPRIL HYDROCHLORIDE	52	BIO-MOXIFLOXACIN	7
BC SHARPS CONTAINER 1.4L	163	BENZACLIN	137	BIO-OMEPRAZOLE	120
BD ALCOHOL SWABS	161	BENZAGEL	142	BIO-PANTOPRAZOLE	120
BD AUTOSHIELD DUO SAFETY PEN NEEDLE	162	BENZAGEL 5	142	BIO-PAROXETINE	80
BD AUTOSHIELD PEN NEEDLES	162	BENZAMYCIN	137	BIO-PRAVASTATIN	41
BD BLUNT 18GX1 1/2 FILTER	162	BENZODIAZEPINE ORAL LIQUID	71	BIO-QUETIAPINE	86
BD BUTTERFLY NEEDLE 21G	163	BENZOYL PEROXIDE	142	BIO-ROSUVASTATIN	42
BD LUER-LOK TIP 10ML SYRINGE	163	BENZTROPINE MESYLATE	95	BIOSENOSIDES	116
BD LUER-LOK TIP 18GX1 1/2 SYRINGE	164	BENZTROPINE OMEGA	95	BIO-SERTRALINE	81
BD LUER-LOK TIP 1ML SYRINGE	163	BENZYLAMINE HYDROCHLORIDE	112	BIO-VALACYCLOVIR	13
BD LUER-LOK TIP 20ML SYRINGE	164	BETADERM	139	BI-PEGLYTE	116
BD LUER-LOK TIP 22GX1 1/2 SYRINGE	164	BETADINE	138	BISACODYL	115
BD LUER-LOK TIP 25GX1 SYRINGE	164	BETAGAN	112	BISACODYL	115
BD LUER-LOK TIP 25GX1 1/2 SYRINGE	164	BETAHISTINE	98	BISACODYL-ODAN	115
BD LUER-LOK TIP 25GX5/8 SYRINGE	164	BETAHISTINE HYDROCHLORIDE	98	BISMUTH	115
BD LUER-LOK TIP 30ML SYRINGE	164	BETAMETHASONE DIPROPIONATE	139	BISMUTH SUBSALICYLATE	115
BD LUER-LOK TIP 3ML SYRINGE	163	BETAMETHASONE DIPROPIONATE, CLOTRIMAZOLE	137	BISMUTH SUBSALICYLATE	115
BD LUER-LOK TIP 5ML SYRINGE	163	BETAMETHASONE DIPROPIONATE, SALICYLIC ACID	139	BISOPROLOL	47
BD LUER-LOK TIP 60ML SYRINGE	164	BETAMETHASONE DIPROPIONATE, PHOSPHATE	121	BISOPROLOL FUMARATE	47
BD MICRO-FINE 0.3CC SYRINGE	163	BETAMETHASONE VALERATE	139	BLEPHAMIDE	111
BD MICRO-FINE 28GX0.5CC SYRINGE	164	BETASERON	153	BOSENTAN MONOHYDRATE	46
BD MICRO-FINE 28GX1CC SYRINGE	164	BETASERON INITIATION KIT	153	BOSULIF	17
BD NANO PRO 32GX4MM PEN NEEDLE	163	BETAXOLOL HYDROCHLORIDE	112	BOSUTINIB	17
BD POSIFLUSH SP	163	BETHANECHOL CHLORIDE	26	BOTOX	159
BD PRECISIONGLIDE 18GX1 1/2	163	BETNESOL	121	BREEZE 2 BG (ON)	100
BD PRECISIONGLIDE 18GX1 NEEDLE	163	BETOPTIC S	112	BRENZYS	155
BD PRECISIONGLIDE 25GX1 NEEDLE	162	BEZAFIBRATE	40	BREO ELLIPTA	29
BD PRECISIONGLIDE 25GX5/8	163	BEZALIP SR	40	BREVICON 0.5/35 (21-DAY PACK)	127
BD PRECISIONGLIDE 25GX7/8	163	BG STAR	100	BREVICON 0.5/35 (28-DAY PACK)	127
BD PRECISIONGLIDE 26GX1/2	163	BG STAR (ON)	100	BREVICON 1/35 (21-DAY PACK)	127
BD PRECISIONGLIDE 26GX3/8	163	BG STAR LANCET	162	BREVICON 1/35 (28-DAY PACK)	127
BD PRECISIONGLIDE 27GX1 1/4	163	BIACNA TOPICAL	142	BREXIPRAZOLE	83
BD PRECISIONGLIDE 27GX1/2	163	BIAXIN	4	BRICANYL TURBUHALER	30
BD SHARPS CONTAINER 3.1L	163	BIAXIN XL	4	BRILINTA	37
BD SHARPS CONTAINER 3L	163	BICALUTAMIDE	16	BRIMONIDINE P	112
BD SLIP TIP 10ML SYRINGE	163	BICALUTAMIDE	17	BRIMONIDINE TARTRATE	112
BD SLIP TIP 1ML SYRINGE	163	BICILLIN	5	BRINZOLAMIDE	113
BD SLIP TIP 20ML SYRINGE	164	BIMATOPROST	113	BRINZOLAMIDE, BRIMONIDINE TARTRATE	113
BD SLIP TIP 30ML SYRINGE	164			BRINZOLAMIDE, TIMOLOL MALEATE	113
BD SLIP TIP 3ML SYRINGE	163			BRIVARACETAM	72

Non-Insured Health Benefits

BRIVLERA	72	CANTHARIDIN, PODOPHYLLIN, SALICYLIC ACID	142	CHILDREN'S ADVIL	63
BRODALUMAB	142	CANTHARONE 07	151	CHILDREN'S EUOPROFEN	63
BROMAZEPAM	91	CANTHARONE PLUS	142	CHILDREN'S MOTRIN	63
BROMAZEPAM	91	CAPECITABINE	17	CHLORAMBUCIL	17
BROMOCRIPTINE	96	CAPRELSA	25	CHLORHEXIDINE	110
BROMOCRIPTINE MESYLATE	96	CAPSAICIN	143	CHLORHEXIDINE GLUCONATE	110
BUDESONIDE	111	CAPSAICIN	143	CHLOROQUINE PHOSPHATE	15
BUDESONIDE, SODIUM CHLORIDE	139	CAPSAISIN	143	CHLORPHENIRAMINE MALEATE	1
BUPRENORPHINE HYDROCHLORIDE, NALOXONE HYDROCHLORIDE	69	CAPTROPRI	52	CHLORPROMAZINE	83
BUPROPION HYDROCHLORIDE (WELLBUTRIN)	77	CARBACHOL	113	CHLORPROMAZINE HYDROCHLORIDE	83
BUPROPION HYDROCHLORIDE (ZYBAN)	77	CARBAMAZEPINE	72	CHLORTHALIDONE	105
BUPROPION SR	77	CARBAMAZEPINE	72	CHLORTHALIDONE	105
BUSCOPAN	28	CARBOCAL	102	CHLOR-TRIPOLON	1
BUSERELIN ACETATE	17	CARBOCAL D	102	CHOLECALCIFEROL	148
BUSPIRONE	93	CARBOLITH	93	CHOLEDYL	146
BUSPIRONE HYDROCHLORIDE	93	CARDIZEM CD	51	CHOLESTYRAMINE RESIN	39
BUSULFAN	17	CARNITOR	104	CHOLESTYRAMINE-ODAN	39
CABERGOLINE	96	CARTRIDGE FOR IR200	160	CHU NICOTINE ANTI SMOKING AID	32
CADUET	50	CARVEDILOL	48	CICLESONIDE	125
CAFFEINE CITRATE	90	CARVEDILOL	48	CICLOPIROX OLAMINE	137
CAFFEINE CITRATE	90	CASODEX	17	CIDOMYCIN	2
CAL500	102	CAYA CONTOURED DIAPHRAGM	99	CILAZAPRIL	52
CALCIMAR	133	CAYA DIAPHRAGM	143	CILAZAPRIL, HYDROCHLOROTHIAZIDE	52
CALCIPOTRIOL	142	CAYSTON	3	CILOXAN	110
CALCIPOTRIOL, BETAMETHASONE DIPROPIONATE	139	CEENU	21	CIMETIDINE	118
CALCITE 500 D 400	102	CEFACTOR	2	CIMETIDINE	118
CALCITE LIQUIDE D 400	102	CEFADROXIL	2	CIMZIA	155
CALCITONIN SALMON (SYNTHETIC)	133	CEFAZOLIN	2	CIPRALEX	79
CALCITRIOL	148	CEFAZOLIN SODIUM	2	CIPRO	6
CALCITRIOL-ODAN	148	CEFAZOLIN STERILE INFUSION	2	CIPRODEX	110
CALCIUM	102	CEFIXIME	2	CIPROFLOXACIN	6
CALCIUM	102	CEFPROZIL	2	CIPROFLOXACIN HYDROCHLORIDE	6
CALCIUM 500	102	CEFTAZIDIME	3	CIPROFLOXACIN HYDROCHLORIDE, DEXAMETHASONE	110
CALCIUM 500 D 400	102	CEFTAZIDIME	3	CITALOPRAM	77
CALCIUM 500 VITAMINE D1000	102	CEFTIN	3	CITALOPRAM HYDROBROMIDE	77
CALCIUM 500 VITAMINE D400	102	CEFTRIAXONE	3	CITRIC ACID, MAGNESIUM OXIDE, SODIUM PICOSULFATE	115
CALCIUM CARBONATE	102	CEFTRIAXONE SODIUM	3	CITRIC ACID, SODIUM CITRATE	102
CALCIUM CARBONATE VITAMINE D	103	CEFTRIAXONE SODIUM FOR BP	3	CITRO MAG	115
CALCIUM CHANNEL BLOCKER IN OINTMENT	150	CEFTRIAXONE STERILE INFUSION	3	CITRODAN	115
CALCIUM GLUCONATE, VIT D	102	CEFUROXIME AXETIL	3	CLARITHROMYCIN	4
CALCIUM POLYSTYRENE SULFONATE	104	CELEBREX	62	CLARITHROMYCIN	4
CALCIUM VITAMIN D LEMON FLAVOUR	102	CELECOXIB	62	CLARITIN	1
CALCIUM, VITAMIN D	102	CELECOXIB	62	CLARITIN KIDS	1
CALD 400	102	CELESTODERM V	139	CLARUS	143
CALODAN D 400	102	CELEXA	78	CLAVULIN 125 F	5
CAMPRAL	97	CELLCEPT	158	CLAVULIN 200	5
CANAGLIFLOZIN	131	CELSENTRI	11	CLAVULIN 250 F	5
CANDESARTAN	56	CENTER-AL	151	CLAVULIN 400	5
CANDESARTAN CILEXETIL	56	CENTRUM	149	CLAVULIN 500 F	5
CANDESARTAN CILEXETIL, HYDROCHLOROTHIAZIDE	57	CENTRUM DHA	149	CLAVULIN 875	5
CANDESARTAN-HCT	57	CENTRUM FOR WOMEN	149	CLEARLAX	115
CANDESARTAN-HCTZ	57	CENTRUM JUNIOR COMPLETE	149	CLICKFINE PEN NEEDLE 31G 4.5MM	162
CANESORAL	9	CENTRUM PRENATAL	149	CLICKFINE PEN NEEDLE 31G 6MM	162
CANESTEN	137	CEPHALEXIN	3	CLICKFINE PEN NEEDLE 31G 8MM	162
CANESTEN COMBI-PAK COMFORTAB 1	138	CEPHALEXIN-500	3	CLIMARA 25	128
CANESTEN COMBI-PAK COMFORTAB 3	138	CERITINIB	17	CLIMARA 50	128
CANESTEN COMFORTAB 1	138	CERTOLIZUMAB PEGOL	155	CLIMARA 75	128
CANTHACUR 07	142	CERVICAL	99	CLINDAMYCIN	7
CANTHARIDIN	142	CESAMET	118	CLINDAMYCIN HYDROCHLORIDE	7
		CETIRIZINE	1	CLINDAMYCIN IN DILUSOL OR DUONALC	137
		CETIRIZINE HYDROCHLORIDE	1	CLINDAMYCIN IV INFUSION	8
		CHAMPIX	32		
		CHAMPIX STARTER PACK	33		

Non-Insured Health Benefits

CLINDAMYCIN PALMITATE HYDROCHLORIDE	7	COMPOUND W GEL	142	CYCLOPENTOLATE HYDROCHLORIDE	112
CLINDAMYCIN PHOSPHATE	7	COMTAN	95	CYCLOPHOSPHAMIDE	18
CLINDAMYCIN PHOSPHATE, BENZOYL PEROXIDE	137	CONCERTA	90	CYCLOSPORINE	157
CLINDAMYCIN PHOSPHATE, TRETINOIN	142	CONDOM	99	CYESTRA-35	158
CLINDAMYCIN STERILE INFUSION	7	CONDOM, LATEX, LUBRICATED	99	CYKLOKAPRON	38
CLINDAMYCINE	7	CONDOM, LATEX, NON-LUBRICATED	99	CYMBALTA	79
CLINDA-T	137	CONDOM, NON-LATEX, LUBRICATED	99	CYPROTERONE	158
CLINDOXYL	137	CONDYLINE	143	CYPROTERONE ACETATE	158
CLINDOXYL ADV	137	CONJUGATED ESTROGENS	128	CYPROTERONE ACETATE, ETHINYL ESTRADIOL	158
CLOBAZAM	71	CONJUGATED ESTROGENS, MEDROXYPROGESTERONE ACETATE	128	CYTOMEL	134
CLOBETASOL PROPIONATE	139	CONTACT DETACH 90 DEGREE 6MMX60CM	160	CYTOVENE	13
CLOBETASONE BUTYRATE	140	CONTACT DETACH 90 DEGREE 8MMX60CM	160	D VI INFANTS	148
CLOMIPRAMINE HYDROCHLORIDE	78	CONTINGENCY ONE	127	D2-DOL	148
CLONAPAM	71	CONTOUR BG (ON)	100	D3-DOL	148
CLONAZEPAM	71	CONTOUR NEXT	100	DABIGATRAN ETEXILATE MESILATE	35
CLONAZEPAM	71	CONTOUR NEXT (ON)	100	DABRAFENIB	18
CLONIDINE HYDROCHLORIDE	44	CONTRACEPTIVE	99	DACLATASVIR	14
CLONIDINE ORAL LIQUID	45	CONTRACEPTIVE DEVICE	99	DAIRY DIGESTIVE	116
CLOPIDOGREL	36	CONTRAGEL GREEN	143	DAIRY AID	117
CLOPIDOGREL BISULFATE	36	COPAXONE	152	DAKLINZA	14
CLOPIXOL	88	CORTATE	140	DALACIN	137
CLOPIXOL DEPOT	88	CORTEF	125	DALACIN C	7
CLOPIXOL-ACUPHASE	88	CORTENEMA	121	DALACIN C PHOSPHATE	8
CLOTIMADERM	137	CORTIFOAM	121	DALACIN T	137
CLOTIMAZOLE	137	CORTISONE	125	DALTEPARIN SODIUM	35
CLOTIMAZOLE	138	CORTISONE ACETATE	125	DANAZOL	126
CLOXACILLIN SODIUM	5	CORTIVERA H	151	DANTRIUM	31
CLOZAPINE	83	CORTODERM	140	DANTROLENE SODIUM	31
CLOZARIL	83	COSENTYX	143	DAPAGLIFLOZIN PROPANEDIOL MONOHYDRATE	132
COAL TAR	142	COSENTYX (STYLO)	143	DAPSONE	10
COAL TAR, SALICYLIC ACID	142	COSENTYX PEN (ON)	143	DAPSONE	10
COBIMETINIB	17	COSOPT	113	DARIFENACIN HYDROBROMIDE	145
CODEINE	66	COTAZYM	117	DARUNAVIR ETHANOLATE	10
CODEINE CONTIN CR	66	COTAZYM ECS 20	117	DARUNAVIR ETHANOLATE, COBICISTAT	10
CODEINE MONOHYDRATE, CODEINE SULFATE TRIHYDRATE	66	COTAZYM ECS 8	117	DDAVP	133
CODEINE PHOSPHATE	66	COTELLIC	17	DDAVP MELT	133
CODEINE PHOSPHATE	66	COUMADIN	36	DDROPS	148
COLCHICINE	152	COVERSYL	54	DDROPS BOOSTER	148
COLCHICINE	152	COVERSYL PLUS	54	DECAXIL	148
COLESEVELAM HYDROCHLORIDE	39	COVERSYL PLUS HD	54	DEGARELIX ACETATE	129
COLESTID	39	COZAAR	58	DELATESTRYL	126
COLESTID ORANGE	39	CREON MINIMICROSPHERES 10	117	DEMULEN 30 (21 DAY PACK)	127
COLESTIPOL HYDROCHLORIDE	39	CREON MINIMICROSPHERES 25	117	DEMULEN 30 (28 DAY PACK)	127
COLISTIMETHATE FOR U.S.P	8	CREON MINIMICROSPHERES MICRO	117	DENOSUMAB (PROLIA)	154
COLISTIN	8	CRESTOR	42	DENOSUMAB (XGEVA)	154
COLLAGENASE	143	CRITIC-AID CLEAR	141	DEPAKENE	77
COLY-MYCIN M PARENTERAL	8	CRIZOTINIB	18	DEPO-MEDROL	126
COLYTE	116	CROMOLYN	110	DEPO-MEDROL WITH LIDOCAINE	126
COMBANTRIN	2	CROMOLYN SODIUM	108	DEPO-PROVERA	133
COMBIGAN	112	CROTAMITON	138	DEPO-TESTOSTERONE	126
COMBIVENT	28	CTP 30	78	DERMAFLEX HC	140
COMBIVENT RESPIMAT	28	CUPRIMINE	124	DERMA-SMOOTHIE	140
COMBIVIR	11	CYANOCOBALAMIN	147	DERMOVATE	139
COMFORT ANGLED INFSET 17MM	160	CYANOCOBALAMIN	147	DESIPRAMINE	78
COMFORT SRT ANGLED INFSET 13	160	CYCLLEN (21 DAY)	127	DESIPRAMINE HYDROCHLORIDE	78
COMPACT SPACE PLUS LARGE MASK	160	CYCLLEN (28 DAY)	127	DESLORATADINE	1
COMPACT SPACE PLUS MEDIUM MASK	160	CYCLOBENZAPRINE	31	DESLORATADINE	1
COMPACT SPACE PLUS NO MASK	160	CYCLOBENZAPRINE HYDROCHLORIDE	31	DESLORATADINE ALLERGY CONTROL	1
COMPACT SPACE PLUS SMALL MASK	160	CYCLOGYL	112	DESMOPRESSIN	133
COMPLERA	12	CYCLOMEN	126	DESMOPRESSIN ACETATE	133
		CYCLOPENTOLATE	112	DESOGESTREL, ETHINYL ESTRADIOL	127
				DESONIDE	140

