



# **P.E.I. Pharmacare Formulary**

<b>Inquiries should be directed to:</b>
PEI Pharmacare Department of Health and Wellness P.O. Box 2000, 20 Fitzroy St. Charlottetown, PEI C1A 7N8

<b>Telephone inquiries should be directed to:</b>	
Patient Eligibility Prescriber Eligibility Medication Eligibility Pharmacy Eligibility Pharmacist Eligibility Claim Inquiries Special Authorization Drug Status Formulary Inquiries	<b>1-902-368-4947 Charlottetown</b> <b>1-877-577-3737 Toll Free in PEI</b> <b>1-902-368-4905 Fax</b>
Technical Support Help Desk for Community Pharmacies	<b>628-3772 Charlottetown</b> <b>1-877-201-6771 Toll Free in PEI</b>  <b>7:00 am to 12:00 midnight</b> <b>7 days per week</b>
PEI Insulin Pump Program Diabetes Glucose Sensor Program Montague Health Center 407 MacIntyre Avenue Montague, PE C0A 1R0	<b>1-902-213-4825 Phone</b> <b>1-833-335-0538 Toll Free in PEI</b> <a href="mailto:diabetesadminofficer@ihis.org">diabetesadminofficer@ihis.org</a> Email

Statements within this document are not intended to override or modify the provisions within an enactment or Minister authority.

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## THE FORMULARY

The Prince Edward Island Pharmacare Formulary is a listing of therapeutically effective medications approved for coverage through the following programs:

HIV Drug Program	High Cost Drug Program
Catastrophic Drug Program	Institutional Pharmacy Program
Community Mental Health Drug Program	Nursing Home Drug Program
Children in Care Program	Phenylketonuria (PKU) Program
Cystic Fibrosis Drug Program	Seniors Drug Program
Diabetes Drug Program	Sexually Transmitted Diseases Program
Erythropoietin Program	Smoking Cessation Drug Program
Family Health Benefit Drug Program	Substance Use Harm Reduction Drug Program
Financial Assistance Drug Program	Transplant Drug Program
Generic Drug Program	Tuberculosis Drug Program
Growth Hormone Drug Program	
Hepatitis Drug Program	

It is compiled on behalf of the Minister of Health and Wellness based upon recommendations from either the Atlantic or Canadian Expert Drug Advisory Committees, or the Joint Oncology Drug Review Committee.

Medications in the Formulary are listed by Therapeutic Categories developed by the American Society of Hospital Pharmacists.

The PEI Pharmacare Formulary is not to be used to determine interchangeability of therapeutic products.

The PEI Pharmacare Formulary may be downloaded from the Department of Health and Wellness website at – [PEI Pharmacare Formulary \(princeedwardisland.ca\)](http://princeedwardisland.ca)

**PRINCE EDWARD ISLAND DRUG PROGRAMS**

<b>Program (Formulary Code)</b>	<b>Beneficiaries</b>	<b>Benefits (Note: A prescription is required for all benefits unless otherwise indicated.)</b>	<b>Fee</b>
<b>Programs Delivered Through Community Retail Pharmacies</b>			
<b>Children-In-Care Program (W)</b>	Persons in temporary or permanent custody of the Director of Child Welfare	All prescription medications. Non-prescription medications approved under the Financial Assistance Program	No fee.
<b>Generic Drug Program (G)</b>	Persons less than 65 years of age with no private drug insurance	Approved generic prescription medications.	Maximum of \$19.95 per prescription.  As part of Prescription Care, copays for eligible medications (identified in the formulary with ⑤) will be \$5 per prescription.
<b>Diabetes Drug Program (D)</b>	Persons eligible for PEI Medicare, diagnosed with diabetes, and registered with the program.	Approved insulin products	\$10.00 per 10 mL vial of insulin. \$20.00 per box of insulin cartridges.  As part of Prescription Care, copays for eligible medications (identified in the formulary with ⑤) will be \$5 per vial or cartridge.
		Approved oral diabetes medications	\$11.00 per prescription.  As part of Prescription Care, copays for eligible medications (identified in the formulary with ⑤) will be \$5 per prescription