Non-Insured Health Benefits

DESOXIMETASONE	140	DIMETHYL FUMARATE	98	DOM-NYSTATIN	9
DETROL	145	DIOVAN	60	DOM-OXYBUTYNIN	145
DETROL LA	145	DIOVAN-HCT	60	DOM-PANTOPRAZOLE	120
DEVICE (METHADONE)	166	DIPENTUM	121	DOM-PAROXETINE	80
DEXAMETHASONE	111	DIPHENHYDRAMINE	1	DOMPERIDONE	121
DEXAMETHASONE	111	DIPHENHYDRAMINE	1	DOMPERIDONE MALEATE	120
DEXAMETHASONE ORAL LIQUID	125	HYDROCHLORIDE		DOMPERIDONE ORAL LIQUID	121
DEXAMETHASONE PHOSPHATE	111	DIPHENIST	1	DOM-PINDOLOL	49
DEXAMETHASONE, TOBRAMYCIN	111	DIPIVEFRIN HYDROCHLORIDE	112	DOM-PIOGLITAZONE	133
DEXAMETHASONE-OMEGA	125	DIPROLENE	139	DOM-PRAMIPEXOLE	96
DEXEDRINE	89	DIPROSALIC	139	DOM-PRAVASTATIN	41
DEXEDRINE SPANSULE	89	DIPROSONE	139	DOM-PREGABALIN	75
DEXIRON	34	DIPYRIDAMOLE	46	DOM-QUETIAPINE	86
DEXTRAN 70, HYDROXYPROPYLMETHYLCELLULOSE	114	DIPYRIDAMOLE, ACETYLSALICYLIC ACID	46	DOM-RABEPRAZOLE EC	120
DEXTROAMPHETAMINE	89	DISOPYRAMIDE	39	DOM-RAMIPRIL	55
DEXTROAMPHETAMINE SULFATE	89	DIVALPROEX	76	DOM-RISEDRONATE	155
DGEL	148	DIVIGEL	128	DOM-RIZATRIPTAN RDT	94
DIABETA	132	DOLICHOVESPULA ARENARIA VENOM PROTEIN	151	DOM-ROSUVASTATIN	42
DIAMICRON	132	DOLICHOVESPULA MACULATA VENOM PROTEIN EXTRACT	151	DOM-SALBUTAMOL	30
DIAMICRON MR	132	DOLORAL 1	68	DOM-SERTRALINE	81
DIANE-35	158	DOLORAL 5	68	DOM-SIMVASTATIN	42
DIAPER RASH	142	DOLUTEGRAVIR SODIUM	11	DOM-SOTALOL	49
DIARR-EZE	115	DOLUTEGRAVIR SODIUM, RILPIVIRINE HYDROCHLORIDE	11	DOM-SUMATRIPTAN	94
DIARRHEA RELIEF	115	DOM-ALENDRONATE	154	DOM-TERAZOSIN	46
DIASTAT	91	DOM-AMIODARONE	39	DOM-TERBINAFINE	9
DIASTAT 2X10MG RECTAL PACK	91	DOM-AMLODIPINE	49	DOM-TIAPROFENIC	65
DIASTAT 2X15MG RECTAL PACK	91	DOM-ATENOLOL	47	DOM-TIMOLOL	113
DIASTIX	101	DOM-ATOMOXETINE	97	DOM-TOPIRAMATE	76
DIAZEPAM	91	DOM-ATORVASTATIN	40	DOM-TRAZODONE	82
DIAZEPAM	91	DOM-AZITHROMYCIN	4	DOM-VALACYCLOVIR	13
DIAZEPAM (DIASTAT)	91	DOM-BACLOFEN	31	DOM-VALPROIC ACID	77
DIAZOXIDE	45	DOM-BROMOCRIPTINE	96	DOM-VENLAFAXINE XR	82
DICETEL	121	DOM-CARBAMAZEPINE	72	DOM-VERAPAMIL SR	52
DICITRATE	102	DOM-CARVEDILOL	48	DOM-ZOLMITRIPTAN	95
DICLECTIN	117	DOM-CEPHALEXIN	3	DONEPEZIL	26
DICLOFENAC	63	DOM-CIPROFLOXACIN	6	DONEPEZIL HYDROCHLORIDE	26
DICLOFENAC EC	63	DOM-CITALOPRAM	78	DORZOLAMIDE HYDROCHLORIDE	113
DICLOFENAC SODIUM	63	DOM-CLARITHROMYCIN	4	DORZOLAMIDE HYDROCHLORIDE, TIMOLOL MALEATE	113
DICLOFENAC SODIUM	63	DOM-CLOPIDOGREL	37	DOSTINEX	96
DICLOFENAC SODIUM (TOPICAL)	63	DOM-CYCLOBENZAPRINE	31	DOVOBET	139
DICLOFENAC TOPICAL	63	DOM-DICLOFENAC	63	DOVONEX	142
DICLOFENAC-SR	63	DOM-DICLOFENAC SR	63	DOXAZOSIN MESYLATE	46
DIDANOSINE	10	DOM-DOMPERIDONE	121	DOXEPIN	78
DIENOGEST	133	DOM-FINASTERIDE	152	DOXEPIN HYDROCHLORIDE	78
DIFFERIN	142	DOM-FLUCONAZOLE	9	DOXYCIN	7
DIFICID	4	DOM-FLUOXETINE	79	DOXYCYCLINE	7
DIFLUCAN	9	DOM-GABAPENTIN	72	DOXYCYCLINE HYCLATE	7
DIFLUNISAL	63	DOM-GEMFIBROZIL	40	DOXYLAMINE SUCCINATE, PYRIDOXINE HYDROCHLORIDE	117
DIFLUNISAL	63	DOM-GLYBURIDE	133	DOXYTAB	7
DIGOXIN	39	DOM-IPRATROPIUM	28	DR SCHOLLS CLEAR AWAY PLANTAR WART REMOVER SYSTEM	142
DIHYDROERGOTAMINE	30	DOM-LANSOPRAZOLE	119	DR SCHOLLS CLEAR AWAY WART REMOVER SYSTEM	142
DIHYDROERGOTAMINE MESYLATE	30	DOM-LEVETIRACETAM	74	DRESSING	160
DILANTIN	71	DOM-LOPERAMIDE	115	DROPLET PEN NEEDLE 10MM 29G	162
DILANTIN INFATABS	71	DOM-LOXAPINE	84	DROPLET PEN NEEDLE 12MM 29G	162
DILAUDID	67	DOM-MEFENAMIC ACID	64	DROPLET PEN NEEDLE 4MM 32G	163
DILTIAZEM CD	51	DOM-MELOXICAM	64	DROPLET PEN NEEDLE 5MM 31G	162
DILTIAZEM HYDROCHLORIDE	51	DOM-METFORMIN	129	DROPLET PEN NEEDLE 5MM 32G	163
DILTIAZEM IN OINTMENT	150	DOM-METOPROLOL-B	48	DROPLET PEN NEEDLE 6MM 31G	162
DILTIAZEM TZ	51	DOM-METOPROLOL-L	48	DROPLET PEN NEEDLE 6MM 32G	163
DIMENHYDRINATE	117	DOM-MIRTAZAPINE	80	DROPLET PEN NEEDLE 8MM 31G	162
DIMENHYDRINATE	117	DOM-MONTELUKAST	108	DROPLET PEN NEEDLE 8MM 32G	163
DIMETHICONE	138				

Non-Insured Health Benefits

DROPLET PERSONAL LANCET 28G	162	ENSTILAR	139	ETHINYL ESTRADIOL, ETONOGESTREL	127
DROPLET PERSONAL LANCET 33G	162	ENTACAPONE	95	ETHINYL ESTRADIOL, LEVONORGESTREL	127
DRSCHOLL'S ATHLETE'S FOOT SPRAY	138	ENTECAVIR MONOHYDRATE	13	ETHINYL ESTRADIOL, NORELGESTROMIN	127
D-TABS	148	ENTOCORT	125	ETHINYL ESTRADIOL, NORETHINDRONE ACETATE	127
DUAKLIR GENUAIR	29	ENTRESTO	61	ETHINYL ESTRADIOL, NORGESTIMATE	127
DULCOLAX	115	ENTROPHEN	62	ETHOPROPAZINE HYDROCHLORIDE	95
DULOXETINE	79	ENTYVIO	158	ETHOSUXIMIDE	71
DULOXETINE DR	79	ENZALUTAMIDE	18	ETIBI	9
DULOXETINE HYDROCHLORIDE	78	EPCLUSA	15	ETIDRONATE DISODIUM	154
DUONALC	138	EPINEPHRINE	30	ETIDRONATE DISODIUM, CALCIUM CARBONATE	154
DUOTRAV PQ	113	EPINEPHRINE	30	ETOPOSIDE	18
DUTASTERIDE	152	EPIPEN	30	ETRAVIRINE	11
DUTASTERIDE	152	EPIPEN JR	30	EUGLUCON	133
DUVOID	26	EPIVAL	76	EURAX	138
DYSPORT THERAPEUTIC	158	EPLERENONE	61	EURO D	148
EDARBI	56	EPOSARTAN MESYLATE	57	EURO K	103
EDECIN	104	EPOSARTAN MESYLATE, HYDROCHLOROTHIAZIDE	57	EURO SENNA	116
EDOXABAN	35	EPURIS	143	EURO VITAMIN B1	147
EDURANT	11	EQUATE DAILY LOW-DOSE	62	EURO-ASA	62
EFAVIRENZ	11	ERDOL	148	EUROCAL	102
EFAVIRENZ, EMTRICITABINE, TENOFIVIR DISOPROXIL FUMARATE	11	ERELZI	156	EURO-D	148
EFFEXOR XR	82	ERGOCALCIFEROL	148	EUROFER	34
EFUDEX	143	ERLOTINIB HYDROCHLORIDE	18	EURO-FERROUS SULFATE	34
EGOZINC-HC	140	ERTAPENEM	3	EUROHYDROCORTISONE	140
ELAVIL	77	ERYC	4	EUTHYROX	134
ELBASVIR, GRAZOPREVIR	14	ERYTHRO BASE	4	EVEROLIMUS	19
ELECTROLYTES	103	ERYTHROMYCIN	4	EVISTA	129
ELIDEL	143	ERYTHROMYCIN	110	EVOLOCUMAB	44
ELIGARD	20	ERYTHROMYCIN STEARATE	4	EVRA	127
ELIQUIS	35	ERYTHROMYCIN, BENZOYL PEROXIDE	137	EXELON	27
ELMIRON	151	ERYTHRO-S	4	EXEMESTANE	19
ELOCOM	141	ESBRIET	107	EXLAX CHOCOLATED	116
ELTROXIN	134	ESCITALOPRAM	79	EXTAVIA	153
EMEND	118	ESCITALOPRAM OXALATE	79	EXTEMPORANEOUS MIXTURE	150
EMEND TRI-PACK	118	ESCULIN, FRAMYCETIN SULFATE, DIBUCAINE HYDROCHLORIDE, HYDROCORTISONE ACETATE	140	EXTEMPORANEOUS MIXTURE (GENDER AFFIRMING)	150
EMLA	141	ESLICARBAZEPINE ACETATE	72	EXTEMPORANEOUS MIXTURE (LU)	150
EMOCORT	140	ESTALIS	128	EXTRA STRENGTH SELSUN	138
EMOLAX	115	ESTRACE	128	EYLEA	114
EMPAGLIFLOZIN	132	ESTRADIOL	128	EZ HEALTH ORACLE	100
EMTRICITABINE, COBICISTAT, ELVITEGRAVIR, TENOFIVIR ALAFENAMIDE	11	ESTRADIOL HEMIHYDRATE	128	EZ HEALTH ORACLE LANCET	162
EMTRICITABINE, RILPIVIRINE HYDROCHLORIDE, TENOFIVIR ALAFENAMIDE	11	ESTRADIOL, NORETHINDRONE ACETATE	128	E-Z JE	163
ENABLEX	145	ESTRADOT 100	128	E-Z SPACER	160
ENALAPRIL	53	ESTRADOT 25	128	E-Z SPACER (MASK ONLY)	160
ENALAPRIL MALEATE	53	ESTRADOT 37.5	128	E-Z SPACER WITH SMALL MASK	160
ENALAPRIL MALEATE, HYDROCHLOROTHIAZIDE	53	ESTRADOT 50	128	EZETIMIBE	39
ENALAPRIL MALEATE/HCTZ	53	ESTRADOT 75	128	EZETIMIBE	40
ENALAPRIL ORAL LIQUID	53	ESTRAGYN	128	EZETROL	40
ENBREL	155	ESTRING	128	FAMCICLOVIR	13
ENBREL SURECLICK	155	ESTROGEL	128	FAMOTIDINE	118
ENEMOL SODIUM PHOSPHATE	116	ESTRONE	128	FAMOTIDINE	118
ENFAMIL A+	166	ETANERCEPT	155	FAMVIR	13
ENFAMIL A+ 663G PDR	166	ETANERCEPT (BRENZYS)	155	FC2 FEMALE CONDOMS	99
ENFAMIL A+ READY TO FEED	166	ETANERCEPT (ERELZI)	156	FEBUXOSTAT	152
ENFAMIL ENFACARE A+	166	ETHACRYNIC ACID	104	FELODIPINE	51
ENFAMIL ENFACARE A+ 363G PDR	166	ETHAMBUTOL HYDROCHLORIDE	9	FEMARA	20
ENFAMIL FERINSOL	34	ETHINYL ESTRADIOL, DESOGESTREL	127	FEMCAP	99
ENFAMIL POLYVISOL	149	ETHINYL ESTRADIOL, DROSPIRENONE	127	FENOFIBRATE	40
ENFAMIL TRIVISOL	149	ETHINYL ESTRADIOL, ETHYNODIOL DIACETATE	127		
ENOXAPARIN SODIUM	35				

Non-Insured Health Benefits

FENOMAX	40	FLUNARIZINE HYDROCHLORIDE	95	FUSIDIC ACID	110
FENO-MICRO	40	FLUOCINONIDE	140	FUSIDIC ACID, HYDROCORTISONE ACETATE	137
FENTANYL	66	FLUOROMETHOLONE	111	GABAPENTIN	72
FENTANYL	66	FLUOROURACIL	143	GABAPENTIN	72
FERAMAX POWDER WATER SOLUBLE POLYSACCHARIDE IRON COMPLEX	34	FLUOXETINE	79	GALANTAMINE	26
FERODAN	34	FLUOXETINE HYDROCHLORIDE	79	GALANTAMINE ER	26
FERODAN INFANT DROPS	34	FLUPENTHIXOL DIHYDROCHLORIDE	83	GALANTAMINE HYDROBROMIDE	26
FERRATE	34	FLUPENTIXOL DECANOATE	84	GANCICLOVIR SODIUM	13
FERRLECIT	34	FLUPHENAZINE	84	GASTROLYTE REGULAR	103
FERROUS FUMARATE	34	FLUPHENAZINE DECANOATE	84	GATIFLOXACIN	110
FERROUS FUMARATE	34	FLUPHENAZINE HYDROCHLORIDE	84	GD-AMLODIPINE	49
FERROUS GLUCONATE	34	FLURBIPROFEN	63	GD-AMLODIPINE-ATORVASTATIN	50
FERROUS GLUCONATE	34	FLUTAMIDE	19	GD-AZITHROMYCIN	4
FERROUS SULFATE	34	FLUTAMIDE	19	GD-CELECOXIB	62
FERROUS SULFATE	34	FLUTICASONE FUROATE	111	GD-DICLOFENAC/MISOPROSTOL	64
FERROUS SULFATE	34	FLUTICASONE FUROATE, VILANTEROL TRIFENATATE	29	GD-GABAPENTIN	72
FERROUS SULPHATE	34	FLUTICASONE FUROATE, VILANTEROL TRIFENATATE (ASTHMA)	29	GD-LATANOPROST	113
FESOTERODINE FUMARATE	145	FLUTICASONE FUROATE, VILANTEROL TRIFENATATE (ASTHMA)	29	GD-LATANOPROST/TIMOLOL	113
FEXOFENADINE HYDROCHLORIDE	1	FLUTICASONE PROPIONATE	111	GD-QUINAPRIL	54
FIBRISTAL	128	FLUVASTATIN SODIUM	41	GD-TRANEXAMIC ACID	38
FIDAXOMICIN	4	FLUVOXAMINE	80	GE200	100
FILGRASTIM	37	FLUVOXAMINE MALEATE	80	GE200 (ON)	100
FINACEA	142	FML	111	GEFITINIB	19
FINASTERIDE	152	FOLIC ACID	147	GEMFIBROZIL	40
FINASTERIDE	152	FOLIC ACID	147	GEN-CLOZAPINE	83
FINGERSTIX LANCET	162	FORADIL	29	GENDER AFFIRMING HORMONES	150
FINGOLIMOD (FINGOLIMOD HYDROCHLORIDE)	157	FORMOTEROL FUMARATE	29	GENDER AFFIRMING TOPICAL HORMONES	150
FIRMAGON	129	FORMOTEROL FUMARATE DIHYDRATE	29	GENTAMICIN	2
FIRST CANADIAN HEALTH LANCETS	162	FORMOTEROL FUMARATE DIHYDRATE, BUDESONIDE	29	GENTAMICIN IV	2
FIRST CANHEALTH 28G LANCET	162	FORMOTEROL FUMARATE DIHYDRATE, MOMETASONE FUROATE	29	GENTAMICIN SULFATE	2
FIRST CANHEALTH 30G LANCET	162	FORTAZ 1G	3	GENTAMYCIN STERILE INFUSION	2
FIRST CANHEALTH 33G LANCET	162	FORTAZ 2G	3	GENTEAL	112
FIRST CANHEALTH SPIRIT	100	FORTAZ 6G	3	GENVOYA	11
FLAGYL	15	FORXIGA	132	GILENYA	157
FLAGYSTATIN	137	FOSAMAX	154	GIOTRIF	16
FLAMAZINE	139	FOSAMPRENAVIR CALCIUM	11	GLATECT	152
FLAREX	111	FOSAVANCE	154	GLATIRAMER ACETATE	152
FLAVOXATE HYDROCHLORIDE	145	FOSFOMYCIN TROMETHAMINE	15	GLECAPREVIR, PIBRENTASVIR	14
FLECAINIDE ACETATE	39	FOSINOPRIL	53	GLEEVEC	19
FLEET ENEMA	116	FOSINOPRIL SODIUM	53	GLICLAZIDE	132
FLEET ENEMA PEDIATRIC	116	FOSRENOL	104	GLICLAZIDE	132
FLEXI-T +300 IUD	99	FRAGMIN	35	GLN-GABAPENTIN	73
FLEXI-T +380 IUD	99	FRAMYCETIN SULFATE, GRAMICIDIN, DEXAMETHASONE	111	GLN-TOPIRAMATE	76
FLEXI-TD	99	FRAXIPARINE	36	GLUCAGEN	133
FLINTSTONES MULTIPLE VITAMINS PLUS IRON	149	FRAXIPARINE FORTE	36	GLUCAGEN HYPOKIT	133
FLINTSTONES MULTIPLE VITAMINS WITH EXTRA C	149	FREESTYLE	100	GLUCAGON	133
FLOCTAFENINE	70	FREESTYLE (ON)	100	GLUCAGON RECOMBINANT DNA ORGIN	133
FLOCTAFENINE	70	FREESTYLE LANCET	162	GLUCOBAY	129
FLOMAX	31	FREESTYLE LITE	100	GLUCONORM	131
FLONASE ALLERGY RELIEF	111	FREESTYLE LITE (ON)	100	GLUCOPHAGE	129
FLORINEF	125	FREESTYLE PRECISION	100	GLUCOSE OXIDASE, PEROXIDASE	100
FLOVENT DISKUS	125	FREESTYLE PRECISION (ON)	100	GLYBURIDE	132
FLOVENT HFA	125	FREYA 21	127	GLYBURIDE	132
FLUANXOL	83	FREYA 28	127	GLYCERIN	115
FLUANXOL DEPOT	84	FUCIDIN	137	GLYCERIN FOR INFANTS CHILDREN	115
FLUCONAZOLE	9	FUCIDIN H	137	GLYCERINE	115
FLUDARA	19	FUCITHALMIC	110	GLYCON	129
FLUDARABINE PHOSPHATE	19	FUROSEMIDE	104	GLYCOPYRRONIUM BROMIDE	28
FLUDROCORTISONE ACETATE	125	FUROSEMIDE	105	GOLIMUMAB	156
FLUMETHASONE PIVALATE, CLIOQUINOL	111	FUSIDATE SODIUM	137	GOLYTELY	115
FLUNARIZINE	95			GOSERELIN ACETATE	129
				GRANISETRON HYDROCHLORIDE	117