Program (Formulary Code)	Beneficiaries	Benefits (Note: A prescription is required for all benefits unless otherwise indicated.)	Fee
		Approved urine testing materials (Diastix and Ketostix – no prescription required)	\$11.00 per prescription
		Blood Glucose test strips. Patients must have used insulin within 150 days (no prescription required).	\$11.00 per dispense. Maximum of 100 strips per 25 days.
		Approved glucagon devices. Patients must have used insulin within 150 days (no prescription required for up to 2 units in 12 months).	\$20.00 per unit  As part of Prescription Care, copays will be \$5 per unit
<b>Financial Assistance Drug Program (W)</b>	Persons eligible under the Social Assistance Act and Regulations.	Approved prescription and non-prescription medications.	No fee.
<b>Family Health Benefit Drug Program (F)</b>	Families (parents, guardians, and children under 25 years of age) eligible for PEI Medicare, with at least one child under 25 years of age who is still attending school full time, and a total annual net family income less than \$24,800, plus \$3,000 for each additional child. Families must apply for coverage on an annual basis and provide income information to the program.	Approved prescription medications.	The pharmacy professional fee for each prescription obtained.  As part of Prescription Care, copays for eligible medications (identified in the formulary with ⑤) will be \$5 per prescription.

<b>Program (Formulary Code)</b>	<b>Beneficiaries</b>	<b>Benefits (Note: A prescription is required for all benefits unless otherwise indicated.)</b>	<b>Fee</b>
<b>High Cost Drug Program (M)</b>	Persons eligible for PEI Medicare and approved for coverage for one or more of the medications included in the program. Patients must apply for coverage on an annual basis and provide income information to the program.	Approved high-cost medications.	An income-based portion of the medication cost plus the pharmacy professional fee for each prescription obtained.
<b>Nursing Home Drug Program (N)</b>	Residents in private nursing homes eligible for coverage under the Social Assistance Act.	Approved prescription and non-prescription medications.	No fee.
<b>Substance Use Harm Reduction Drug Program (L)</b>	Persons eligible for PEI Medicare and assessed by a medical practitioner or nurse practitioner and determined to require treatment for an opioid use disorder or alcohol use disorder	Approved prescription medications.	No fee.
<b>Smoking Cessation Drug Program (Z)</b>	Persons eligible for PEI Medicare and having received smoking cessation counselling through Primary Care. For more information, please visit <a href="http://www.princeedwardisland.ca/quitsmoking">www.princeedwardisland.ca/quitsmoking</a>	12 weeks of approved prescription or non-prescription medications.	No fee.
<b>Seniors Drug Program (S)</b>	Persons eligible for PEI Medicare and 65 years of age or older. Eligibility is effective upon a person becoming 65 years	Approved prescription medications.	First \$8.25 of the medication cost plus the first \$7.69 of the pharmacy dispensing fee for each prescription obtained.

Program (Formulary Code)	Beneficiaries	Benefits (Note: A prescription is required for all benefits unless otherwise indicated.)	Fee
	of age.		As part of Prescription Care, copays for eligible medications (identified in the formulary with ⑤) will be \$5 per prescription
<b>Catastrophic Drug Program (Q)</b>	PEI permanent residents with a PEI Health card whose household members have up to date tax filings and are experiencing out of pocket eligible drug expenses that exceed their annual household limit. Eligible drug expenses are expenses incurred for drugs designated as having coverage under the Catastrophic Drug Program- (Q) listed on the PEI formulary.	Out of pocket costs for eligible drug expenses	This is an income based program. Once an applicant's out of pocket eligible drug expenses exceed the annual household limit the program will cover any further eligible drug expenses in the program year.
<b>Sexually Transmitted Diseases (STD) Program (V)</b>	Persons diagnosed with a sexually transmitted disease or identified contacts of a person diagnosed with a sexually transmitted disease	Approved antibiotics	No fee.
<b>Programs Delivered Through the Provincial Pharmacy</b> <b>Note: Beneficiaries are responsible for arranging for and paying for delivery of medications obtained through the Provincial Pharmacy.</b>			
<b>HIV Drug Program (A)</b>	Persons diagnosed as HIV positive, diagnosed with AIDS, or with a non work related needle-stick injury and no	Approved antiretroviral agents and adjunctive therapies.	No fee.