Non-Insured Health Benefits

GRASTOFIL	37	HYDROCORTISONE ACETATE, ZINC SULFATE, PRAMOXINE HYDROCHLORIDE	141	INLYTA	16
GRAVOL	117	HYDROCORTISONE ACETATE-UREA	141	INNOHEP	36
GUM PAROEX	110	HYDROCORTISONE POWDER AND CLOTRIMAZOLE CREAM	150	INSET 30 INFUSION SETS	160
HABITROL	32	HYDROCORTISONE VALERATE	141	INSET 6MMX43"	161
HALOBETASOL PROPIONATE	140	HYDROMORPH CONTIN	67	INSET II 90 DEGREE 6MMX110CM	160
HALOPERIDOL	84	HYDROMORPHONE HYDROCHLORIDE	67	INSET II 90 DEGREE 6MMX60CM	160
HALOPERIDOL	84	HYDROSONE	140	INSET II 90 DEGREE 9MMX110CM	160
HALOPERIDOL DECANOATE	84	HYDROVAL	141	INSET II 90 DEGREE 9MMX60CM	160
HALOPERIDOL LA	84	HYDROXYCHLOROQUINE SULFATE	15	INSPIOLTO RESPIMAT	30
HARVONI	14	HYDROXYPROPYL CELLULOSE	114	INSPIRA CHAMBER W LARGE MASK	160
HEMANGIOL	49	HYDROXYPROPYLMETHYLCELLULOSE	112	INSPIRA CHAMBER W MEDIUM MASK	160
HEPARIN LEO	35	HYDROXYUREA	19	INSPIRA CHAMBER W MOUTHPIECE	160
HEPARIN SODIUM	35	HYDROXYZINE	93	INSPIRA CHAMBER W SMALL MASK	160
HEPARIN SODIUM	35	HYDROXYZINE HYDROCHLORIDE	93	INSPRA	61
HEPARIN SODIUM (MULTIDOSE VIAL-WITH PRESERVATIVE)	35	HYMENOPTERA VENOM PRODUCT HONEY BEE VENOM	151	INSULIN (30% NEUTRAL & 70% ISOPHANE) HUMAN BIOSYNTHETIC	130
HEPARIN SODIUM (SINGLE USE VIAL-PRESERVATIVE FREE)	35	HYMENOPTERA VENOM PRODUCT MIXED VESPID VENOM PROTEIN	151	INSULIN (40% NEUTRAL & 60% ISOPHANE) HUMAN BIOSYNTHETIC	130
HEPSERA	13	HYMENOPTERA VENOM PRODUCT WASP VENOM PROTEIN	151	INSULIN (50% NEUTRAL & 50% ISOPHANE) HUMAN BIOSYNTHETIC	130
HEPTOVIR	11	HYMENOPTERA VENOM PRODUCT YELLOW JACKET VENOM PROTEIN	152	INSULIN (ISOPHANE) HUMAN BIOSYNTHETIC	130
HI POTENCY MAGNESIUM OXIDE	115	HYMENOPTERA VENOM PRODUCTS YELLOW HORNET VENOM PROTEIN	152	INSULIN (ZINC CRYSTALLINE) HUMAN BIOSYNTHETIC (RDNA ORIGIN)	130
HONEY BEE VENOM PROTEIN EXTRACT	151	HYOSCINE BUTYLBROMIDE	28	INSULIN 31GX0.3CC	164
HP-PAC	119	HYZAAR	58	INSULIN 31GX0.5CC	164
HUMALOG	131	HYZAAR DS	59	INSULIN 31GX1CC	164
HUMALOG (CARTRIDGE)	131	IBAVYR	14	INSULIN ASPART	131
HUMALOG (KWIKPEN)	131	IBRANCE	22	INSULIN BIOSYNTHETIC HUMAN BR	131
HUMALOG 100U/ML CARTRIDGE	131	IBRUTINIB	19	INSULIN DEGLUDEC	131
HUMALOG 200U/ML KWIKPEN	131	IBUPROFEN	63	INSULIN DETEMIR	131
HUMALOG MIX 25 (CARTRIDGE)	131	IBUPROFEN	63	INSULIN GLARGINE	131
HUMALOG MIX 25 (KWIKPEN)	131	ICLUSIG	23	INSULIN GLULISINE	131
HUMALOG MIX 50 (CARTRIDGE)	131	IDELALISIB	19	INSULIN HUMAN BIOSYNTHETIC	131
HUMALOG MIX 50 (KWIKPEN)	131	ILEVRO	112	INSULIN LISPRO	131
HUMATIN	15	IMATINIB MESYLATE	19	INSULIN LISPRO, INSULIN LISPRO PROTAMINE	131
HUMIRA	155	IMBRUVICA	19	INSULIN PEN NEEDLE 31GX6MM	162
HUMULIN 30/70	130	IMDUR	45	INSULIN PEN NEEDLE 31GX8MM	162
HUMULIN 30/70 CARTRIDGE	130	IMIPRAMINE	80	INSULIN PEN NEEDLE 32GX4MM	163
HUMULIN N	130	IMIPRAMINE HYDROCHLORIDE	80	INSULIN PEN NEEDLE 32GX6MM	163
HUMULIN N (CARTRIDGE)	130	IMIQUIMOD	143	INSULIN PEN NEEDLE 32GX8MM	163
HUMULIN N (KWIKPEN)	130	IMITREX	94	INSULIN PUMP BATTERY	160
HUMULIN N 100U/ML (CARTRIDGE)	130	IMITREX DF	94	INSULIN PUMP SUPPLIES	160
HUMULIN R	130	IMITREX STAT DOSE KIT	94	INSULIN SYR W/NEEDL 0.25CC	163
HUMULIN R (KWIKPEN)	131	IMODIUM CALMING	115	INSULIN SYR W/NEEDLE 0.3CC	163
HUMULIN R 100U/ML (CARTRIDGE)	130	IMURAN	157	INSULIN SYR W/NEEDLE 0.5CC	163
HUMULIN R CARTRIDGE	131	INCOBOTULINUMTOXINA	159	INSULIN SYR W/NEEDLE 1CC	163
HYDERM	140	INCRUSE ELLIPTA	28	INSUPEN 29GX12MM NEEDLE	162
HYDRALAZINE HYDROCHLORIDE	45	INDACATEROL MALEATE	30	INSUPEN 30GX8MM NEEDLE	162
HYDRALYTE ELECTROLYTE	103	INDACATEROL MALEATE, GLYCOPYRRONIUM BROMIDE	28	INSUPEN 31GX6MM NEEDLE	162
HYDREA	19	INDAPAMIDE	105	INSUPEN 31GX8MM NEEDLE	162
HYDROCHLOROTHIAZIDE	105	INDAYO	127	INSUPEN 32GX4MM NEEDLE	163
HYDROCHLOROTHIAZIDE	105	INDERAL LA	49	INSUPEN 32GX6MM NEEDLE	163
HYDROCHLOROTHIAZIDE ORAL LIQUID	105	INDOMETHACIN	64	INSUPEN 32GX8MM NEEDLE	163
HYDROCHLOROTHIAZIDE, PINDOLOL	48	INFANT FORMULA	166	INSUPEN 33GX4MM NEEDLE	163
HYDROCHLOROTHIAZIDE, SPIRONOLACTONE	61	INFLECTRA	156	INTELENCE	11
HYDROCORTISONE	125	INFLIXIMAB (INFLECTRA)	156	INTERFERON ALFA-2B	12
(HYDROCORTISONE SODIUM SUCCINATE)		INFLIXIMAB (REMICADE)	156	INTERFERON BETA-1A	153
HYDROCORTISONE ACETATE	121	INFUFER	34	INTERFERON BETA-1B	153
HYDROCORTISONE ACETATE	140	INHIBACE	52	INTRAUTERINE DEVICE	99
HYDROCORTISONE ACETATE, UREA	140	INHIBACE PLUS	52	INTRON A	12
HYDROCORTISONE ACETATE, ZINC SULFATE	140			INVANZ	3
HYDROCORTISONE ACETATE, ZINC SULFATE MONOHYDRATE	141			INVEGA SUSTENNA	85
				INVEGA TRINZA	85

Non-Insured Health Benefits

INVIRASE	12	JAMP OLANZAPINE ODT	85	JAMPLACTASE ENZYME	116
INVOKANA	131	JAMP PERINDOPRIL	54	JAMP-LACTULOSE	102
IOPIDINE	114	JAMP POTASSIUM CHLORIDE ER	103	JAMP-LETROZOLE	20
IPECAC	117	JAMP REHYDRALYTE	103	JAMP-LEVETIRACETAM	74
IPRATROPIUM BROMIDE	28	JAMP REPAGLINIDE	131	JAMP-LISINAPRIL	53
IPRATROPIUM BROMIDE, SALBUTAMOL SULFATE	28	JAMP SENNAQUIL	116	JAMP-LOSARTAN	58
IPRAVENT	28	JAMP VITAMIN A, D AND C	149	JAMP-LOSARTAN HCTZ	58
IRBESARTAN	57	JAMP VITAMIN B12	147	JAMP-MAGNESIUM	103
IRBESARTAN	57	JAMP VITAMIN D	148	JAMP-METFORMIN	129
IRBESARTAN HCT	57	JAMP-ALENDRONATE	154	JAMP-METFORMIN BLACKBERRY	129
IRBESARTAN, HYDROCHLOROTHIAZIDE	57	JAMP-ALLOPURINOL	152	JAMP-METHOTREXATE	21
IRBESARTAN/HCTZ	57	JAMP-ALPRAZOLAM	90	JAMP-METOPROLOL-L	48
IRBESARTAN-HCTZ	57	JAMP-AMITRIPTYLINE	77	JAMP-MONTELUKAST	108
IRESSA	19	JAMP-AMLODIPINE	49	JAMP-MOXIFLOXACIN	7
IRON	34	JAMP-AMOXICILLIN	5	JAMP-MYCOPHENOLATE	158
IRON	34	JAMP-ANASTROZOLE	16	JAMP-NYSTATIN	9
IRON DEXTRAN	34	JAMP-ASA	62	JAMPOCAINE VISCOUS	141
IRON FERROUS GLUCONATE	34	JAMP-ASA EC	62	JAMP-OLANZAPINE	84
IRON SUCROSE	34	JAMP-ATENOLOL	47	JAMP-OLMESARTAN	59
IRON SUCROSE STERILE INFUSION	34	JAMP-ATORVASTATIN	40	JAMP-OLOPATADINE	110
ISDN	45	JAMP-AZITHROMYCIN	4	JAMPOLYCIN	137
ISENTRESS	11	JAMP-BACITRACINE	137	JAMP-OMEPRAZOLE DR	120
ISONIAZID	9	JAMP-BEZFIBRATE	40	JAMP-ONDANSETRON	117
ISONIAZID ORAL LIQUID	10	JAMP-BICALUTAMIDE	17	JAMP-OXCARBAZEPINE	74
ISOPROPYL ALCOHOL	138	JAMP-BISACODYL	115	JAMP-PANTOPRAZOLE	120
ISOPROPYL MYRISTATE	138	JAMP-CALCIUM + VITAMIN D	102	JAMP-PAROXETINE	80
ISOPTIN SR	52	JAMP-CALCIUM CARBONATE	102	JAMP-PIOGLITAZONE	133
ISOPTO ATROPINE	112	JAMP-CALCIUM VITAMIN D	103	JAMP-POTASSIUM CHLORIDE	103
ISOPTO CARPINE	113	JAMP-CANDESARTAN	56	JAMP-PRAVASTATIN	41
ISOPTO TEARS	114	JAMP-CARVEDILOL	48	JAMP-PREGABALIN	75
ISOSORBIDE DINITRATE	45	JAMP-CELECOXIB	62	JAMP-PYRANTEL PAMOATE	2
ISOSORBIDE-5-MONONITRATE	45	JAMP-CETIRIZINE	1	JAMP-QUETIAPINE	86
ISOTAMINE	9	JAMP-CIPROFLOXACIN	6	JAMP-RAMIPRIL	55
ISOTRETINOIN	143	JAMP-CITALOPRAM	78	JAMP-RANITIDINE	118
ITEST	100	JAMP-CLOPIDOGREL	37	JAMP-RISEDRONATE	155
ITEST SAFETY 28G LANCET	162	JAMP-COLCHICINE	152	JAMP-RISPERIDONE	87
ITEST ULTRA-THIN 33G LANCET	162	JAMP-CYANOCOBALAMIN	147	JAMP-RIZATRIPTAN	94
ITRACONAZOLE	9	JAMP-CYCLOBENZAPRINE	31	JAMP-RIZATRIPTAN IR	94
ITRACONAZOLE PDR	9	JAMP-DIMENHYDRINATE	117	JAMP-RIZATRIPTAN ODT	94
IV3000	160	JAMP-DOMPERIDONE	121	JAMP-RAPINIROLE	96
IV3000 STANDARD	161	JAMP-DONEPEZIL	26	JAMP-ROSUVASTATIN	42
IVABRADINE (IVABRADINE HYDROCHLORIDE)	39	JAMP-DULOXETINE	79	JAMP-SERTRALINE	81
IXEKIZUMAB	143	JAMP-ESCITALOPRAM	79	JAMP-SIMVASTATIN	42
JAKAVI	24	JAMP-EZETIMIBE	40	JAMP-SODIUM PHOSPHATE	116
JAMP ACETAMINOPHEN BLAZON	70	JAMP-FER	34	JAMP-SOLIFENACIN	145
JAMP CALCIUM CARBONATE VITAMIN D	102	JAMP-FERROUS FUMARATE	34	JAMP-SOTALOL	49
JAMP CALCIUM CITRATE VITAMIN D	103	JAMP-FINASTERIDE	152	JAMP-TENOFOVIR	12
JAMP CALCIUM LACTOGLUCONATE VITAMIN D	102	JAMP-FLUCONAZOLE	9	JAMP-TERBINAFINE	9
JAMP DICLOFENAC TOPICAL	63	JAMP-FLUOXETINE	79	JAMP-TIMOLOL	113
JAMP ENALAPRIL	60	JAMP-FOLIC ACID	147	JAMP-TOBRAMYCIN	2
JAMP ENTECAVIR	13	JAMP-FOSFOMYCIN	15	JAMP-TOPIRAMATE	76
JAMP FERROUS FUMARATE	34	JAMP-FOSINOPRIL	53	JAMP-URSODIOL	116
JAMP FERROUS SULFATE	34	JAMP-GABAPENTIN	72	JAMP-VALACYCLOVIR	13
JAMP FERROUS SULFATE LIQUID5	34	JAMP-HC	140	JAMP-VANCOMYCIN	8
JAMP FOLIC ACID	147	JAMP-HYDRALAZINE	45	JAMP-VITAMIN A	147
JAMP GLICLAZIDE-MR	132	JAMP-HYDROCORTISONE	140	JAMP-VITAMIN B12	147
JAMP GLYCERIN	115	JAMP-HYDROCORTISONE UREA	141	JAMP-VITAMIN D	148
JAMP K	103	JAMP-IBUPROFEN	64	JAMP-ZINC-HC	141
JAMP MAGNESIUM GLUCONATE	103	JAMP-INDAPAMIDE	105	JAMP-ZOLMITRIPTAN	95
JAMP NEVIRAPINE	11	JAMP-IRBESARTAN	57	JAMP-ZOLMITRIPTAN ODT	95
		JAMP-IRBESARTAN AND HYDROCHLOROTHIAZIDE	57	JANUMET	130
		JAMP-K 8	103	JANUMET XR	130
		JAMP-K EFFERVESCENT	103	JANUVIA	130
		JAMPKCITRATE	103	JARDIANCE	132

Non-Insured Health Benefits

J-CAL+D	102	LATANOPROST, TIMOLOL MALEATE	113	LISINOPRIL	53
JENCYCLA	128	LATUDA	84	LISINOPRIL	53
JENTADUETO	130	LAX-A-DAY	115	LISINOPRIL, HYDROCHLOROTHIAZIDE	54
JULUCA	11	LAX-A-DAY PHARMA	116	LISINOPRIL/HCTZ (TYPE Z)	54
K LYTE	103	LCD IN CORTICOSTEROID CREAM	150	LITHANE	93
K20 POTASSIUM	103	LCD IN CORTICOSTEROID OINTMENT	150	LITHIUM CARBONATE	93
KADIAN	69	LCD IN NON-MEDICATED CREAM	150	LITHIUM CITRATE	93
KALETRA	11	LCD IN NON-MEDICATED OINTMENT	150	LITHMAX	93
KAYEXALATE	104	LEDERLE LEUCOVORIN	152	LIVOSTIN	110
KCITRA 10	102	LEFLUNOMIDE	156	LIXIANA	35
KEFLEX	3	LEFLUNOMIDE	156	LOCACORTEN VIOFORM	111
KENALOG-10	126	LEMTRADA	157	LODALIS	39
KENALOG-40	126	LENALIDOMIDE	20	LODOXAMIDE TROMETHAMINE	110
KEPPRA	74	LENVATINIB	20	LOESTRIN	127
KETOCONAZOLE	9	LENVIMA	20	LOLO	127
KETODERM	138	LESCOL XL	41	LOMUSTINE	21
KETOPROFEN	64	LETROZOLE	20	LONITEN	45
KETOPROFEN	64	LETROZOLE	20	LOPERAMIDE	115
KETOPROFEN SR	64	LEUCOVORIN CALCIUM	152	LOPERAMIDE HYDROCHLORIDE	115
KETOPROFEN-E	64	LEUKERAN	17	LOPINAVIR, RITONAVIR	11
KETOROLAC TROMETHAMINE	112	LEUPROLIDE ACETATE	20	LOPRESOR	48
KETOSTIX	101	LEVATE	77	LOPRESOR SR	48
KETOTIFEN	110	LEVEMIR FLEXTOUCH	131	LOPROX	137
KETOTIFEN FUMARATE	1	LEVEMIR PENFILL	131	LORATADINE	1
K-EXIT	104	LEVETIRACETAM	74	LORATADINE	1
KIVEXA	10	LEVETIRACETAM	74	LORAZEPAM	92
KOMBOGLYZE	130	LEVETIRACETAM ORAL LIQUID	74	LORAZEPAM	92
KWELLADA-P	138	LEVOBUNOLOL HYDROCHLORIDE	112	LORAZEPAM SUBLINGUAL	92
KYLEENA	127	LEVOCABASTINE HYDROCHLORIDE	110	LOSARTAN	58
LABETALOL HYDROCHLORIDE	48	LEVOCARNITINE	104	LOSARTAN HCT	58
LACOSAMIDE	73	LEVODOPA, BENSERAZIDE	95	LOSARTAN POTASSIUM	58
LACRISERT	114	HYDROCHLORIDE		LOSARTAN POTASSIUM,	58
LACTAID	117	LEVODOPA, CARBIDOPA	95	HYDROCHLOROTHIAZIDE	
LACTAID EXTRA STRENGTH	117	LEVODOPA, CARBIDOPA,	96	LOSARTAN/HCTZ	58
LACTAID ULTRA	117	ENTACAPONE		LOSARTAN-HCTZ	58
LACTASE	116	LEVOFLOXACIN	6	LOSEC	120
LACTEEZE DROPS	116	LEVOFLOXACIN HEMIHYDRATE	6	LOTRIDERM	137
LACTOMAX	117	LEVONORGESTREL	127	LOVASTATIN	41
LACTOMAX EXTRA	117	LEVONORGESTREL INTRAUTERINE	127	LOVASTATIN	41
LACTULOSE	102	INSERT		LOVENOX	35
LACTULOSE	102	LEVONORGESTREL, ETHINYL	127	LOVENOX HP	35
LAMICTAL	73	ESTRADIOL		LOWPRIN	62
LAMISIL	9	LEVOTHYROXINE SODIUM	134	LOXAPINE HYDROCHLORIDE	84
LAMIVUDINE	11	LIBERTE UT380 SHORT IUD	99	LOXAPINE SUCCINATE	84
LAMIVUDINE, ZIDOVUDINE	11	LIBERTE UT380 STANDARD IUD	99	LOZIDE	105
LAMOTRIGINE	73	LIDEMOL	140	LUBRICANT	143
LAMOTRIGINE	73	LIDEX	140	LUBRICATING	114
LANCET	162	LIDOCAINE HYDROCHLORIDE	112	LUBRICATING NASAL MIST	114
LANCORA	39	LIDOCAINE, PRILOCAINE	141	LUCENTIS	114
LANREOTIDE ACETATE	159	LIDODAN VISCOUS	135	LUCENTIS PFS	114
LANSOPRAZOLE	119	LIFE BRAND PEN NEEDLE 31G 8MM	163	LUMIGAN RC	113
LANSOPRAZOLE	119	LINAGLIPTIN	129	LUMIGAN RC (ON)	113
LANSOPRAZOLE ODT	119	LINAGLIPTIN, METFORMIN	130	LUPIN-ESTRADIOL	128
LANSOPRAZOLE ORAL LIQUID	119	HYDROCHLORIDE		LUPRON DEPOT	129
LANSOYL	115	LINCTUS CODEINE	66	LURASIDONE HYDROCHLORIDE	84
LANSOYL SUGAR FREE	115	LINESSA 21	127	LUVOX	80
LANTHANUM CARBONATE HYDRATE	104	LINESSA 28	127	LYDERM	140
LANTUS	131	LINEZOLID	8	LYNPARZA	22
LANTUS SOLOSTAR	131	LIORESAL	31	LYRICA	75
LANVIS	24	LIOTHYRONINE SODIUM	134	LYSODREN	21
LAPELGA	37	LIPASE, AMYLASE, PROTEASE	117	M CALCIUM VITAMINE D	102
LASIX	104	LIPIDIL EZ	40	M SENNOSIDES	116
LASIX SPECIAL	105	LIPIDIL SUPRA	40	MACROBID	15
LATANOPROST	113	LIPITOR	40		
		LISDEXAMFETAMINE DIMESYLATE	89		

Non-Insured Health Benefits

MACROGOL, POTASSIUM CHLORIDE, SODIUM BICARBONATE, SODIUM CHLORIDE, SODIUM SULFATE	115	MAR-VALACYCLOVIR	13	MESTINON-SR	27
MACROGOL, PROPYLENE GLYCOL	114	MARVELON 21	127	METADOL	68
MAGIC MOUTHWASH	150	MARVELON 28	127	METADOL-D	67
MAGLUCATE	103	MAR-ZOLMITRIPTAN	95	METAMUCIL FIBRE THERAPY ORIGINAL TEXTURE UNFLAVOURED	116
MAGNESIUM	103	M-ASA	62	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE	151
MAGNESIUM	103	MATERNA	149	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE ORANGE FLAVOUR	116
MAGNESIUM CITRATE	115	M-ATORVASTATIN	40	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE SUGAR FREE	116
MAGNESIUM COMPLEX	103	MATULANE	23	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE UNFLAVOURED	151
MAGNESIUM GLUCOHEPTONATE	103	MAVIK	56	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE SUGAR FREE	116
MAGNESIUM GLUCONATE	103	MAVIRET	14	METAMUCIL FIBRE THERAPY SMOOTH TEXTURE UNFLAVOURED	151
MAGNESIUM HYDROXIDE	115	MAXALT	94	METAMUCIL SMOOTH TEXTURE UNFLAVOURED UNSWEETENED	129
MAGNESIUM OXIDE	115	MAXALT RPD	94	METFORMIN	129
MAGNESIUM OXIDE	115	MAXIDEX	111	METFORMIN HYDROCHLORIDE	129
MAGNESIUM-ODAN	103	MAXIMUM STRENGTH ACID REDUCER	119	METFORMIN HYDROCHLORIDE, DAPAGLIFLOZIN	132
MAGNIFIER	162	MAZEPINE	72	METFORMIN HYDROCHLORIDE, EMPAGLIFLOZIN	132
MAJEPTIL	88	M-B1	147	METHADONE HYDROCHLORIDE	67
M-AMLODIPINE	49	M-B12	147	METHADONE HYDROCHLORIDE (BC ONLY)	67
MANERIX	80	M-B6	147	METHADONE HYDROCHLORIDE (METADOL)	68
MAR-ALLOPURINOL	152	M-CAL	102	METHADONE LOCK BOX	166
MAR-AMITRIPTYLINE	77	M-CAL D	102	METHADONE POWDER (OAT)	67
MAR-AMLODIPINE	49	M-CLARITHROMYCIN	4	METHADOSE	67
MAR-ANASTROZOLE	16	M-D	148	METHADOSE DEL. W DIRECT INTER (OAT)	67
MAR-ATENOLOL	47	M-DONEPEZIL	26	METHADOSE DEL. W/OUT DIR INTER (OAT)	67
MAR-ATORVASTATIN	40	M-DULOXETINE	79	METHADOSE W DIRECT INTERACTION (OAT)	67
MARAVIROC	11	MEBENDAZOLE	2	METHADOSE W/OUT DIRECT INTER (OAT)	67
MAR-AZITHROMYCIN	4	MED-ANASTROZOLE	16	METHAZOLAMIDE	113
MAR-CELECOXIB	62	MED-CYPROTERONE	158	METHAZOLAMIDE	113
MAR-CETIRIZINE	1	MED-DORZOLAMIDE-TIMOLOL	113	METHOPRAZINE	84
MAR-CIPROFLOXACIN	6	MED-DUTASTERIDE	152	METHOTREXATE	21
MAR-CITALOPRAM	78	MED-EXEMESTANE	19	METHOTREXATE SODIUM	21
MAR-CLOPIDOGREL	37	MEDI+SURE	100	METHOTRIMEPRAZINE MALEATE	84
MAR-DAPSONE	10	MEDI+SURE (ON)	100	METHYLDOPA	45
MAR-DILTIAZEM T	51	MEDI+SURE SOFT 30G TWIST	162	METHYLDOPA	45
MAR-DOMPERIDONE	121	MEDI+SURE SOFT 33G TWIST	162	METHYLDOPA, HYDROCHLOROTHIAZIDE	45
MAR-DONEPEZIL	26	MED-LATANOPROST	113	METHYLPHENIDATE	90
MAR-DULOXETINE	79	MED-LETROZOLE	20	METHYLPHENIDATE HYDROCHLORIDE	90
MAR-ENALAPRIL	53	MED-MOXIFLOXACIN	7	METHYLPREDNISOLONE	125
MAR-ESCITALOPRAM	79	MED-RIVASTIGMINE	27	METHYLPREDNISOLONE	126
MAR-EZETIMIBE	40	MEDROL	125	METHYLPREDNISOLONE (METHYLPREDNISOLONE SODIUM SUCCINATE)	125
MAR-FLUCONAZOLE	9	MED-ROSUVASTATIN	42	METHYLPREDNISOLONE ACETATE	126
MAR-GABAPENTIN	72	MEDROXY	133	METHYLPREDNISOLONE ACETATE, LIDOCAINE HYDROCHLORIDE	126
MAR-GALANTAMINE ER	26	MEDROXYPROGESTERONE	133	METHYLPREDNISOLONE SODIUM SUCCINATE	126
MAR-LETROZOLE	20	MEDROXYPROGESTERONE ACETATE	133	METOCLOPRAMIDE HYDROCHLORIDE	121
MAR-METHIMAZOLE	134	MED-SOLIFENACIN	145	METOJECT	21
MAR-MIDODRINE	28	MEFENAMIC	64	METOJECT SUBCUTANEOUS	21
MAR-MODAFINIL	90	MEFENAMIC ACID	64	METOLAZONE	105
MAR-MONTELUKAST	108	MEGESTROL	21		
MAR-MOXIFLOXACIN	7	MEGESTROL ACETATE	21		
MAR-OLANZAPINE ODT	85	MEKINIST	25		
MAR-ONDANSETRON	117	MELOXICAM	64		
MAR-PANTOPRAZOLE	120	MELOXICAM	64		
MAR-PAROXETINE	80	MELPHALAN	21		
MAR-PREGABALIN	75	MENTHOL & CAMPHOR IN CORTICOSTEROID LOTION	150		
MAR-QUETIAPINE	87	MENTHOL &/OR CAMPHOR IN STEROID	150		
MAR-RAMIPRIL	55	MEPOLIZUMAB	157		
MAR-RANITIDINE	118	MEPRON	15		
MAR-RISPERIDONE	87	MERCAPTOPYRINE	21		
MAR-RIZATRIPTAN	94	MERCAPTOPYRINE	21		
MAR-RIZATRIPTAN ODT	94	MEROPENEM	3		
MAR-ROSUVASTATIN	42	MEROPENEM	3		
MAR-SERTRALINE	81	MESALAZINE	121		
MAR-SIMVASTATIN	42	M-ESLON	68		
MAR-TOPIRAMATE	76	MESTINON	27		

Non-Insured Health Benefits

METONIA	121	MINT-CIPROFLOX	6	MISCELLANEOUS COMPOUNDED EXTERNAL LOTION	150
METOPROLOL	48	MINT-CITALOPRAM	78	MISCELLANEOUS COMPOUNDED EXTERNAL POWDER	150
METOPROLOL ORAL LIQUID	49	MINT-CLONIDINE	44	MISCELLANEOUS COMPOUNDED EYE/EAR DROP	150
METOPROLOL SR	48	MINT-DORZOLAMIDE/TIMOLOL	113	MISCELLANEOUS COMPOUNDED INJECTION/INFUSION	150
METOPROLOL TARTRATE	48	MINT-DULOXETINE	79	MISCELLANEOUS COMPOUNDED INTERNAL LIQUID	150
METOPROLOL-L	48	MINT-DUTASTERIDE	152	MISCELLANEOUS COMPOUNDED INTERNAL POWDER	150
METROGEL	137	MINT-EPLERENONE	61	MISCELLANEOUS COMPOUNDED SUPPOSITORY	150
METROLOTION	137	MINT-ESCITALOPRAM	79	MISCELLANEOUS COMPOUNDED TOPICAL CREAM	150
METRONIDAZOLE	15	MINT-EZETIMIBE	40	MISCELLANEOUS COMPOUNDED TOPICAL OINTMENT	150
METRONIDAZOLE	15	MINT-FENOFIBRATE E	40	MISOPROSTOL	119
METRONIDAZOLE ORAL LIQUID	15	MINT-FINASTERIDE	152	MISOPROSTOL	119
METRONIDAZOLE, NYSTATIN	137	MINT-FLUOXETINE	79	MISOPROSTOL, DICLOFENAC SODIUM	64
MEXILETINE HYDROCHLORIDE	39	MINT-FUROSEMIDE	105	MISOPROSTOL, MIFEPRISTONE	136
MEZAVANT	121	MINT-GLICLAZIDE MR	132	MITOTANE	21
MEZERA	121	MINT-HYDRALAZINE	45	MK 10	103
M-EZETIMIBE	40	MINT-HYDROCHLOROTHIAZIDE	105	MK 20	103
MFER FUMARATE	34	MINT-HYDROXYCHLOROQUINE	15	MK 8	103
M-FOLIQUE	147	MINT-INDOMETHACIN	64	MK20 SOLUBLE	103
M-HC	140	MINT-IRBESARTAN	57	MMAGNESIUM GLUCONATE	103
M-HC UREA	140	MINT-IRBESARTAN/HCTZ	57	M-MOXIFLOXACIN	7
MICARDIS	59	MINT-ITRACONAZOLE	9	MMT-174 ADHESIVE	161
MICARDIS PLUS	59	MINT-LEVOCARB	95	MOCLOBEMIDE	80
MICATIN	138	MINT-LOSARTAN	58	MOCLOBEMIDE	80
MICONAZOLE 3 DAY OVULE TREATMENT	138	MINT-LOSARTAN/HCTZ	58	MODAFINIL	90
MICONAZOLE NITRATE	138	MINT-METFORMIN	129	MOGADON	92
MICOZOLE	138	MINT-MONTELUKAST	108	MOMETASONE CREAM	141
MICRO K	103	MINT-OLANZAPINE ODT	85	MOMETASONE FUROATE	111
MICROLAX	116	MINT-OLOPATADINE	110	MONA LISA 10	99
MICROLET LANCET	162	MINT-ONDANSETRON	117	MONA LISA 5	99
MICRONOR 28-DAY	128	MINT-PANTOPRAZOLE	120	MONA LISA N	99
MICTORYL PEDIATRIC	145	MINT-PAROXETINE	80	MONISTAT 3	138
MIDAMOR	105	MINT-PERINDOPRIL	54	MONISTAT 3 DUAL-PAK	138
MIDODRINE HYDROCHLORIDE	28	MINT-PIGLITAZONE	133	MONISTAT 7	138
MIDOSTAURIN	21	MINT-PRAVASTATIN	41	MONISTAT 7 DUAL-PAK	138
MIFEGYMISO	136	MINT-PREGABALIN	75	MONISTAT DERM	138
MIGRANAL	30	MINT-QUETIAPINE	86	MONOJECT	163
MILK OF MAGNESIA	115	MINT-RAMIPRIL	55	MONOJECT ALCOHOL WIPES	161
MINERAL OIL	115	MINT-RISPERIDON	87	MONOLET 21G LANCET	162
MINERAL OIL (HEAVY)	115	MINT-ROSUVASTATIN	42	MONOLET THIN (MONOJECT) 28G	162
MINERAL OIL, WHITE PETROLATUM	114	MINT-SERTRALINE	81	MONTELUKAST	108
MINESTRIN 1/20 (21-DAY)	127	MINT-SIMVASTATIN	42	MONTELUKAST SODIUM	107
MINESTRIN 1/20 (28-DAY)	127	MINT-SOLIFENACIN	145	MONTELUKAST SODIUM	108
MINIMS ATROPINE	112	MINT-TOLTERODINE	145	MONTKIDDY BLUE NEEDLE 32GX4MM	163
MINIMS CYCLOPENTOLATE	112	MINT-TOPIRAMATE	76	MONTKIDDY GREEN NEEDLE 32GX4MM	163
MINIMS PHENYLEPHRINE	112	MINT-ZOLMITRIPTAN	95	MONTKIDDY PINK NEEDLE 32GX4MM	163
MINIMS PILOCARPINE	113	MIO BLUE 6MMX18	160	MONTKIDDY YELLOW NEEDLE 32GX4MM	163
MINIMS PREDNISOLONE	111	MIO BLUE 6MMX23	160	MONUROL	15
MINIPRESS	46	MIO CLEAR 6MMX32	160	MORPHINE HYDROCHLORIDE	68
MINITRAN	45	MIO CLEAR 9MMX32	160	MORPHINE SR	68
MINOCYCLINE	7	MIO PINK 6MMX18	160	MORPHINE SULFATE	68
MINOCYCLINE HYDROCHLORIDE	7	MIO PINK 6MMX23	160	MORPHINE SULFATE (KADIAN)	69
MIN-OVRAL 21	127	MIOSTAT	113	MOTION SICKNESS	117
MIN-OVRAL 28	127	MIRABEGRON	146	MOTRIN	63
MINOXIDIL	45	MIRAPEX	96	MOVISSE	128
MINT-ABACAVIR	10	MIRAPEX (ON)	96	MOXIFLOXACIN	7
MINT-ACITRETIN	142	MIRENA	127		
MINT-ALENDRONATE	154	MIRTAZAPINE	80		
MINT-AMLODIPINE	50	MIRTAZAPINE	80		
MINT-ANASTROZOLE	16	MIRVALA 21	127		
MINT-ATENOL	47	MIRVALA 28	127		
MINT-BISOPROLOL	47	MISC LIMITED USE COMPOUND INTERNAL	150		
MINT-CANDESARTAN	56	MISC LIMITED USE EXTERNAL	150		
MINT-CELECOXIB	62	COMPOUND MIXTURE			

Non-Insured Health Benefits

MOXIFLOXACIN HYDROCHLORIDE	7	MYLAN-LAMOTRIGINE	73	NAT-OSELTAMIVIR	13
MOXIFLOXACIN HYDROCHLORIDE (OPHTHALMIC)	110	MYLAN-LANSOPRAZOLE	119	NAT-QUETIAPINE	86
MOZOBIL	37	MYLAN-MINOCYCLINE	7	NAT-RIZATRIPTAN ODT	94
M-PANTOPRAZOLE	120	MYLAN-MIRTAZAPINE	80	NAT-TENOFOVIR	12
M-PAROXETINE	80	MYLAN-MYCOPHENOLATE	158	NATURAL HEALTH PRODUCT	151
MPD THIN LANCET (NS)	162	MYLAN-NEVIRAPINE	11	NATURES BOUNTY PRENATAL VITAMINS	149
MPD ULTRA THIN LANCET (100)	162	MYLAN-NIFEDIPINE	51	NAT-ZOLMITRIPTAN	95
MPD ULTRA THIN LANCET (200)	162	MYLAN-NITRO	45	NAVANE	88
M-PEG 3350	115	MYLAN-ONDANSETRON	117	NELFINAVIR MESYLATE	11
M-PRAVASTATIN	41	MYLAN-PANTOPRAZOLE T	120	NEOCATE W/ DHA & ARA 400G PDR	166
M-PREGABALIN	75	MYLAN-PERINDOPRIL/INDAPAMIDE	54	NEO-FER	34
M-RANITIDINE	119	MYLAN-PROPAFENONE	39	NEORAL	157
MS CONTIN SR	68	MYLAN-RISPERIDONE ODT	88	NEOSTIGMINE BROMIDE	27
MS IR	68	MYLAN-RIZATRIPTAN ODT	94	NEO-ZOL	138
M-SENNOSIDES	116	MYLAN-SIMVASTATIN	42	NEPAFENAC	112
M-SULFATE FERREUX	34	MYLAN-SUMATRIPTAN	94	NESTL MATERNA	149
MUCILLIUM	116	MYLAN-TENOFOVIR DISOPROXIL	12	NEULASTA	37
MULTIVITAMINS (CHILDREN AND YOUTH)	149	MYLAN-TOLTERODINE ER	145	NEULEPTIL	86
MULTIVITAMINS (PRENATAL)	149	MYLAN-TOPIRAMATE	76	NEUPOGEN	37
MUPIROCIN	137	MYLAN-VALACYCLOVIR	13	NEUPOGEN (ON)	37
MUPIROCIN CALCIUM	137	MYLAN-VANCOMYCIN	8	NEUPOGEN (QC)	37
MURO 128	114	MYLAN-VERAPAMIL	52	NEUPRO	97
M-VENLAFAXINE XR	82	MYLAN-VERAPAMIL SR	52	NEURONTIN	72
MYA	127	MYLERAN	17	NEUTROGENA	142
MYCOBUTIN	10	MYOCHRYSSINE	123	NEVANAC	112
MYCOPHENOLATE	158	MYRBETRIQ	146	NEVIRAPINE	11
MYCOPHENOLATE MOFETIL	158	NABILONE	118	NIACIN	147
MYCOPHENOLATE MOFETIL	158	NACL SALINE PF	103	NIACIN	147
MYCOPHENOLATE SODIUM	158	NADOLOL	49	NICHIT	32
MYDFRIN	112	NADOLOL	49	NICODERM	32
MYDRIACYL	112	NADROPARIN CALCIUM	36	NICORETTE GUM	31
MYFORTIC	158	NADRYL	1	NICORETTE INHALER	31
MYHEALTH SYRINGE CASE-7	165	NAFARELIN ACETATE	129	NICORETTE LOZENGE	32
MYHEALTH SYRINGE CASE-SINGLE	165	NALCROM	108	NICORETTE QUICKMIST	32
MYLAN-ABACAVIR/LAMIVUDINE	10	NALOXONE	70	NICOTINE (GUM)	31
MYLAN-ACYCLOVIR	13	NALOXONE HYDROCHLORIDE	70	NICOTINE (INHALER)	31
MYLAN-ALMOTRIPTAN	93	NALOXONE KIT	70	NICOTINE (LOZENGE)	32
MYLAN-AMLODIPINE	50	NALTREXONE HYDROCHLORIDE	71	NICOTINE (PATCH)	32
MYLAN-AMOXICILLIN	5	NALTREXONE HYDROCHLORIDE	71	NICOTINE (SPRAY)	32
MYLAN-ATAZANAVIR	10	NAPHAZOLINE HYDROCHLORIDE	112	NICOTINE GUM	31
MYLAN-ATORVASTATIN	40	NAPROSYN	65	NICOTINE TRANSDERMAL	32
MYLAN-BACLOFEN	31	NAPROXEN	64	NICOTINE TRANSDERMAL SYSTEM	32
MYLAN-BECLO AQ	111	NAPROXEN	64	NIDAGEL	137
MYLAN-BUDESONIDE AQ	111	NAPROXEN EC	65	NIFEDIPINE	51
MYLAN-BUPRENORPHINE/NALOXONE	69	NAPROXEN SODIUM	64	NIFEDIPINE	51
MYLAN-BUPROPION XL	77	NAPROXEN SODIUM DS	65	NILOTINIB	21
MYLAN-CILAZAPRIL	52	NAPROXEN-NA	64	NILUTAMIDE	21
MYLAN-CIMETIDINE	118	NAPROXEN-NA DF	65	NIMODIPINE	51
MYLAN-CLOBETASOL	139	NARATRIPTAN HYDROCHLORIDE	93	NIMOTOP	51
MYLAN-DIVALPROEX	76	NARCAN	70	NINTEDANIB ESILATE	107
MYLAN-EFAVIRENZ	11	NARDIL	81	NITOMAN	98
MYLAN-EFAVIRENZ/EMTRICITABINE/TENOFOVIR DISOPROXIL FUMARATE	12	NASACORT AQ	111	NITRAZEPAM	92
MYLAN-ENALAPRIL	53	NASONEX	111	NITRO-DUR	45
MYLAN-ESCITALOPRAM	79	NAT-ANASTROZOLE	16	NITROFURANTOIN	15
MYLAN-FLUCONAZOLE	9	NAT-CITALOPRAM	78	NITROFURANTOIN	15
MYLAN-GALANTAMINE ER	26	NAT-DONEPEZIL	26	NITRO-FURANTOIN ORAL LIQUID	15
MYLAN-GLICLAZIDE MR	132	NAT-ERLOTINIB	18	NITROGLYCERIN	45
MYLAN-HYDROXYUREA	19	NAT-ESCITALOPRAM	79	NITROLINGUAL PUMPSPRAY	45
MYLAN-INDAPAMIDE	105	NAT-GRANISETRON	117	NITROSTAT	45
		NAT-IMATINIB	19	NIX	138
		NAT-LETROZOLE	20	NIX DERMAL	138
		NAT-LEVETIRACETAM	74	NIZATIDINE	118
		NAT-OMEPRAZOLE DR	120	NIZORAL	138
		NAT-ONDANSETRON	117		