<b>Program (Formulary Code)</b>	<b>Beneficiaries</b>	<b>Benefits (Note: A prescription is required for all benefits unless otherwise indicated.)</b>	<b>Fee</b>
	private insurance; and registered with the program through the Chief Health Officer.		
<b>Community Mental Health Drug Program (B)</b>	Approved long-term psychiatric patients living in the community.	Approved long- acting injectable antipsychotic medications provided through an approved out- patient psychiatric program.	No fee.
<b>Cystic Fibrosis Drug Program (C)</b>	Persons eligible for PEI Medicare, diagnosed with cystic fibrosis, and who are registered with the program.	Approved prescription and non-prescription medications.	No fee.
<b>Growth Hormone Drug Program (Y)</b>	Children eligible for PEI Medicare, with a proven growth hormone deficiency or Turners Syndrome, and who are registered with the program.	Approved growth hormone supplements.	No fee.
<b>Hepatitis Drug Program (H)</b>	Persons diagnosed with hepatitis.	Approved prescription medications.	No fee
<b>Institutional Pharmacy Program (N)</b>	Residents in government manors.	Approved prescription and non-prescription medications.	No fee.
<b>Phenylketonuria (PKU) Program (P)</b>	Persons eligible for PEI Medicare, diagnosed with phenylketonuria, and who are registered with the program.	Special low protein formula. Up to \$3600 annually for low protein food items.	No fee.
<b>Transplant Drugs Program</b>	Persons eligible for PEI Medicare, who	Approved immunosuppressant	No fee.



<b>Program (Formulary Code)</b>	<b>Beneficiaries</b>	<b>Benefits (Note: A prescription is required for all benefits unless otherwise indicated.)</b>	<b>Fee</b>
(T)	received a bone marrow or solid organ transplant, and are registered with the program.	medications	
<b>Tuberculosis (TB) Drug Program (X)</b>	Persons diagnosed with tuberculosis or who have been in close contact with a person diagnosed with tuberculosis, and who have registered with the program through the Chief Health Officer.	Approved antibiotics	No fee.
<b>Programs Delivered Through Hospitals</b>			
<b>Erythropoietin Program (E)</b>	Persons eligible for PEI Medicare, have been diagnosed with chronic renal failure or are receiving kidney dialysis.	Approved erythropoietin injections	No fee.

**PEI Diabetes Glucose Sensor Program**

<b>Program</b>	<b>Beneficiaries</b>	<b>Benefits (Note: A prescription is required for all benefits unless otherwise indicated.)</b>	<b>Fee</b>
<b>Program Delivered Through Community Retail Pharmacies</b>			
Diabetes Glucose Sensor Program	Persons eligible for PEI Medicare, diagnosed with diabetes, eligible for program enrollment and registered with the program. <a href="http://www.healthpei.ca/glucose-sensor-program">www.healthpei.ca/glucose-sensor-program</a>	Approved diabetes glucose sensors and transmitters.  <a href="#">See Appendix F</a>	This is an income based program. Fee per dispense period is based on household income to a maximum coverage of \$2,400 per program year.

**PEI Diabetes Insulin Pump Program**

Program	Beneficiaries	Benefits	Fee
<p><b>Programs Delivered Through Approved Vendors (who have entered into an agreement with Department of Health and Wellness) <a href="#">See Appendix D For Approved Vendors</a></b></p>			
<p>Insulin Pump Program</p>	<p>Children / Youth up to the age of 25 years living with type 1 diabetes who meet eligibility requirements.</p> <p><a href="http://www.healthpei.ca/insulin-pump">www.healthpei.ca/insulin-pump</a></p>	<p>Insulin pump and pump supplies from the approved vendors list (<a href="#">see appendix D</a>)</p> <p>The following list details the supplies that are eligible for coverage under the PEI Insulin Pump Program:</p> <ul style="list-style-type: none"> <li>• Insulin pump (one pump every 5 years)</li> <li>• Infusion sets (maximum of 140 sets per year)</li> <li>• Reservoirs (maximum of 140 per year)</li> <li>• Site inserts (maximum of one replacement device per year)</li> <li>• Skin adhesive wipes (maximum of 150 per year)</li> <li>• Sterile transparent dressings (maximum of 200 per year)</li> </ul>	<p>An income-based program.</p> <p>Funding through the program varies depending on household income and private health insurance coverage</p>

**PEI Ostomy Supplies Program**

Program	Beneficiaries	Benefits	Fee
<p><b>Programs Delivered Through Ostomy Supply Vendors</b>  <b>See <a href="#">Appendix E</a> For Eligible Supplies</b></p>			
<p>Ostomy Supplies Program</p>	<p>Persons eligible for PEI Medicare, with permanent abdominal ostomies who meet requirements  <a href="#">Ostomy Supplies Program</a></p>	<p>Ostomy supplies (see appendix E for examples)</p> <p>Coverage is in the form of reimbursement, and will be based on the patient’s household income.</p> <p>Coverage is not retroactive. Patients must be enrolled in the Ostomy Supplies Program at the time of ostomy supply purchase to be eligible for reimbursement.</p> <p>The following list details the categories that are eligible for coverage under the PEI Ostomy Supplies Program:</p> <ul style="list-style-type: none"> <li>• Skin wafers</li> <li>• Ostomy pouches</li> <li>• Adhesive removers</li> <li>• Skin barrier wipes</li> <li>• Stoma powders, pastes, and barrier rings</li> <li>• Ostomy belts</li> </ul> <p><a href="#">Appendix E</a></p>	<p>An income-based program.</p> <p>Funding through the program varies depending on household income and private health insurance coverage</p> <p><a href="#">Ostomy Supplies Program</a></p>