Non-Insured Health Benefits

NOLVADEX-D	24	OCRELIZUMAB	153	OPTIMYXIN	110
NON POLLEN	151	OCREVUS	153	OPTION 2	127
NORETHINDRONE	128	OCTREOTIDE ACETATE	150	OPUS CAL D	102
NORETHINDRONE, ETHINYL ESTRADIOL	128	OCTREOTIDE ACETATE OMEGA	150	OPUS SENNOSIDES	116
NORFLOXACIN	7	OCUFLOX	110	ORACORT DENTAL PASTE	141
NORFLOXACIN	7	ODAN K20	103	ORAP	86
NORGESTIMATE, ETHINYL ESTRADIOL	128	ODAN K8	103	ORCIPRENALINE	30
NORITATE	137	ODAN LIQUOR CARBONIS DETERGENT	142	ORCIPRENALINE SULFATE	30
NORTRIPTYLINE HYDROCHLORIDE	80	ODAN SODIUM CHLORIDE	114	ORENCIA	155
NORVASC	50	ODAN-ERYTHROMYCIN	110	OSELTAMIVIR	13
NORVIR	12	ODAN-FLUOXETINE	80	OSIMERTINIB	22
NOVA MAX	100	ODAN-SODIUM CHLORIDE	114	OVIMA 21	127
NOVAMILOR	105	ODEFSEY	11	OVIMA 28	127
NOVAMOXIN	5	OESCLIM	128	OXAZEPAM	92
NOVASEN	62	OFEV	107	OXAZEPAM	92
NOVA-T	99	OFLOXACIN	110	OXCARBAZEPINE	74
NOVO-CLINDAMYCIN	8	OLANZAPINE	84	OXCARBAZEPINE (SUSPENSION)	74
NOVO-ETIDRONATECAL	154	OLANZAPINE	84	OXEZE TURBUHALER	29
NOVOFINE 30GX 6MM NEEDLE	162	OLANZAPINE ODT	85	OXPAM	92
NOVOFINE 30GX 8MM NEEDLE	162	OLAPARIB	22	OXTRIPHYLLINE	146
NOVOFINE 32G TIP PEN NEEDLE	163	OLESTYR	39	OXYBUTYNIN	145
NOVOFINE PLUS 4MM NEEDLE	163	OLMESARTAN MEDOXOMIL	59	OXYBUTYNIN CHLORIDE	145
NOVO-FLUCONAZOLE	9	OLMESARTAN MEDOXOMIL, HYDROCHLOROTHIAZIDE	59	OXYCODONE	69
NOVO-GESIC	70	OLMETEC	59	OXYCODONE HYDROCHLORIDE	69
NOVO-GESIC FORTE	70	OLMETEC PLUS	59	OXYCODONE/ACET	65
NOVO-HYDROXYZIN	93	OLMETEC PLUS	59	OXYCODONE-ACET	65
NOVOLIN GE 30/70	130	OLODATEROL HYDROCHLORIDE, TITROPIUM BROMIDE MONOHYDRATE	30	OXY-IR	69
NOVOLIN GE 30/70 PENFILL	130	OLOPATADINE HYDROCHLORIDE	110	OYSTER SHELL CALCIUM	102
NOVOLIN GE 40/60 PENFILL	130	OLSALAZINE SODIUM	121	PALAFER	34
NOVOLIN GE 50/50 PENFILL	130	OMALIZUMAB	109	PALBOCICLIB	22
NOVOLIN GE NPH	130	OMEPRAZOLE	120	PALIPERIDONE PALMITATE	85
NOVOLIN GE NPH 100U/ML PENFILL	130	OMEPRAZOLE MAGNESIUM	120	PAL-TIZANIDINE	31
NOVOLIN GE NPH PENFILL	130	OMEPRAZOLE ORAL LIQUID	120	PAMIDRONATE	154
NOVOLIN GE TORONTO	131	OMEPRAZOLE-20	120	PAMIDRONATE DISODIUM	154
NOVOLIN GE TORONTO PENFILL	131	ONABOTULINUMTOXINA	159	PAMIDRONATE DISODIUM	154
NOVOLIN-PEN NEEDLE	162	ONBREZ BREEZHALER	30	PAMIDRONATE DISODIUM OMEGA	154
NOVO-PENICILLIN G POTASSIUM	5	ONDANSETRON	117	PANTOLOC	120
NOVO-PHENIRAM	1	ONDANSETRON HYDROCHLORIDE	117	PANTOPRAZOLE	120
NOVO-PROFEN	63	ONDISSOLVE ODF	117	PANTOPRAZOLE MAGNESIUM	120
NOVORAPID	131	ONE A DAY WOMEN	149	PANTOPRAZOLE MAGNESIUM	120
NOVOTWIST TIP 30G NEEDLE	163	ONE ALPHA	148	PANTOPRAZOLE SODIUM	120
NOVOTWIST TIP 32G NEEDLE	163	ONE TOUCH DELICA 30G LANCET	162	PANTOPRAZOLE T	120
NRA-AMLODIPINE	49	ONE TOUCH ULTRA	100	PANTOPRAZOLE-40	120
NRA-CITALOPRAM	78	ONE-ALPHA	148	PARADIGM SILHOUETTE 13MMX 43	161
NRA-ESCITALOPRAM	79	ONETOUCH DELICA 33G LANCET	162	PARADIGM SILHOUETTE 13MMX18"	161
NRA-EZETIMIBE	40	ONETOUCH ULTRASOFT LANCET	162	PARADIGM SILHOUETTE 13MMX23	161
NRA-PANTOPRAZOLE	120	ONETOUCH VERIO	100	PARADIGM SILHOUETTE 13MMX32"	161
NRA-PAROXETINE	80	ONETOUCH VERIO (ON)	100	PARADIGM SILHOUETTE 17MMX23	161
NRA-PREGABALIN	75	ONGLYZA	130	PARADIGM SILHOUETTE 17MMX32"	161
NU-CAL	102	OPIOID COMPOUNDED	150	PARADIGM SILHOUETTE 17MMX43	161
NU-CAL D	102	OPTICHAMBER	160	PARADIGM SILHOUETTE CANNULA 13MM	161
NUCALA	157	OPTICHAMBER DIAMOND (CHAMBER)	160	PARADIGM SILHOUETTE CANNULA 17MM	161
NUTRAMIGEN A+	166	OPTICHAMBER DIAMOND LARGE MASK	160	PARADIGM SURE-T 29G 6MMX18	161
NUTRAMIGEN A+ LGG 566G PDR	166	OPTICHAMBER DIAMOND MEDIUM MASK	160	PARADIGM SURE-T 29G 6MMX23	161
NUTRITIONAL SUPPLEMENT	166	OPTICHAMBER DIAMOND SMALL MASK	160	PARADIGM SURE-T 29G 8MMX23	161
NUVARING	127	OPTICHAMBER LARGE MASK	160	PARIENT	120
NYADERM	138	OPTICHAMBER MEDIUM MASK	160	PARNATE	82
NYDA	138	OPTICHAMBER SMALL MASK	160	PAROMOMYCIN SULFATE	15
NYSTATIN	9	OPTICROM	110	PAROXETINE	80
OBETICHOLIC ACID	121	OPTIHALER	160	PAROXETINE HYDROCHLORIDE	80
O-CALCIUM	102			PARSITAN	95
OCALIVA	121			PATANOL	110
OCCLUSAL HP	142			PATE D'IHLE	142

Non-Insured Health Benefits

PÂTE D'IHLE	142	PHARMALGEN MIXED VESPID VENOM PROTEIN	151	PMS-AMITRIPTYLINE	77
PAT-GALANTAMINE ER	26	PHARMALGEN WASP VENOM PROTEIN	151	PMS-AMLODIPINE	49
PAXIL	80	PHARMALGEN WHITE FACED HORNET VENOM	151	PMS-AMLODIPINE-ATORVASTATIN	50
PAZOPANIB	22	PHARMALGEN YELLOW HORNET VENOM PROTEIN	151	PMS-AMOXICILLIN	5
PDP-ACETAMINOPHEN	69	PHARMALGEN YELLOW JACKET VENOM PROTEIN	151	PMS-AMPHETAMINES XR	89
PDP-BENZTROPINE	95	PHARMA-RAMIPRIL	55	PMS-ANAGRELIDE	36
PDP-DESONIDE	140	PHARMA-SIMVASTATIN	42	PMS-ANASTROZOLE	16
PDP-DIPHENHYDRAMINE	1	PHENAZOPYRIDINE COMPOUNDED	150	PMS-ARIPIRAZOLE	82
PDP-ERYTHROMYCIN	110	PHENELZINE SULFATE	81	PMS-ASA EC	62
PDP-ISONIAZID	10	PHENOBARB	71	PMS-ATENOLOL	47
PDP-PROCYCLIDINE	95	PHENOBARBITAL	71	PMS-ATOMOXETINE	97
PDP-PYRAZINAMIDE	10	PHENYLEPHRINE	112	PMS-ATORVASTATIN	40
PEDIAFER	34	PHENYLEPHRINE HYDROCHLORIDE	112	PMS-AZITHROMYCIN	4
PEDIALYTE	103	PHENYTOIN	71	PMS-BACLOFEN	31
PEDIAPHEN	69	PHILIPS MAGNESIA	115	PMS-BENZTROPINE	95
PEDIAPRED	126	PHILLIPS MILK OF MAGNESIA	115	PMS-BENZYDAMINE	112
PEDIATRIC ELECTROLYTE	103	PHOSLAX	116	PMS-BETAHISTINE	98
PEDIATRIX	69	PHOSPHATES	116	PMS-BEZAFIBRATE	40
PEDIAVIT	149	PHYTONADIONE	149	PMS-BICALUTAMIDE	17
PEDIAVIT D	148	PICO-SALAX	115	PMS-BISACODYL	115
PEG 3350	116	PILOCARPINE	113	PMS-BISOPROLOL	47
PEGASYS	12	PILOCARPINE HYDROCHLORIDE	27	PMS-BOSENTAN	46
PEGETRON KIT	12	PILOCARPINE HYDROCHLORIDE	27	PMS-BRIMONIDINE	112
PEGFILGRASTIM	37	PILOCARPINE NITRATE	113	PMS-BROMOCRIPTINE	96
PEGFILGRASTIM (LAPELGA)	37	PIMECROLIMUS	143	PMS-BUPRENORPHINE-NALOXONE	69
PEGINTERFERON ALFA-2A	12	PIMOZIDE	86	PMS-BUPROPION SR	77
PEGINTERFERON ALFA-2B, RIBAVIRIN	12	PIMOZIDE	86	PMS-BUSPIRONE	93
PEGINTERFERON BETA-1A	12	PINAVERIUM BROMIDE	121	PMS-CANDESARTAN	56
PEGLYTE	115	PINDOLOL	49	PMS-CANDESARTAN HCTZ	57
PEN NEEDLE	162	PINDOLOL	49	PMS-CAPTOPRIL	52
PENICILLAMINE	124	PIOGLITAZONE HYDROCHLORIDE	133	PMS-CARBAMAZEPINE	72
PENICILLIN G	5	PIPERACILLIN SODIUM/TAZOBACTAM SODIUM	6	PMS-CARVEDILOL	48
PENICILLIN G BENZATHINE	5	PIPERACILLIN, TAZOBACTAM	6	PMS-CELECOXIB	62
PENICILLIN G POTASSIUM	5	PIPERONYL BUTOXIDE, PYRETHRINS	138	PMS-CEPHALEXIN	3
PENICILLIN G SODIUM	5	PIPORTIL L4	86	PMS-CETIRIZINE	1
PENICILLIN G SODIUM	5	PIPOTIAZINE PALMITATE	86	PMS-CILAZAPRIL	52
PENICILLIN G STERILE INFUSION	5	PIRFENIDONE	107	PMS-CIPROFLOXACIN	6
PENICILLIN V POTASSIUM	5	PIROXICAM	65	PMS-CITALOPRAM	78
PENTASA	121	PIZOTIFEN MALATE	95	PMS-CITRALOPRAM	4
PENTOSAN POLYSULFATE SODIUM	151	PLAN B	127	PMS-CLARITHROMYCIN	4
PENTOXIFYLLINE	37	PLAQUENIL	15	PMS-CLOBAZAM	71
PENTOXIFYLLINE	37	PLASTIPAK MICRO	163	PMS-CLOBETASOL	139
PEN-VK	5	PLAVIX	37	PMS-CLONAZEPAM	71
PEPTO BISMOL	115	PLEGRIDY	12	PMS-CLONAZEPAM-R	71
PERICHLOR	110	PLENDIL	51	PMS-CLOPIDOGREL	37
PERICYAZINE	86	PLERIXAFOR	37	PMS-COLCHICINE	152
PERIDEX	110	PMS DESIPRAMINE	78	PMS-CYCLOBENZAPRINE	31
PERINDOPRIL ERBUMINE	54	PMS DEXAMETHASONE	125	PMS-DESMOPRESSIN	133
PERINDOPRIL ERBUMINE	54	PMS FLUPHENAZINE	84	PMS-DEXAMETHASONE	111
PERINDOPRIL ERBUMINE, INDAPAMIDE	54	PMS HYDROMORPHONE	67	PMS-DIAZEPAM	91
PERMETHRIN	138	PMS HYDROXYZINE	93	PMS-DICLOFENAC	63
PERPHENAZINE	86	PMS PERPHENAZINE	86	PMS-DICLOFENAC-MISOPROSTOL	64
PERPHENAZINE	86	PMS PROCHLORPERAZINE	86	PMS-DILTIAZEM CD	51
PETROLATUM, MINERAL OIL	114	PMS TRAZODONE	82	PMS-DIMENHYDRINATE	117
PHARIXIA	112	PMS-ABACAVIR/LAMIVUDINE	10	PMS-DIPHENHYDRAMINE	1
PHARMA-AMLODIPINE	49	PMS-ACETAMINOPHEN	65	PMS-DIVALPROEX	76
PHARMA-CAL	102	PMS-ALENDRONATE	154	PMS-DOMPERIDONE	121
PHARMA-D	148	PMS-AMANTADINE	10	PMS-DONEPEZIL	26
PHARMA-ESCITALOPRAM	79	PMS-AMIODARONE	39	PMS-DORZOLAMIDE-TIMOLOL	113
PHARMA-K20	103			PMS-DOXAZOSIN	46
PHARMA-LACOSAMIDE	73			PMS-DULOXETINE	79
PHARMA-LACTULOSE	102			PMS-DUTASTERIDE	152
PHARMALGEN HONEY BEE VENOM	151			PMS-EMTRICITABINE-TENOFOVIR	12
				PMS-ENTECAVIR	13
				PMS-ERLOTINIB	18

Non-Insured Health Benefits

PMS-ESCITALOPRAM	79	PMS-NITROFURANTOIN	15	PMS-VALPROIC ACID	77
PMS-EZETIMIBE	40	PMS-NIZATIDINE	118	PMS-VANCOMYCIN	8
PMS-FAMCICLOVIR	13	PMS-NYSTATIN	9	PMS-VANCOMYCIN 1 G	8
PMS-FENTANYL MTX	66	PMS-OLANZAPINE	84	PMS-VENLAFAXINE XR	82
PMS-FERROUS SULFATE	34	PMS-OLANZAPINE ODT	85	PMS-VERAPAMIL SR	52
PMS-FINASTERIDE	152	PMS-OLMESARTAN	59	PMS-ZOLMITRIPTAN	95
PMS-FLUCONAZOLE	9	PMS-OMEPRAZOLE	120	PMS-ZOLMITRIPTAN ODT	95
PMS-FLUOXETINE	79	PMS-ONDANSETRON	117	POCKET CHAMBER	160
PMS-FLUPHENAZINE	84	PMS-OXYBUTYNIN	145	POCKET CHAMBER WITH ADULT MASK	160
PMS-FLUTAMIDE	19	PMS-OXYCODONE	69	POCKET CHAMBER WITH INFANT MASK	160
PMS-FOSINOPRIL	53	PMS-PAMIDRONATE	154	POCKET CHAMBER WITH MEDIUM MASK	160
PMS-FUROSEMIDE	105	PMS-PANTOPRAZOLE	120	POCKET CHAMBER WITH SMALL MASK	160
PMS-GABAPENTIN	72	PMS-PAROXETINE	81	PODOFILM	143
PMS-GALANTAMINE ER	27	PMS-PERINDOPRIL	54	PODOFILOX	143
PMS-GEMFIBROZIL	40	PMS-PINDOLOL	49	PODOPHYLLIN	143
PMS-GLYBURIDE	133	PMS-PIOGLITAZONE	133	PODS	160
PMS-HALOPERIDOL	84	PMS-POLYTRIMETHOPRIM	110	POLISTES SPP VENOM PROTEIN EXTRACT	151
PMS-HYDROCHLOROTHIAZIDE	105	PMS-POTASSIUM	103	POLLEN	151
PMS-HYDROMORPHONE	67	PMS-PRAMIPEXOLE	96	POLLEN AND NON POLLEN	151
PMS-IBUPROFEN	64	PMS-PRAVASTATIN	41	POLLINEX R	151
PMS-IMATINIB	19	PMS-PREDNISOLONE	126	POLYETHYLENE GLYCOL	115
PMS-IPRATROPIUM	28	PMS-PREGABALIN	75	POLYETHYLENE GLYCOL 3350	115
PMS-IRBESARTAN	57	PMS-PROCHLORPERAZINE	86	POLYETHYLENE GLYCOL 3350	115
PMS-IRBESARTAN-HCTZ	57	PMS-PROGESTERONE	134	POLYETHYLENE GLYCOL 3350, SODIUM SULFATE, SODIUM BICARBONATE, SODIUM CHLORIDE, POTASSIUM CHLORIDE	116
PMS-ISMN	45	PMS-PROPAFENONE	39	POLYETHYLENE GLYCOL 3350, SODIUM SULFATE, SODIUM BICARBONATE, SODIUM CHLORIDE, POTASSIUM CHLORIDE, BISACODYL	116
PMS-ISOSORBIDE	45	PMS-PROPRANOLOL	49	POLYMYXIN B SULFATE, BACITRACIN ZINC	110
PMS-KETOPROFEN	64	PMS-QUETIAPINE	86	POLYMYXIN B SULFATE, BACITRACIN ZINC, GRAMICIDIN	137
PMS-LACTULOSE	102	PMS-QUINAPRIL	54	POLYMYXIN B SULFATE, GRAMICIDIN	110
PMS-LACTULOSE-PHARMA	102	PMS-RABEPRAZOLE	120	POLYMYXIN B SULFATE, TRIMETHOPRIM SULFATE	110
PMS-LAMOTRIGINE	73	PMS-RALOXIFENE	129	POLYSACCHARIDE IRON COMPLEX	34
PMS-LANSOPRAZOLE	119	PMS-RAMIPRIL	55	POLYSPORIN	110
PMS-LATANOPROST	113	PMS-RAMIPRIL-HCTZ	56	POLYSPORIN ANTIBIOTIC	137
PMS-LATANOPROST-TIMOLOL	113	PMS-RANITIDINE	119	POLYSPORIN EYE AND EAR	110
PMS-LEFLUNOMIDE	156	PMS-RISEDRONATE	155	POLYSPORIN TRIPLE	137
PMS-LETROZOLE	20	PMS-RISPERIDONE	87	POLYTOPIC	137
PMS-LEVETIRACETAM	74	PMS-RIVASTIGMINE	27	POLYTRIM	110
PMS-LEVOCARB	95	PMS-RIZATRIPTAN RDT	94	POLYVINYL ALCOHOL	114
PMS-LEVOFLOXACIN	6	PMS-ROPINIROLE	96	POMALIDOMIDE	22
PMS-LIDOCAINE VISCOSUS	135	PMS-ROSUVASTATIN	42	POMALYST	22
PMS-LISINAPRIL	53	PMS-SALBUTAMOL	30	PONATINIB HYDROCHLORIDE	23
PMS-LITHIUM CARBONATE	93	PMS-SENNOSIDES	116	PONSTAN	64
PMS-LITHIUM CITRATE	93	PMS-SERTRALINE	81	PORTIA 21	127
PMS-LOPERAMIDE	115	PMS-SILDENAFIL R	45	PORTIA 28	127
PMS-LORAZEPAM	92	PMS-SIMVASTATIN	42	POTASSIUM CHLORIDE	103
PMS-LOSARTAN	58	PMS-SODIUM CROMOGLYCATE	108	POTASSIUM CITRATE	102
PMS-LOSARTAN-HCTZ	58	PMS-SOLIFENACIN	145	POTASSIUM CITRATE	103
PMS-LOVASTATIN	41	PMS-SOTALOL	49	POVIDONE-IODINE	138
PMS-MELOXICAM	64	PMS-SULFASALAZINE	7	PRADAXA	35
PMS-METFORMIN	129	PMS-SUMATRIPTAN	94	PRALUENT	44
PMS-METHOTREXATE	21	PMS-TELMISARTAN	59	PRAMIPEXOLE	96
PMS-METHYLPHENIDATE	90	PMS-TELMISARTAN-HCTZ	59	PRAMIPEXOLE DIHYDROCHLORIDE	96
PMS-METHYLPHENIDATE ER	90	PMS-TENOFOVIR	12	PRAVACHOL	41
PMS-METOPROLOL-B	48	PMS-TERAZOSIN	46	PRAVASTATIN	41
PMS-METOPROLOL-L	48	PMS-TERBINAFINE	9		
PMS-MINOCYCLINE	7	PMS-TESTOSTERONE	126		
PMS-MIRTAZAPINE	80	PMS-TETRABENAZINE	98		
PMS-MOCLOBEMIDE	80	PMS-TIAPROFENIC	65		
PMS-MOMETASONE	141	PMS-TIMOLOL	113		
PMS-MONTELUKAST	108	PMS-TOPIRAMATE	76		
PMS-MOXIFLOXACIN	110	PMS-TRANDOLAPRIL	56		
PMS-NABILONE	118	PMS-TRAZODONE	82		
PMS-NAPROXEN	64	PMS-TRIHENXYPHENIDYL	95		
PMS-NAPROXEN EC	65	PMS-URSODIOL	116		
PMS-NIFEDIPINE	51	PMS-VALACYCLOVIR	13		