## **PRESCRIPTION CARE**

As of June 1<sup>st</sup>, 2023, under the joint federal-provincial *Prescription Care* Initiative, copays for commonly prescribed, eligible medications will be reduced to \$5 for residents covered under Seniors Drug, Family Health Benefit, Generic Drug and Diabetes Drug programs.

Medications eligible for the \$5 copay are identified in the formulary with a ⑤ preceding the non-proprietary or generic name.

## **FORMULARY REVIEW PROCESS**

The coverage of new pharmaceutical products, new dosage forms and new strengths of existing products, and new uses for existing products must be approved on the authority of the Minister of Health and Wellness. The approval is based, in part, upon review by and recommendations received from either the Canadian Expert Drug Advisory Committee (CEDAC), the Atlantic Expert Advisory Committee (AEAC) or the pan-Canadian Oncology Drug Review (pCODR). Prioritization of listing for products is under the direction of the Provincial Drugs and Therapeutics (PD&T) Committee.

The membership of these committees includes practicing physicians, pharmacists, and experts in drug evaluation. They review and evaluate scientific and economic information on new pharmaceutical products and make a recommendation to participating federal, provincial, and territorial government drug programs on whether a drug should be listed as a program benefit, including any conditions and/or criteria for coverage.

The Drug review process involves the following steps:

### **Health Canada Approval**

Before a manufacturer can sell a drug in Canada, they must receive Health Canada approval. Health Canada assesses the drug's safety, efficacy (usually compared to taking no drug at all) and quality of the manufacturing process used to make the drug. When a drug has met all the regulatory requirements, Health Canada issues a Notice of Compliance (NOC) and/or a Drug Identification Number (DIN).

Information on the Health Canada drug review process is available [here](#).

### **Canadian Drug Expert Committee Review**

PEI is a participant in the national Canadian Drug Expert Committee (CDEC) process. CDEC provides participating federal, provincial, and territorial drug benefit programs with a systematic review of the best available clinical evidence, a critique of manufacturer-submitted pharmacoeconomic studies, and a formulary listing recommendation made by the CDEC.

Submissions for new chemical entities, new combination products, and resubmissions related to these products should be filed with the CEDAC Directorate. Information on the CEDAC requirements and procedures are posted at: [www.cadth.ca](http://www.cadth.ca)

### **Pan Canadian Oncology Drug Review**

PEI is a participant in the pan-Canadian Oncology Drug Review (pCODR) process. This process provides participating federal, provincial and territorial drug benefit programs with a systematic review of the best available clinical evidence, and a formulary listing recommendation for oncology medications by an Expert Advisory Committee.

Submissions for new oncology medications and re-submissions related to these products should be directed through this process. For more information on pCODR, please reference the following web site:

<https://cadth.ca/pcodr>

### **Atlantic Common Drug Review**

PEI is a participant in the Atlantic Common Drug Review (ACDR). The ACDR provides the provincial drug benefit programs in New Brunswick, Newfoundland and Labrador, Nova Scotia, and Prince Edward Island with a systematic review of the best available clinical evidence, and a formulary listing recommendation made by the Atlantic Expert Advisory Committee (AEAC), for drugs that do not fall under the mandate of CEDAC or pCODR.

Submissions for new single source products that do not contain new chemical entities, line extensions, new indications for products released prior to CEDAC, and resubmissions for products reviewed prior to CEDAC should be sent to the drug programs within each of the four Atlantic provinces. The Prince Edward Island copy should be sent to:

PEI Pharmacare  
Department of Health and Wellness  
P.O. Box 2000, 20 Fitzroy St.  
Charlottetown, PE C1A 7N8

Products are normally reviewed in the order of receipt of complete submissions. However, there can be exceptions to this. There is no fast tracking of products or pre-NOC reviews.

Information on the ACDR requirements and procedures is available [here](#).

### **pan-Canadian Pharmaceutical Alliance (pCPA)**

Price negotiations are conducted through the pCPA to achieve greater value for publicly funded drug plans. All brand name drugs reviewed through the Canadian Expert Drug Advisory Committee (CEDAC) and pan-Canadian Oncology Drug Review (pCODR) are considered for negotiation. Generic drugs are considered for negotiation through the pCPA Tiered Pricing Framework.

Information on pCPA is available [here](#).

**Provincial Drug and Therapeutics Committee**

Prioritization of listing for products is under the guidance of the PD&T Committee. The prioritized list is based on expert advisory committees' recommendation and other factors such as drug plan mandates, jurisdictional priorities, budget impact, and resources. This list is submitted to the Department of Health and Wellness where final formulary listing decision is under the framework of the Drug Cost Assistance Act and Regulations.

Drug formulary listing decisions for PEI Pharmacare are announced in a Bulletin which is posted on the PEI Pharmacare webpage.

**Biosimilars**

Health Canada authorizes biosimilars for sale. It is recognized that there are no expected clinically meaningful differences in efficacy and safety between a biosimilar and the biologic drug that was already authorized for sale.

When a biosimilar version of a drug is added to the PEI Pharmacare formulary:

- The originator biologic will be delisted after a 12-month period.
- To maintain Pharmacare coverage, a patient must switch to a biosimilar before the originator biologic is delisted.

## **Maximum Reimbursable Price List**

The process for adding medications to the PEI Pharmacare Maximum Reimbursable Price (MRP) list has been revised effective August 1, 2019.

### **Submission Types:**

A manufacturer may file a submission for a generic drug if:

1. The originator brand and strength of the drug is listed on the PEI Pharmacare Formulary.
2. The originator brand of the drug is listed on the PEI Pharmacare Formulary, but not the strength of the generic drug being submitted.
3. The originator brand is not listed but a generic brand of the drug is listed on the PEI Pharmacare Formulary.
4. The generic product was previously listed on the PEI Pharmacare Formulary and was delisted or was withdrawn from the market and is being re-introduced.
5. PEI Pharmacare requests a submission for a generic drug that is being considered for listing.
6. There is a change in DIN for a generic drug that is listed on the PEI Pharmacare Formulary.

A manufacturer must file a new submission for a generic drug if:

7. There is a change in product ownership for a generic product that is listed on the PEI Pharmacare Formulary.

In cases where none of the submission types described above apply, or if there is doubt as to whether a submission should be made, please contact PEI Pharmacare by email at [pharmservices@ihis.org](mailto:pharmservices@ihis.org) for guidance.

### **Submission Requirements (MRP List)**

Submissions filed by manufacturers to have a generic drug product listed on the PEI Pharmacare Formulary must include the requirements outlined below. PEI Pharmacare may request additional information from the manufacturer, Health Canada, or any other source, or take other factors into consideration when reviewing the submission.

The following information must be contained in the submission and should be compiled in the following order:

1. **Cover Letter or Executive Summary.**
  - Indicate which of the submission types is being filed.
  - The names and contact information (email and phone number) for the primary and backup contact(s) who can be contacted regarding the submission. The manufacturer may designate the consultant(s) preparing the submission as primary and/or backup contact(s). Any changes in contacts should be communicated as soon as possible, by emailing [pharmservices@ihis.org](mailto:pharmservices@ihis.org)

- Additional information can be included in the Cover Letter. Please limit this to an explanation of unexpected situations or unusual features of a particular submission. For example, if any strengths of a product listed in the NOC will not be marketed, include this information as a comment in the Cover Letter.
- An electronic signature is acceptable.

2. **Submission Summary Form**

- Include the completed form as part of the whole PDF file and also as a separate attachment in MS Word format.

3. **Copy of the Notice of Compliance (NOC)** issued by Health Canada or, for drug products without a Notice of Compliance, the Drug Notification Form.

4. Copy of the Health Canada approved **Product Monograph**.

5. **Price**

- Indicate the submitted price in the Submission Summary Form.
- Confirm that the price has been submitted to the pan-Canadian Pharmaceutical Alliance (pCPA) Centralized Price Confirmation Process.

6. a) **Drug Notification Form**

b) A signed letter stating that the manufacturer is able to supply the drug product in quantities sufficient to meet the anticipated demand in the province.