Non-Insured Health Benefits

PRAVASTATIN SODIUM	41	PROCYCLIDINE HYDROCHLORIDE	95	QUICK-SET 6MMX32	161
PRAVASTATIN-10	41	PROCYTOX	18	QUICK-SET 6MMX43 TUBING	161
PRAVASTATIN-20	41	PRO-DEXAMETHASONE	125	QUICK-SET 9MMX23 TUBING	161
PRAVASTATIN-40	42	PRO-ENALAPRIL	53	QUICK-SET 9MMX32	161
PRAxis ASA DAILY LOW DOSE	62	PRO-FENO-SUPER	40	QUICK-SET 9MMX43 TUBING	161
PRAZOSIN HYDROCHLORIDE	46	PRO-FLUCONAZOLE	9	QUINAPRIL	54
PRECISION XTRA	100	PRO-FLUOXETINE	79	QUINAPRIL, HYDROCHLOROTHIAZIDE	55
PRED FORTE	111	PRO-GABAPENTIN	72	QUINSAIR	6
PRED MILD	111	PROGESTERONE	134	QVAR	125
PREDNISOLONE ACETATE	111	PROGLYCEM	45	R & C SHAMPOO WITH CONDITIONER	138
PREDNISOLONE ACETATE, SULFACETAMIDE SODIUM	111	PROGRAF	158	RABEPRAZOLE	120
PREDNISOLONE SODIUM PHOSPHATE	111	PRO-HYDROXYQUINE	15	RABEPRAZOLE EC	120
PREDNISOLONE/SULFACETAMIDE	111	PRO-INDAPAMIDE	105	RABEPRAZOLE SODIUM	120
PREDNISONE	126	PRO-ISMN	45	RALOXIFENE HYDROCHLORIDE	129
PREDNISONE ORAL LIQUID	126	PRO-K 20	103	RALTEGRAVIR POTASSIUM	11
PREGABALIN	75	PRO-LEVETIRACETAM	74	RAMIPRIL	55
PREGABALIN	75	PRO-LEVOCARB	95	RAMIPRIL	55
PREMARIN	128	PROLIA	154	RAMIPRIL, HYDROCHLOROTHIAZIDE	55
PREPLUS	128	PRO-LISINOPRIL	53	RAMIPRIL-HCTZ	56
PRENATAL AND POSTPARTUM VITAMINS AND MINERALS	149	PROLOPA	95	RAN-ALENDRONATE	154
PREVACID	119	PRO-LORAZEPAM	92	RAN-AMLODIPINE	50
PREVACID FASTAB	119	PRO-METFORMIN	129	RAN-ANASTROZOLE	16
PREVEX HC	140	PROMETRIUM	134	RAN-ATENOLOL	47
PREZCOBIX	10	PRO-MIRTAZAPINE	80	RAN-ATORVASTATIN	40
PREZISTA	10	PRO-NAPROXEN	65	RAN-BICALUTAMIDE	17
PRIMAQUINE	15	PROPADERM	139	RAN-CANDESARTAN	56
PRIMAQUINE PHOSPHATE	15	PROPAFENONE	39	RAN-CARVEDILOL	48
PRIMIDONE	71	PROPAFENONE HYDROCHLORIDE	39	RAN-CEFPROZIL	2
PRIMIDONE	71	PRO-PIOGLITAZONE	133	RAN-CELECOXIB	62
PRINIVIL	53	PROPIVERINE HYDROCHLORIDE	145	RAN-CIPROFLOX	6
PRIVA-CELECOXIB	62	PROPRANOLOL (HEMANGIOL)	49	RAN-CITALO	78
PRIVA-CETIRIZINE	1	PROPRANOLOL HYDROCHLORIDE	49	RAN-CLARITHROMYCIN	4
PRIVA-ESCITALOPRAM	79	PROPRANOLOL ORAL LIQUID	49	RAN-CLOPIDOGREL	37
PRIVA-EZETIMIBE	40	PROPYLTHIOURACIL	134	RAN-CYPROTERONE/ETHINYL ESTRADIOL	158
PRIVA-FLUCONAZOLE	9	PROPYL-THYRACIL	134	RAN-DOMPERIDONE	121
PRIVA-PANTOPRAZOLE	120	PRO-QUETIAPINE	86	RAN-DONEPEZIL	26
PRIVA-VALACYCLOVIR	13	PRO-RABEPRAZOLE	120	RAN-DULOXETINE	79
PRO AMOX	5	PRO-RAMIPRIL	55	RAN-ENALAPRIL	53
PRO-AAS	62	PRO-RISPERIDONE	87	RAN-ESCITALOPRAM	79
PRO-AMIODARONE	39	PROSCAR	152	RAN-EZETIMIBE	40
PRO-AMOX	5	PRO-SOTALOL	49	RAN-EZETIMIBE	40
PRO-AZITHROMYCINE	4	PROSTIGMIN	27	RAN-FENTANYL MATRIX	66
PRO-BICALUTAMIDE	17	PROTOPIC	143	RAN-FINASTERIDE	152
PRO-BISOPROLOL	47	PRO-TOPIRAMATE	76	RAN-FLUOXETINE	79
PROCAINAMIDE HYDROCHLORIDE	39	PROTRIN DF	7	RAN-FOSINOPRIL	53
PROCAL 500	102	PRO-VALACYCLOVIR	13	RAN-GABAPENTIN	72
PROCALD 400	102	PROVERA	133	RAN-GLICLAZIDE	132
PROCAN SR	39	PROZAC	79	RAN-GLICLAZIDE MR	132
PROCARBAZINE HYDROCHLORIDE	23	PSYLLIUM MUCILLOID	116	RANIBIZUMAB	114
PRO-CEFADROXIL	2	PULMICORT NEBUAMP	125	RAN-IRBESARTAN	57
PRO-CEFUROXIM	3	PULMICORT TURBUHALER	125	RAN-IRBESARTAN HCTZ	57
PROCET-30	65	PULMOPHYLLINE	146	RANITIDINE	119
PROCHLORAZINE	86	PURAMINO A+ PDR	166	RANITIDINE HYDROCHLORIDE	118
PROCHLORPERAZINE	86	PURG-ODAN	115	RAN-LANSOPRAZOLE	119
PROCHLORPERAZINE MALEATE	86	PURINETHOL	21	RAN-LETROZOLE	20
PROCHLORPERAZINE MESYLATE	86	PYRANTEL PAMOATE	2	RAN-LEVETIRACETAM	74
PRO-CIPROFLOXACIN	6	PYRAZINAMIDE	10	RAN-LISINOPRIL	53
PRO-CLONAZEPAM	71	PYRIDOSTIGMINE BROMIDE	27	RAN-LOSARTAN	58
PROCTODAN-HC	141	PYRIDOXINE HYDROCHLORIDE	147	RAN-METFORMIN	129
PROCTOL	140	QUETIAPINE	86	RAN-MONTELUKAST	108
PROCTOSEDYL	140	QUETIAPINE FUMARATE	86	RAN-NABILONE	118
PROCYCLIDINE (PQ)	95	QUETIAPINE XR	87	RAN-OLANZAPINE	84
PROCYCLIDINE HCL	95	QUICK-SET 6MMX18	161	RAN-OLANZAPINE ODT	85
		QUICK-SET 6MMX23 TUBING	161	RAN-OMEPRAZOLE	120

Non-Insured Health Benefits

RAN-ONDANSETRON	117	REPAGLINIDE	131	RIVA-CLARITHROMYCIN	4
RAN-PANTOPRAZOLE	120	REPATHA	44	RIVA-CLINDAMYCIN	7
RAN-PIOGLITAZONE	133	RESERVOIR PARADIGM 5X1.8ML	161	RIVA-CLONAZEPAM	71
RAN-PRAVASTATIN	41	RESERVOIR PARADIGM 7X3.0ML	161	RIVA-CLOPIDOGREL	37
RAN-PREGABALIN	75	RESONIUM CALCIUM	104	RIVACOCET	65
RAN-QUETIAPINE	86	RESOURCE THICKEN CLEAR	166	RIVA-CYCLOBENZAPRINE	31
RAN-RABEPRAZOLE	120	RESOURCE THICKEN UP 6.4G	166	RIVA-CYPROTERONE	158
RAN-RAMIPRIL	55	RESPICHAMBER SILICONE MEDIUM MASK	160	RIVA-D	148
RAN-RAMIPRIL HCTZ	55	RESPICHAMBER SILICONE SMALL MASK	160	RIVA-DONEPEZIL	26
RAN-RANITIDINE	119	RESPICHAMBER VHC W MOUTHPIECE	160	RIVA-DORZOLAMIDE/TIMOLOL	113
RAN-RISPERIDONE	87	RESTORALAX	116	RIVA-DULOXETINE	79
RAN-ROPINIROLE	96	RESTORIL	93	RIVA-DUTASTERIDE	152
RAN-SERTRALINE	81	RESULTZ	138	RIVA-ENALAPRIL	53
RAN-SIMVASTATIN	42	RETIN-A	141	RIVA-ESCITALOPRAM	79
RAN-SOLIFENACIN	145	RETROVIR	12	RIVA-EZETIMIBE	40
RAN-TOPIRAMATE	76	REVATIO	45	RIVA-FINASTERIDE	152
RAN-VALSARTAN	60	REVA	71	RIVA-FLUCONAZOLE	9
RAN-VENLAFAXINE XR	82	REVLIMID	20	RIVA-FLUOXETINE	80
RAPAMUNE	158	REXULTI	83	RIVA-FLUVOX	80
RAPID-D 10MM/110CM	161	REYATAZ	10	RIVA-GABAPENTIN	72
RAPID-D 10MM/60CM	161	RHINARIS NASAL	114	RIVA-HC	140
RAPID-D 10MM/80CM	161	RHINARIS NASAL MIST	114	RIVA-INDAPAMIDE	105
RAPID-D 6MM/110CM	161	RHINARIS-CS	108	RIVA-K 20	103
RAPID-D 6MM/60CM	161	RHINOCORT AQUA	111	RIVA-K 8	103
RAPID-D 6MM/80CM	161	RHINOCORT TURBUHALER	111	RIVA-LANSOPRAZOLE	119
RAPID-D 8MM/110CM	161	RHO-NITRO PUMPSPRAY	45	RIVA-LATANOPROST	113
RAPID-D 8MM/60CM	161	RIBAVIRIN	14	RIVA-LETROZOLE	20
RAPID-D 8MM/80CM	161	RIDAURA	123	RIVA-LOPERAMIDE	115
RATIO-AMCINONIDE	139	RIFABUTIN	10	RIVA-METFORMIN	129
RATIO-BACLOFEN	31	RIFADIN	10	RIVA-METOPROLOL L	48
RATIO-CEFUROXIME	3	RIFAMPIN	10	RIVA-MIRTAZAPINE	80
RATIO-ECTOSONE	139	RIFAMPIN ORAL LIQUID	10	RIVA-MONTELUKAST	108
RATIO-FENOFIBRATE	40	RIFAXIMIN	8	RIVA-MOXIFLOXACIN	7
RATIO-FLUTICASON	111	RILPIVIRINE HYDROCHLORIDE	11	RIVANASE AQ	111
RATIO-HEMCORT-HC	140	RIOCIGUAT	108	RIVA-OLANZAPINE	84
RATIO-IPRA SAL	28	RISEDRONATE	155	RIVA-OLMESARTAN	59
RATIO-IPRATROPIUM	28	RISEDRONATE SODIUM	154	RIVA-OMEPRAZOLE DR	120
RATIO-LACTULOSE	102	RISEDRONATE-35	155	RIVA-OXYBUTYNIN	145
RATIO-LENOLTEC NO 2	65	RISPERDAL	87	RIVA-PANTOPRAZOLE	120
RATIO-LENOLTEC NO 3	65	RISPERDAL CONSTA	88	RIVA-PAROXETINE	81
RATIO-METFORMIN	129	RISPERIDONE	87	RIVA-PERINDOPRIL	54
RATIO-NYSTATIN	138	RISPERIDONE	87	RIVA-PREGABALIN	75
RATIO-SALBUTAMOL	30	RISPERIDONE (CONSTA)	88	RIVA-QUETIAPINE	86
RATIO-TAMSULOSIN	31	RITONAVIR	12	RIVA-RABEPRAZOLE	120
RATIO-TOPISALIC	139	RITUXAN	23	RIVA-RABEPRAZOLE EC	120
REACTINE	1	RITUXIMAB	23	RIVA-RANITIDINE	119
REBIF	153	RIVA OXAZEPAM	92	RIVA-RISEDRONATE	155
REDDY-ATORVASTATIN	40	RIVA SENNA	116	RIVA-RISPERIDONE	87
REDDY-PROGESTERONE	134	RIVA-ALENDRONATE	154	RIVA-ROSUVASTATIN	42
REFRESH CELLUVISC	114	RIVA-AMIODARONE	39	RIVAROXABAN	36
REFRESH LACRI-LUBE	114	RIVA-AMLODIPINE	50	RIVAROXABAN (10)	36
REFRESH LIQUIGEL	114	RIVA-ANASTROZOLE	16	RIVASA	62
REFRESH PLUS	114	RIVA-ARIPIRAZOLE	82	RIVASA EC	62
REFRESH TEARS	114	RIVA-ATENOLOL	47	RIVA-SERTRALINE	81
REFUSAL TO FILL	166	RIVA-ATOMOXETINE	97	RIVASOL-HC	141
REGORAFENIB	23	RIVA-ATORVASTATIN	40	RIVASONE	139
RELAXA	116	RIVA-AZITHROMYCIN	4	RIVASTIGMINE	27
REMERON	80	RIVA-BACLOFEN	31	RIVASTIGMINE HYDROGEN TARTRATE	27
REMERON RD	80	RIVA-BISOPROLOL	47	RIVA-TERBINAFINE	9
REMICADE	156	RIVA-CAL D	102	RIVA-VALACYCLOVIR	13
RENAGEL	104	RIVA-CELECOX	62	RIVA-VENLAFAXINE XR	82
RENFLEXIS	156	RIVA-CIPROFLOXACIN	6	RIVA-VERAPAMIL SR	52
REVELA	104	RIVA-CITALOPRAM	78	RIVA-ZOLMITRIPTAN	95
REPAGLINIDE	131			RIVOTRIL	71

Non-Insured Health Benefits

RIZATRIPTAN BENZOATE	94	SANDOZ CAPECITABINE	17	SANDOZ NARATRIPTAN	94
RIZATRIPTAN ODT	94	SANDOZ CEFPROZIL	2	SANDOZ OFLOXACIN	110
RIZATRIPTAN RDT	94	SANDOZ CELECOXIB	62	SANDOZ OLANZAPINE	84
ROCALTROL	148	SANDOZ CIPROFLOXACIN	6	SANDOZ OLANZAPINE ODT	85
ROFACT	10	SANDOZ CITALOPRAM	78	SANDOZ OLMESARTAN	59
ROLENE	139	SANDOZ CLARITHROMYCIN	4	SANDOZ OLOPATADINE	110
ROPINIROLE	96	SANDOZ CLOPIDOGREL	37	SANDOZ OMEPRAZOLE	120
ROPINIROLE HYDROCHLORIDE	96	SANDOZ COLCHICINE	152	SANDOZ ONDANSETRON	117
ROSONE	139	SANDOZ CYCLOSPORINE	157	SANDOZ	65
ROSUVASTATIN	42	SANDOZ D-FORTE	148	OXYCODONE/ACETAMINOPHEN	
ROSUVASTATIN CALCIUM	42	SANDOZ DICLOFENAC MISOPROSTOL	64	SANDOZ PANTOPRAZOLE	120
ROTIGOTINE	97	SANDOZ DICLOFENAC OPHTHA	111	SANDOZ PAROXETINE	81
ROUGIER-MAGNESIUM	103	SANDOZ DILTIAZEM CD	51	SANDOZ PERINDOPRIL ERBUMINE	54
RUFINAMIDE	76	SANDOZ DILTIAZEM T	51	SANDOZ PERINDOPRIL ERBUMINE/ INDAPAMIDE	54
RUGBY NICOTINE POLACRILEX GUM	31	SANDOZ DIMENHYDRINATE	117	SANDOZ PERINDOPRIL ERBUMINE/ INDAPAMIDE HD	54
RUXOLITINIB	24	SANDOZ DONEPEZIL	26	SANDOZ PIOGLITAZONE	133
RYDAPT	21	SANDOZ DORZOLAMIDE	113	SANDOZ POLYTRIMETHOPRIM	110
RYTHMODAN	39	SANDOZ DORZOLAMIDE/TIMOLOL	113	SANDOZ PRAMIPEXOLE	96
RYTHMOL	39	SANDOZ DULOXETINE	79	SANDOZ PRAVASTATIN	41
S.O.S NALOXONE HYDROCHLORIDE	70	SANDOZ DUTASTERIDE	152	SANDOZ PREDNISOLONE	111
SABRIL	77	SANDOZ ENALAPRIL	53	SANDOZ PREGABALIN	75
SALAGEN	27	SANDOZ ENTACAPONE	95	SANDOZ PROCHLORPERAZINE	86
SALAZOPYRIN	7	SANDOZ ESCITALOPRAM	79	SANDOZ PROCTOMYXIN HC	140
SALAZOPYRIN EN	7	SANDOZ ESTRADIOL DERM	128	SANDOZ QUETIAPINE	86
SALBUTAMOL HFA	30	SANDOZ EZETIMIBE	40	SANDOZ QUETIAPINE XRT	87
SALBUTAMOL SULFATE	30	SANDOZ FAMCICLOVIR	13	SANDOZ RABEPRAZOLE	120
SALICYLIC ACID	142	SANDOZ FELODIPINE	51	SANDOZ RAMIPRIL	55
SALICYLIC ACID IN CORTICOSTEROID CREAM	139	SANDOZ FENOFIBRATE E	40	SANDOZ RANITIDINE	119
SALICYLIC ACID IN NON-MEDICATED OINTMENT	139	SANDOZ FENOFIBRATE S	40	SANDOZ REPAGLINIDE	131
		SANDOZ FENTANYL	66	SANDOZ RISEDRONATE	155
SALICYLIC ACID, FLUOROURACIL	143	SANDOZ FINASTERIDE	152	SANDOZ RISPERIDONE	87
SALINEX	114	SANDOZ FLUOROMETHOLONE	111	SANDOZ RIVASTIGMINE	27
SALMETEROL XINAFOATE	30	SANDOZ FLUOXETINE	79	SANDOZ RIZATRIPTAN ODT	94
SALMETEROL XINAFOATE, FLUTICASONE PROPIONATE	30	SANDOZ FLUVOXAMINE	80	SANDOZ ROSUVASTATIN	42
SALOFALK	121	SANDOZ FOLIC ACID	147	SANDOZ SERTRALINE	81
SANDOMIGRAN	95	SANDOZ GLICLAZIDE MR	132	SANDOZ SIMVASTATIN	42
SANDOMIGRAN DS	95	SANDOZ INDOMETHACIN	64	SANDOZ SOLIFENACIN	145
SANDOSTATIN	150	SANDOZ IRBESARTAN	57	SANDOZ SUMATRIPTAN	94
SANDOSTATIN LAR	150	SANDOZ IRBESARTAN HCT	57	SANDOZ TACROLIMUS	158
SANDOZ ALENDRONATE	154	SANDOZ LACOSAMIDE	73	SANDOZ TAMSULOSIN	31
SANDOZ	154	SANDOZ LANSOPRAZOLE	119	SANDOZ TELMISARTAN	59
ALENDRONATE/CHOLECALCIFEROL		SANDOZ LATANOPROST	113	SANDOZ TELMISARTAN HCT	59
SANDOZ ALFUZOSIN	31	SANDOZ LATANOPROST/TIMOLOL	113	SANDOZ TIMOLOL	113
SANDOZ ALMOTRIPTAN	93	SANDOZ LEFLUNOMIDE	156	SANDOZ TOBRAMYCIN	110
SANDOZ AMIODARONE	39	SANDOZ LETROZOLE	20	SANDOZ TOLTERODINE LA	145
SANDOZ AMLODIPINE	50	SANDOZ LEVETIRACETAM	74	SANDOZ TOPIRAMATE	76
SANDOZ AMOXI-CLAV	5	SANDOZ LEVOFLOXACIN	6	SANDOZ TRANDOLAPRIL	56
SANDOZ AMPHETAMINE XR	89	SANDOZ LINEZOLID	8	SANDOZ TRAVOPROST	113
SANDOZ ANAGRELIDE	36	SANDOZ LISINOPRIL	53	SANDOZ TRAVOPROST / TIMOLOL PQ	113
SANDOZ ANASTROZOLE	16	SANDOZ LISINOPRIL HCT	54	SANDOZ VALACYCLOVIR	13
SANDOZ ANUZINC HC	140	SANDOZ LOSARTAN	58	SANDOZ VALPROIC	77
SANDOZ ANUZINC HC PLUS	141	SANDOZ LOSARTAN HCT	58	SANDOZ VALSARTAN	60
SANDOZ ARIPIPRAZOLE	82	SANDOZ LOVASTATIN	41	SANDOZ VALSARTAN HCT	60
SANDOZ ATOMOXETINE	97	SANDOZ METFORMIN	129	SANDOZ VENLAFAXINE XR	82
SANDOZ ATORVASTATIN	40	SANDOZ METFORMIN FC	129	SANDOZ VORICONAZOLE	9
SANDOZ AZITHROMYCIN	4	SANDOZ METHYLPHENIDATE SR	90	SANDOZ ZOLMITRIPTAN	95
SANDOZ BICALUTAMIDE	17	SANDOZ METOPROLOL SR	49	SANDOZ ZOLMITRIPTAN ODT	95
SANDOZ BISOPROLOL	47	SANDOZ MINOCYCLINE	7	SANDOZ-CARBAMAZEPINE	72
SANDOZ BOSENTAN	46	SANDOZ MIRTAZAPINE	80	SANDOZ-DICLOFENAC	63
SANDOZ BRIMONIDINE	112	SANDOZ MOMETASONE	111	SANDOZ-DICLOFENAC SR	63
SANDOZ BUPROPION SR	77	SANDOZ MONTELUKAST	107	SANDOZ-FELODIPINE	51
SANDOZ CANDESARTAN	56	SANDOZ MORPHINE SR	68	SANTYL	143
SANDOZ CANDESARTAN PLUS	57	SANDOZ MOXIFLOXACIN	7	SAPHRIS	83
		SANDOZ MYCOPHENOLATE	158		

Non-Insured Health Benefits

SAQUINAVIR MESYLATE	12	SIMVASTATIN-10	43	SPIRIVA	28
SARNA HC	140	SIMVASTATIN-20	43	SPIRIVA RESPIMAT	28
SAXAGLIPTIN HYDROCHLORIDE	130	SIMVASTATIN-40	43	SPIRONOLACTONE	61
SAXAGLIPTIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE	130	SIMVASTATIN-80	43	SPIRONOLACTONE ORAL LIQUID	61
SDZ CELECOXIB	62	SINECATECHINS	137	SPIRONOLACTONE, HYDROCHLOROTHIAZIDE	105
SEASONALE	127	SINEMET	95	SPORANOX	9
SEASONIQUE	127	SINEQUAN	78	STALEVO	96
SEBCUR	142	SINGULAIR	107	STATEX	68
SEBCUR-T	142	SINTROM	34	STAVUDINE	12
SECARIS	114	SIROLIMUS	158	STELARA	151
SECUKINUMAB	143	SITAGLIPTIN PHOSPHATE MONOHYDRATE	130	STERILE EXTEMPORANEOUS MIXTURE (QC)	150
SEEBRI BREEZHALER	28	SITAGLIPTIN PHOSPHATE MONOHYDRATE, METFORMIN HYDROCHLORIDE	130	STERILE WATER	106
SELECT 1/35 (21-DAY)	127	SKIN PREP ADHESIVE WIPES SLOWK	160	STERILE WATER PF	166
SELECT 1/35 (28-DAY)	127	SN IV3000 1-HAND TRANS	103	STEROID AND ANTIFUNGAL CREAM	150
SELEGILINE HYDROCHLORIDE	97	SODIUM AUROTHIOMALATE	123	STIEVA-A	141
SELENIUM SULFIDE	138	SODIUM AUROTHIOMALATE	123	STIVARGA	23
SELEXIPAG	109	SODIUM BICARBONATE	102	STRATTERA	97
SENNA	116	SODIUM BICARBONATE	115	STRESSTABS FOR WOMEN	149
SENNA LAXATIVE	116	SODIUM CARBOXYMETHYL CELLULOSE	114	STRIBILD	12
SENNA SENNOSIDES	116	SODIUM CHLORIDE	103	SUBOXONE	69
SENNA SENNOSIDES NATURALS	116	SODIUM CHLORIDE	103	SUCRALFATE	119
SENNACE	116	SODIUM CHLORIDE (SMALL VOL.)	103	SULCRATE	119
SENNALAX	116	SODIUM CHLORIDE 1G	103	SULCRATE PLUS	119
SENNAPREP	116	SODIUM PHOSPHATE	116	SULFAMETHOXAZOLE, TRIMETHOPRIM	7
SENNOSIDES	116	SODIUM POLYSTYRENE SULFONATE	104	SULFASALAZINE	7
SENNOSIDES	116	SOFOSBUVIR	14	SULFATRIM	7
SEKOKOT	116	SOFOSBUVIR, LEDIPASVIR	14	SULFATRIM DS	7
SEPTA DONEPEZIL	26	SOFOSBUVIR, VELPATASVIR	15	SULFATRIM PEDIATRIC	7
SEPTA-AMLODIPINE	50	SOFOSBUVIR, VELPATASVIR, VOXILAPREVIR	15	SULFINPYRAZONE	105
SEPTA-ATENOLOL	47	SOFRACORT EAR/EYE	111	SULFINPYRAZONE	105
SEPTA-CIPROFLOXACIN	6	SOLIFENACIN	145	SULFUR IN NON-MEDICATED CREAM	150
SEPTA-CITALOPRAM	78	SOLIFENACIN SUCCINATE	145	SULFUR IN NON-MEDICATED OINTMENT	150
SEPTA-LOSARTAN	58	SOLUCAL	102	SULINDAC	65
SEPTA-LOSARTAN HCTZ	58	SOLUCAL D	102	SUMATRIPTAN	94
SEPTA-METFORMIN	129	SOLUCAL D CITRUS	102	SUMATRIPTAN DF	94
SEPTA-ONDANSETRON	117	SOLUCAL D FORT	102	SUMATRIPTAN HEMISULFATE	94
SEPTA-ZOLMITRIPTAN-ODT	95	SOLUCAL D FORT CITRUS	102	SUMATRIPTAN SUCCINATE	94
SERC	98	SOLUCAL D FORT GREEN APPLE	102	SUNITINIB MALATE	24
SEREVENT DISKUS	30	SOLUCAL D RASPBERRY	102	SUNPHARMA ROSUVASTATIN	42
SEROQUEL	86	SOLUCAL GREEN APPLE	102	SUPER-FINE MICRO 31G-5MM NEEDLE	162
SEROQUEL XR	87	SOLUCAL RASPBERRY	102	SUPER-FINE STANDARD 29G-12.7MM	162
SERTRALINE	81	SOLU-CORTEF ACT-O-VIAL	125	SUPER-FINE XTRA 31G-8MM NEEDLE	163
SERTRALINE HYDROCHLORIDE	81	SOLU-MEDROL	125	SUPEUDOL	69
SERTRALINE-100	81	SOLUVER	142	SUPRAX	2
SERTRALINE-25	81	SOLUVER PLUS	142	SUPREFACT	17
SERTRALINE-50	81	SOLYSTAT	104	SUPREFACT (NASAL)	17
SEVELAMER CARBONATE	104	SOMATULINE AUTOGEL	159	SUPREFACT DEPOT 2 MONTHS	17
SEVELAMER HYDROCHLORIDE	104	SOOthe NIGHT TIME	114	SUPREFACT DEPOT 3 MONTHS	17
SHARPS CONTAINER	163	SORBITOL, SODIUM CITRATE, SODIUM LAURYL SULFOACETATE	116	SURE STEP	100
SHARPS NESTABLE YELLOW LARGE 22.7L	163	SORIATANE	142	SURECOMFORT 1/2 IN 28GX0.5CC	164
SIALOR	114	SOTALOL HYDROCHLORIDE	49	SURECOMFORT 1/2 IN 28GX1CC	164
SIDEKICK	100	SOTALOL ORAL LIQUID	49	SURECOMFORT 1/2 IN 29GX0.3CC	164
SILDENAFIL CITRATE	45	SOURCE THICKEN UP 227G	166	SURECOMFORT 1/2 IN 29GX1CC	164
SILIQ	142	SOVALDI	14	SURECOMFORT 1/2 IN 30GX0.3CC	164
SILVER SULFADIAZINE	139	SPACER DEVICE	160	SURECOMFORT 1/2 IN 30GX0.5CC	164
SIMBRINZA	113	SPECTRO ACNECARE WASH	142	SURECOMFORT 1/2 IN 30GX1CC	164
SIMILAC ADVANCE NEOSURE 363G	166	SPECTRO ECZEMACARE	140	SURECOMFORT 29GX1/2 NEEDLE	162
SIMPLY THICK 64OZ BOTTLE PUMP	166	SPIRIT TEST STRIP (ON)	100	SURECOMFORT 30GX5/16 NEEDLE	162
SIMPLY THICK HONEY	166			SURECOMFORT 31GX3/16 NEEDLE	162
SIMPLY THICK NECTAR	166				
SIMPONI	156				
SIMVASTATIN	42				
SIMVASTATIN	42				

Non-Insured Health Benefits

SURECOMFORT 31GX5/16 NEEDLE	162	TARO-TERCONAZOLE	138	TESTOSTERONE UNDECANOATE	126
SURECOMFORT 32GX1/4 NEEDLE	162	TARO-TESTOSTERONE	126	TETRABENAZINE	98
SURECOMFORT 32GX5/32 NEEDLE	162	TARO-WARFARIN	36	TETRABENAZINE	98
SURECOMFORT 5/16 IN 30GX0.3CC	163	TARO-ZOLEDRONIC ACID	155	TETRACYCLINE	7
SURECOMFORT 5/16 IN 30GX0.5CC	163	TASIGNA	21	TETRACYCLINE HYDROCHLORIDE	7
SURECOMFORT 5/16 IN 30GX1CC	163	TAZAROTENE	144	TEVA-5 ASA	121
SURECOMFORT 5/16 IN 31GX0.3CC	163	TAZORAC	144	TEVA-ABACAVIR/LAMIVUDINE	10
SURECOMFORT 5/16 IN 31GX0.5CC	163	TEARS NATURALE FREE	114	TEVA-ACEBUTOLOL	47
SURECOMFORT 5/16 IN 31GX1CC	163	TEARS NATURALE II	114	TEVA-ACYCLOVIR	13
SURETEST (ON)	100	TEARS PLUS	114	TEVA-ALENDRONATE	154
SUSTIVA	11	TEBRAZID	10	TEVA-	154
SUTENT	24	TECFIDERA	98	ALENDRONATE/CHOLECALCIFEROL	
SYMBICORT 100 TURBUHALER	29	TECTA	120	TEVA-ALMOTRIPTAN	93
SYMBICORT 200 TURBUHALER	29	TEGRETOL	72	TEVA-ALPRAZOLAM	90
SYNALAR	140	TELMISARTAN	59	TEVA-AMIODARONE	39
SYNAREL	129	TELMISARTAN	59	TEVA-AMITRIPTYLINE	77
SYNJARDY	132	TELMISARTAN HCTZ	59	TEVA-AMLODIPINE	50
SYNPHASIC 21	128	TELMISARTAN, HYDROCHLOROTHIAZIDE	59	TEVA-AMPICILLIN	5
SYNPHASIC 28	128	TELMISARTAN/HCTZ	59	TEVA-ARIPIPRAZOLE	82
SYNTHROID	134	TELMISARTAN-HCTZ	59	TEVA-ATAZANAVIR	10
SYRINGE & NEEDLE	163	TELMISARTAN-HCTZ	59	TEVA-ATENOLOL	47
SYRINGE CASE	165	TELZIR	11	TEVA-ATENOLOL/CHLOROTHALIDONE	47
SYRINGE SCALE MAGNIFIER	162	TEMAZEPAM	93	TEVA-ATOMOXETINE	97
SYSTANE	114	TEMAZEPAM	93	TEVA-ATORVASTATIN	40
T : SLIM X2 CARTRIDGE (SK)	161	TEMODAL	24	TEVA-AZATHIOPRINE	157
T/ THERAPEUTIC SHAMPOO EXTRA STRENGTH	142	TEMOZOLOMIDE	24	TEVA-AZITHROMYCIN	4
TACROLIMUS (PROTOPIC)	143	TEMPRA CHILDREN'S	69	TEVA-BETAHISTINE	98
TACROLIMUS MONOHYDRATE	158	TEMPRA CHILDREN'S DOUBLE STRENGTH	69	TEVA-BICALUTAMIDE	17
TADALAFIL	46	TEMPRA INFANT	69	TEVA-BISOPROLOL	47
TAFINLAR	18	TENDER-1 17MM/110CM	161	TEVA-BOSENTAN	46
TAGRISSO	22	TENDER-1 17MM/60CM	161	TEVA-BROMAZEPAM	91
TALTZ	143	TENDER-1 17MM/80CM	161	TEVA-BUDESONIDE	125
TAMBOCOR	39	TENDER-1 MINI INF SET 13MM/110CM	161	TEVA-BUSPIRONE	93
TAMOXIFEN CITRATE	24	TENDER-1 MINI INFSET 13MM/60CM	161	TEVA-CANDESARTAN	56
TAMSULOSIN	31	TENDER-1 MINI INFSET 13MM/80CM	161	TEVA-CANDESARTAN/HCTZ	57
TAMSULOSIN HYDROCHLORIDE	31	TENDER-2 17MM/110CM	161	TEVA-CAPECITABINE	17
TAPAZOLE	134	TENDER-2 17MM/60CM	161	TEVA-CAPTROPIL	52
TARCEVA	18	TENDER-2 17MM/80CM	161	TEVA-CARBAMAZEPINE	72
TARGEL	142	TENDER-2 17MM/80CM	161	TEVA-CARVEDILOL	48
TARGEL SA	142	TENDER-2 MINI INF SET 13MM/110CM	161	TEVA-CEFADROXIL	2
TARO-ACITRETIN	142	TENDER-2 MINI INFSET 13MM/60CM	161	TEVA-CEPHALEXIN	3
TARO-AMCINONIDE	139	TENDER-2 MINI INFSET 13MM/80CM	161	TEVA-CHLOROQUINE	15
TARO-ANASTROZOLE	16	TENOFOVIR DISOPROXIL FUMARATE	12	TEVA-CHLORPROMAZINE	83
TARO-BENZOYL PEROXIDE / CLINDAMYCIN KIT	137	TENOFOVIR DISOPROXIL FUMARATE, EMTRICITABINE	12	TEVA-CILAZAPRIL/HCTZ	52
TARO-CAPECITABINE	17	TENOFOVIR DISOPROXIL FUMARATE, EMTRICITABINE, COBICISTAT, ELVITEGRAVIR	12	TEVA-CITALOPRAM	78
TARO-CARBAMAZEPINE	72	TENOFOVIR DISOPROXIL FUMARATE, EMTRICITABINE, RILPIVIRINE HYDROCHLORIDE	12	TEVA-CLARITHROMYCIN	4
TARO-CIPROFLOXACIN	6	TENORETIC	47	TEVA-CLINDAMYCIN	7
TARO-CLARITHROMYCIN	4	TENORMIN	47	TEVA-CLOBAZAM	71
TARO-CLINDAMYCIN	137	TERAZOSIN	46	TEVA-CLOBETASOL	139
TARO-CLINDAMYCIN/BENZOYL PEROXIDE	137	TERAZOSIN HYDROCHLORIDE	46	TEVA-CLONAZEPAM	71
TARO-CLOBETASOL	139	TERBINAFINE	9	TEVA-CLONIDINE	44
TARO-DICLOFENAC	63	TERBINAFINE HYDROCHLORIDE	9	TEVA-CLOPIDOGREL	37
TARO-DIPYRIDAMOLE/ ASA	46	TERBUTALINE SULFATE	30	TEVA-CLOXACILLIN	5
TARO-ENALAPRIL	53	TERCONAZOLE	138	TEVA-CODEINE	66
TARO-FLUCONAZOLE	9	TERIFLUNOMIDE	153	TEVA-COMBO STERINEBS	28
TARO-IMIQUIMOD PUMP	143	TESTIM	126	TEVA-CYCLOBENZAPRINE	31
TARO-MOMETASONE	141	TESTOSTERONE (TOPICAL)	126	TEVA-CYPROTERONE / ETHINYL ESTRADIOL	158
TARO-MUPIROCIN	137	TESTOSTERONE CYPIONATE	126	TEVA-DESMOPRESSIN	133
TARO-PHENYTOIN	71	TESTOSTERONE CYPIONATE	126	TEVA-DICLOFENAC	63
TARO-SONE	139	TESTOSTERONE ENANTHATE	126	TEVA-DICLOFENAC SR	63
TARO-SUMATRIPTAN	94			TEVA-DILTIAZEM	51
TARO-TEMOZOLOMIDE	24			TEVA-DILTIAZEM CD	51
				TEVA-DIVALPROEX	76
				TEVA-DOMPERIDONE	121

Non-Insured Health Benefits

TEVA-DONEPEZIL	26	TEVA-MIRTAZAPINE OD	80	TEVA-TIAPROFENIC	65
TEVA-DOXAZOSIN	46	TEVA-MODAFINIL	90	TEVA-TOBRAMYCIN	2
TEVA-DOXYCYCLINE	7	TEVA-MOMETASONE	111	TEVA-TOLTERODINE	145
TEVA-DUTASTERIDE	152	TEVA-MONTELUKAST	108	TEVA-TOLTERODINE LA	145
TEVA-ECTOSONE	139	TEVA-MORPHINE SR	68	TEVA-TOPILENE	139
TEVA-EFAVIRENZ	11	TEVA-MOXIFLOXACIN	7	TEVA-TOPIRAMATE	76
TEVA-EFAVIRENZ/EMTRICITABINE/TENOFOVIR	11	TEVA-MYCOPHENOLATE	158	TEVA-TOPISONE	139
TEVA-EMTEC-30	65	TEVA-NABILONE	118	TEVA-TRANDOLAPRIL	56
TEVA-EMTRICITABINE/TENOFOVIR	12	TEVA-NAPROXEN	64	TEVA-TRAVOPROST Z	113
TEVA-ENALAPRIL	53	TEVA-NAPROXEN DS	65	TEVA-TRAZODONE	82
TEVA-ENTACAPONE	95	TEVA-NARATRIPTAN	94	TEVA-TRIAMTERENE/HCTZ	105
TEVA-ERLOTINIB	18	TEVA-NITROFURANTOIN	15	TEVA-TRIMEL	7
TEVA-ESCITALOPRAM	79	TEVA-NYSTATIN	9	TEVA-TRIMEL DS	7
TEVA-EXEMESTANE	19	TEVA-OLANZAPINE	84	TEVA-VALACYCLOVIR	13
TEVA-EZETIMIBE	40	TEVA-OMEPRAZOLE	120	TEVA-VALGANCICLOVIR	13
TEVA-FAMOTIDINE	118	TEVA-OXYBUTYNIN	145	TEVA-VALSARTAN	60
TEVA-FENTANYL	66	TEVA-OXYCOCET	65	TEVA-VALSARTAN/HCTZ	60
TEVA-FINASTERIDE	152	TEVA-OXYCODAN	66	TEVA-VENLAFAXINE XR	82
TEVA-FLUCONAZOLE	9	TEVA-PANTOPRAZOLE	120	TEVA-VORICONAZOLE	9
TEVA-FLUOXETINE	79	TEVA-PANTOPRAZOLE MAGNESIUM	120	TEVA-ZOLMITRIPTAN	95
TEVA-FLURBIPROFEN	63	TEVA-PAROXETINE	81	TEVA-ZOLMITRIPTAN OD	95
TEVA-FLUTICASONONE	111	TEVA-PERINDOPRIL	54	TEVETEN	57
TEVA-FLUVASTATIN	41	TEVA-PERINDOPRIL/INDAPAMIDE	54	TEVETEN PLUS	57
TEVA-FOSINOPRIL	53	TEVA-PINDOLOL	49	THE MAGIC BULLET	115
TEVA-FUROSEMIDE	105	TEVA-PIROXICAM	65	THEO ER	146
TEVA-GABAPENTIN	72	TEVA-PRAVASTATIN	41	THEOLAIR	146
TEVA-GEMFIBROZIL	40	TEVA-PRAZOSIN	46	THEOPHYLLINE	146
TEVA-GLICLAZIDE	132	TEVA-PREDNISOLONE	111	THEOPHYLLINE	146
TEVA-GLYBURIDE	132	TEVA-PREDNISONONE	126	THIAMAZOLE	134
TEVA-HALOPERIDOL	84	TEVA-PREGABALIN	75	THIAMJECT	147
TEVA-HYDROCHLOROTHIAZIDE	105	TEVA-PROCTOSONE	140	THIAMINE	147
TEVA-HYDROMORPHONE	67	TEVA-PROFEN	64	THIAMINE HYDROCHLORIDE	147
TEVA-IMATINIB	19	TEVA-PROGESTERONE	134	THICKENING AGENT	166
TEVA-INDOMETHACIN	64	TEVA-PROPRANOLOL	49	THICKENING GEL	166
TEVA-IPRATROPIUM STERINEBS	28	TEVA-QUETIAPINE	86	THIOGUANINE	24
TEVA-IRBESARTAN	57	TEVA-QUETIAPINE XR	87	THIOPROPERAZINE MESYLATE	88
TEVA-IRBESARTAN HCTZ	57	TEVA-RABEPRAZOLE	120	THIOTHIXENE	88
TEVA-KETOCONAZOLE	9	TEVA-RAMIPRIL	55	THRIVE GUM (NS)	32
TEVA-LACOSAMIDE	73	TEVA-RANITIDINE	118	THRIVE NICOTINE LOZENGES	32
TEVA-LACTULOSE	102	TEVA-RISEDRONATE	154	THRIVE NICOTINELL GUM	31
TEVA-LAMIVUDINE/ZIDOVUDINE	11	TEVA-RISPERIDONE	87	THYROGEN	101
TEVA-LAMOTRIGINE	73	TEVA-RIZATRIPTAN ODT	94	THYROID	134
TEVA-LANSOPRAZOLE	119	TEVA-ROSUVASTATIN	42	THYROID	134
TEVA-LATANOPROST	113	TEVA-SALBUTAMOL	30	THYROTROPIN ALFA	101
TEVA-LEFLUNOMIDE	156	TEVA-SALBUTAMOL HFA	30	TIAMOL	140
TEVA-LETROZOLE	20	TEVA-SELEGILINE	97	TIAPROFENIC ACID	65
TEVA-LEVOCARBDOPA	95	TEVA-SERTRALINE	81	TIAZAC	51
TEVA-LEVOFLOXACIN	6	TEVA-SILDENAFIL R	45	TIAZAC XC	52
TEVA-LISINOPRIL (TYPE P)	53	TEVA-SIMVASTATIN	42	TICAGRELOR	37
TEVA-LISINOPRIL (TYPE Z)	53	TEVA-SOLIFENACIN	145	TICLOPIDINE	37
TEVA-LISINOPRIL/HCTZ (TYPE P)	54	TEVA-SPIRONOLACTONE	61	TICLOPIDINE HYDROCHLORIDE	37
TEVA-LISINOPRIL/HCTZ (TYPE Z)	54	TEVA-SPIRONOLACTONE/HCTZ	105	TIMOLOL	49
TEVA-LOPERAMIDE	115	TEVA-SUCRALFATE	119	TIMOLOL MALEATE	49
TEVA-LORAZEPAM	92	TEVA-SULINDAC	65	TIMOLOL MALEATE, BRIMONIDINE TARTRATE	112
TEVA-LOSARTAN	58	TEVA-SUMATRIPTAN	94	TIMOLOL MALEATE, TRAVOPROST	113
TEVA-LOSARTAN/HCTZ	58	TEVA-SUMATRIPTAN DF	94	TIMOLOL MALEATE-EX	112
TEVA-MEDROXYPROGESTERONE	133	TEVA-TAMOXIFEN	24	TIMOPTIC	112
TEVA-MELOXICAM	64	TEVA-TAMSULOSIN	31	TIMOPTIC-XE	113
TEVA-METHYLPHENIDATE	90	TEVA-TELMISARTAN	59	TINACTIN	138
TEVA-METOPROLOL	48	TEVA-TELMISARTAN HCTZ	59	TINACTIN AEROSOL	138
TEVA-MEXILETINE	39	TEVA-TEMAZEPAM	93	TINZAPARIN SODIUM	36
TEVA-MINOCYCLINE	7	TEVA-TENOFOVIR	12	TIOTROPIUM BROMIDE MONOHYDRATE	28
TEVA-MIRTAZAPINE	80	TEVA-TERAZOSIN	46		
		TEVA-TERBINAFINE	9		

Non-Insured Health Benefits

TIPRANAVIR	12	TRIDESILON	140	ULTICARE 5/16 IN 31GX1CC SYRINGE	164
TIVICAY	11	TRIFLUOPERAZINE	88	ULTICARE LOW DEAD SPACE SYRINGE	163
TIZANIDINE	31	TRIFLUOPERAZINE HYDROCHLORIDE	88	ULTILET CLASSIC LANCET	162
TIZANIDINE HYDROCHLORIDE	31	TRIFLURIDINE	110	ULTRA 29G3/10CC	164
TOBI PODHALER	2	TRIHEXYPHENIDYL	95	ULTRA-FINE II 30G.1CC	164
TOBRADEX	111	TRIHEXYPHENIDYL HYDROCHLORIDE	95	ULTRA-FINE II 30GX0.3 CC SYRINGE	164
TOBRAMYCIN	2	TRILEPTAL	74	ULTRAFINE III NEEDLE 31G 8MM	163
TOBRAMYCIN	2	TRIMEBUTINE	28	ULTRAFLEX 1 10MM/110CM	161
TOBRAMYCIN (OPHTHALMIC)	110	TRIMEBUTINE MALEATE	28	ULTRAFLEX 1 10MM/60CM	161
TOBRAMYCIN INHALATION	2	TRIMETHOPRIM	15	ULTRAFLEX 1 10MM/80CM	161
TOBRAMYCINE	2	TRIMETHOPRIM	15	ULTRAFLEX 1 8MM/110CM	161
TOBREX	110	TRIMETHOPRIM ORAL LIQUID	15	ULTRAFLEX 1 8MM/60CM	161
TOCILIZUMAB (IV)	156	TRIMIPRAMINE	82	ULTRAFLEX 1 8MM/80CM	161
TOCILIZUMAB (SC)	156	TRIMIPRAMINE MALEATE	82	ULTRAFLEX 1 8MM/80CM	161
TODAY SPONGE VAGINAL CONTRACEPTIVE	99	TRINIPATCH	45	ULTRAVATE	140
TOFACITINIB CITRATE	156	TRIPTORELIN PAMOATE	25	UMECLIDINIUM BROMIDE	28
TOLNAFTATE	138	TRIQUILAR 21	127	UMECLIDINIUM BROMIDE, VILANTEROL TRIFENATATE	28
TOLOXIN	39	TRIQUILAR 28	127	UNIFINE 29G 12MM NEEDLE	162
TOLTERODINE TARTRATE	145	TRIUMEQ	10	UNIFINE 31G.6MM NEEDLE	162
TOPAMAX	76	TRIZIVIR	10	UNIFINE 31G.8MM NEEDLE	163
TOPICORT	140	TROPICAMIDE	112	UNIFINE PENTIPS 31GX5MM	162
TOPICORT MILD	140	TROSEC	145	UNIPHYL	146
TOPIRAMATE	76	TROSPIUM CHLORIDE	145	UPTRAVI	109
TOPIRAMATE	76	TRUE TRACK	101	UREA	141
TOPIRAMATE ORAL LIQUID	76	TRUETEST	100	UREMOL	141
TOUJEO SOLOSTAR	131	TRUSOPT	113	UREMOL 10	141
TOVIAZ	145	TRUVADA	12	URINE TEST STRIP	101
TRACLEER	46	TUDORZA GENUAIR	27	URISEC 12	141
TRAJENTA	129	TWYNSTA	50	URISEC 22	141
TRAMETINIB	25	TYLENOL	69	URISEC10	141
TRANDATE	48	TYLENOL EXTRA STRENGTH	70	URISPAS	145
TRANDOLAPRIL	56	TYLENOL JR STRENGTH FASTMELTS	70	UROSODIOL ORAL LIQUID	116
TRANEXAMIC ACID	38	TYLENOL JUNIOR STRENGTH	70	URSO	116
TRANEXAMIC ACID	38	TYLENOL WITH CODEINE NO.2	65	URSO DS	116
TRANEXAMIC DENTAL MOUTHWASH	38	TYLENOL WITH CODEINE NO.3	65	URSODIOL	116
TRANSDERMAL NICOTINE	32	ULIPRISTAL ACETATE	128	URSODIOL	116
TRANSDERMAL NICOTINE PATCHDAY	32	ULORIC	152	USTEKINUMAB	151
TRANSDERM-NITRO	45	ULTI SYG 1/2 IN 29GX0.3CC	164	VAGIFEM 10	128
TRANLYCYPROMINE SULFATE	82	ULTI SYG 1/2 IN 29GX0.5CC	164	VALACYCLOVIR	13
TRAVATAN Z	113	ULTI SYG 1/2 IN 29GX1CC SYRINGE	164	VALACYCLOVIR HYDROCHLORIDE	13
TRAVEL	117	ULTI SYG 1/2 IN 30GX0.3CC	164	VALCYTE	13
TRAVEL ON	117	ULTI SYG 1/2 IN 30GX0.5CC	164	VALGANCICLOVIR HYDROCHLORIDE	13
TRAVOPROST	113	ULTI SYG 1/2 IN 30GX1CC SYRINGE	164	VALISONE	139
TRAZODONE	82	ULTI SYG 5/16 IN 30GX0.3CC	164	VALIUM	91
TRAZODONE HYDROCHLORIDE	82	ULTI SYG 5/16 IN 30GX0.5CC	164	VALPROIC ACID (DIVALPROEX SODIUM)	76
TRELSTAR	25	ULTI SYG 5/16 IN 30GX1CC SYRINGE	164	VALPROIC ACID (SODIUM VALPROATE)	77
TRESIBA	131	ULTI SYG 5/16 IN 31GX0.5CC	164	VALSARTAN	59
TRETINOIN	25	ULTI SYG 5/16 IN 31GX1CC SYRINGE	164	VALSARTAN	60
TRIADERM	141	ULTIBRO BREEZHALER	28	VALSARTAN HCT	60
TRIAMCINOLONE	126	ULTICARE 1/2 IN 28GX0.5CC SYRINGE	164	VALSARTAN, HYDROCHLOROTHIAZIDE	60
TRIAMCINOLONE ACETONIDE	111	ULTICARE 1/2 IN 28GX1CC SYRINGE	164	VALSARTAN, SACUBITRIL	61
TRIAMCINOLONE DIACETATE	126	ULTICARE 29GX0.1CC	164	VALSARTAN-HCTZ	60
TRIAMTERENE, HYDROCHLOROTHIAZIDE	105	ULTICARE 29GX0.3CC	164	VALTRESX	13
TRIA TEC-30	65	ULTICARE 29GX0.5CC	164	VAN-ALENDRONATE	154
TRIAZOLAM	93	ULTICARE 29GX12MM PEN NEEDLE	162	VAN-AMLODIPINE	50
TRIAZOLAM	93	ULTICARE 30GX0.1CC	164	VAN-ANASTROZOLE	16
TRICIRA LO 21	128	ULTICARE 30GX0.3CC	164	VAN-BICALUTAMIDE	17
TRICIRA LO 28	128	ULTICARE 30GX0.5CC	164	VANCOCIN	8
TRI-CYCLEN 21-DAY	128	ULTICARE 31GX6MM PEN NEEDLE	162	VANCOMYCIN	8
TRI-CYCLEN 28-DAY	128	ULTICARE 31GX8MM PEN NEEDLE	163	VANCOMYCIN HYDROCHLORIDE	8
TRI-CYCLEN LO (21 DAY)	128	ULTICARE 32GX4MM PEN NEEDLE	163	VANCOMYCIN HYDROCHLORIDE	8
TRI-CYCLEN LO (28 DAY)	128	ULTICARE 5/16 IN 31GX0.3CC SYRINGE	164		
		ULTICARE 5/16 IN 31GX0.5CC SYRINGE	164		

Non-Insured Health Benefits

VANCOMYCIN HYDROCHLORIDE (INJECTION)	8	VISANNE	133	XATRAL	31
VANDETANIB	25	VISKAZIDE	48	XELJANZ	156
VAN-FINASTERIDE	152	VISKEN	49	XELODA	17
VAN-FLUOXETINE	79	VISTITAN	113	XENEX IPECAC	117
VAN-IRBESARTAN	57	VISUDYNE	114	XENEX SODIUM BICARBONATE	102
VAN-LOSARTAN	58	VIT D 1000	148	XEOMIN	159
VAN-MYCOPHENOLATE	158	VIT D 400	148	XGEVA	154
VAN-OMEPRAZOLE	120	VITAMIN A	147	XIGDUO	132
VAN-ONDANSETRON	117	VITAMIN A	147	XOLAIR	109
VAN-PIOGLITAZONE	133	VITAMIN A ACID	141	XTANDI	18
VAN-QUETIAPINE	86	VITAMIN B1	147	XYLAC	84
VAN-RAMIPRIL	55	VITAMIN B12	147	XYLOCAINE VISCOUS	112
VAN-RIZATRIPTAN	94	VITAMIN B12 SUBLINGUAL	147	YASMIN 21	127
VAN-RIZATRIPTAN ODT	94	VITAMIN B6	147	YASMIN 28	127
VAN-ZOLMITRIPTAN ODT	95	VITAMIN C	147	YAZ	127
VARENICLINE TARTRATE	32	VITAMIN C	148	YELLOW HORNET VENOM PROTEIN	151
VASERETIC	53	VITAMIN D	148	YELLOW JACKET VENOM PROTEIN	152
VASOTEC	53	VITAMIN D	148	ZADITEN	1
VCF FOAM VAGINAL CONTRACEPTIVE	99	VITAMIN D3	148	ZAMINE 21	127
VCF VAGINAL CONTRACEPTIVE FILM	99	VITAMIN E	149	ZAMINE 28	127
VEDOLIZUMAB	158	VITAMIN E	149	ZARONTIN	71
VEMURAFENIB	25	VITAMIN K1	149	ZAROXOLYN	105
VENCLEXTA	25	VITAMINE C	148	ZAXINE	8
VENETOCLAX	25	VITAMINE D	148	ZELBORAF	25
VENLAFAXINE HYDROCHLORIDE	82	VOLIBRIS	46	ZELDOX	88
VENLAFAXINE XR	82	VOLTAREN	63	ZENHALE	29
VENOFER	34	VOLTAREN OPHTHA	111	ZEPATIER	14
VENOM PROTEIN EXTRACT	151	VOLTAREN SR	63	ZERIT	12
VENOMIL HONEY BEE VENOM	151	VORICONAZOLE	9	ZESTORETIC	54
VENOMIL MIXED VESPID VENOM PROTEIN	151	VOSEVI	15	ZESTRIL	53
VENOMIL WASP VENOM PROTEIN	151	VOTRIENT	22	ZIAGEN	10
VENOMIL WHITE-FACED HORNET VENOM PROTEIN	151	VPI-ONDANSETRON ODT	118	ZIDOVUDINE	12
VENOMIL YELLOW HORNET VENOM PROTEIN	151	VYVANSE	89	ZINC OXIDE	142
VENOMIL YELLOW JACKET VENOM PROTEIN	152	WAMPOLE CALCIUM	102	ZINC OXIDE	142
VENTOLIN DISKUS	30	WAMPOLE CALCIUM AND D	103	ZINC OXIDE, WHITE PETROLATUM	142
VENTOLIN HFA	30	WAMPOLE CALCIUM VITAMIN D	102	ZINCOFAX EXTRA STRENGTH	142
VENTOLIN P.F	30	WAMPOLE COMPLETE MULT-PRE AND POST NATAL WITH FOLIC ACID	149	ZINDA-LETROZOLE	20
VENTOLIN RESPIRATOR	30	WAMPOLE FERROUS GLUCONATE	34	ZIPRASIDONE HYDROCHLORIDE	88
VEPESID	18	WAMPOLE FOLIC ACID	147	MONOHYDRATE	
VERAPAMIL HYDROCHLORIDE	52	WAMPOLE MINERAL CALCIUM	102	ZITHROMAX	4
VEREGEN	137	WAMPOLE VITAMIN C	148	ZOCOR	43
VERELAN	52	WAMPOLE VITAMIN D	148	ZODERM	143
VERMOX	2	WARFARIN SODIUM	36	ZOFRAN	117
VERSEL	138	WASP VENOM PROTEIN	151	ZOFRAN ODT	118
VERTEPORFIN	114	WATER	105	ZOLADEX	129
VESANOID	25	WEBCOL ALCOHOL PREP	161	ZOLADEX LA	150
VESICARE	145	WELLBUTRIN SR	77	ZOLEDRONIC ACID	155
VESPULA SPP VENOM PROTEIN EXTRACT	151	WELLBUTRIN XL	77	ZOLEDRONIC ACID MONOHYDRATE	155
VFEND	9	WHITE FACED HORNET VENOM PROTEIN	151	ZOLMITRIPTAN	95
VIDEX EC	10	WHITE FACED HORNET VENOM PROTEIN, YELLOW HORNET VENOM PROTEIN, YELLOW JACKET VENOM PROTEIN	151	ZOLMITRIPTAN	95
VIDEXTRA	149	WHITE PETROLATUM	141	ZOLMITRIPTAN ODT	95
VIGABATRIN	77	WHITE PETROLATUM, LANOLIN, MINERAL OIL	114	ZOLOFT	81
VIGAMOX	110	WINPRED	126	ZOMIG	95
VIMPAT	73	XALACOM	113	ZOMIG RAPIMELT	95
VIRACEPT	11	XALATAN	113	ZOSTRIX	143
VIRAMUNE	11	XALKORI	18	ZOSTRIX HP	143
VIRAMUNE XR	11	XANAX	90	ZOVIRAX	13
VIREAD	12	XANAX TS	91	ZUCLOPENTHIXOL ACETATE	88
VIROPTIC	110	XARELTO	36	ZUCLOPENTHIXOL DIHYDROCHLORIDE	88
				ZYBAN	77
				ZYDELIG	19
				ZYKADIA	17
				ZYLOPRIM	152
				ZYMAR	110

Non-Insured Health Benefits

ZYPREXA	84
ZYPREXA ZYDIS	85
ZYTIGA	16
ZYVOXAM	8