7. **A signed letter authorizing unrestricted communication** regarding the drug product between PEI Pharmacare and

- Other federal, provincial, and territorial (F/P/T) drug programs
- F/P/T health authorities and related facilities
- Health Canada
- Patented Medicine Prices Review Board (PMPRB)
- Canadian Agency for Drugs and Technologies in Health (CADTH)

**All submissions for the addition of products to the PEI Pharmacare Maximum Reimbursable Price (MRP) list must be made by email to [pharmservices@ihis.org](mailto:pharmservices@ihis.org)**

The subject of all email submissions must be “MRP List Submission”.

The email must contain the following attachments:

- A single Portable Document Format (PDF) document that contains all the submission requirements with appropriate bookmarks for each component of the submission
- Submission Summary form (in MS Word format)

Submissions must not be made until there is product ready for sale and shipment to PEI pharmacies.

Pre-Notice of Compliance (NOC) submissions will not be accepted.

Products will not be listed until pCPA pricing is received.

Email submissions must not exceed 5 megabytes in size. Submissions may be sent as compressed



“zip” files.

An email confirmation will be sent to manufacturers to notify them that submissions are considered to be complete and to confirm availability and pricing. Questions regarding the submission will also be sent to manufacturers by email.

Submissions will be reviewed by drug program staff.

**Bookmark Names:**

The following are suggested bookmark names:

- Cover Letter
- Submission Summary
- Notice of Compliance (or Product License for NHPs)
- Drug Notification Form
- Product Monograph
- Unrestricted Sharing of Information Letter
- Notification of Changes Letter

## **SUBMISSION REQUIREMENTS (BRAND PRODUCTS)**

**Manufacturers must complete the CDR, pCODR, pCPA, and ACDR process (as applicable) prior to submitting to PEI Pharmacare for consideration of listing a brand drug. All submissions should be made by email only.** The email should contain an attachment in Portable Document Format (PDF) that contains all of the submission requirements with appropriate bookmarks for each component of the submission. Due to technical limitations individual email submissions must not exceed 5 megabytes in size. Submissions may be sent as compressed “zip” files.

### **Submission Requirements**

The following information must be contained in the submission and should be compiled in the following order:

1. Cover Letter or Executive Summary.
  - The names and contact information (email and phone number) for the primary and backup contact(s) who can be contacted regarding the submission. The manufacturer may designate the consultant(s) preparing the submission as primary and/or backup contact(s). Any changes in contacts should be communicated as soon as possible, by emailing [pharmservices@ihis.org](mailto:pharmservices@ihis.org)
  - Additional information can be included in the Cover Letter. Please limit this to an explanation of unexpected situations or unusual features of a particular submission. For example, if any strengths of a product listed in the NOC will not be marketed, include this information as a comment in the Cover Letter.
  - An electronic signature is acceptable.
2. Copy of the Notice of Compliance (NOC) issued by Health Canada.
3. Copy of the Health Canada approved Product Monograph.
4.
  - a) Drug Notification Form
  - b) Current price for all marketed dosage forms and strengths.
5. A signed letter stating that the manufacturer is able to supply the drug product in quantities sufficient to meet the anticipated demand in the province.
6. A signed letter authorizing unrestricted communication regarding the drug product between PEI Pharmacare and
  - a. Other federal, provincial, and territorial (F/P/T) drug programs
  - b. F/P/T health authorities and related facilities
  - c. Health Canada
  - d. Patented Medicine Prices Review Board (PMPRB)
  - e. Canadian Agency for Drugs and Technologies in Health (CADTH)
7. A Budget Impact Analysis (BIA).

### **For More Information**

For more information on the submission process, please contact PEI Pharmacare at [pharmservices@ihis.org](mailto:pharmservices@ihis.org).

## **PRODUCT DELETIONS**

Except where the manufacture of a product is discontinued or approval for sale of a product in Canada is withdrawn, the deletion of products from the Formulary must be approved on the authority of the Minister of Health and Wellness.

## **SPECIAL AUTHORIZATION DRUG STATUS**

Under the HIV, Diabetes, Family Health Benefit, Financial Assistance, High Cost Drugs, Institutional Pharmacy, Nursing Home, Seniors, Transplant, and Catastrophic Drug Programs, certain drug products may be considered for Special Authorization (SA) coverage under the following circumstances:

1. Therapeutic alternatives listed in the Formulary are contraindicated or have been found to be ineffective; or
2. Drugs for which there is no alternative listed in the Formulary.

SA coverage will not be considered for medications that have not yet been reviewed for coverage by the Atlantic Expert Advisory Committee (AEAC), the Canadian Expert Drug Advisory Committee (CEDAC), the pan-Canadian Oncology Drug Review (pCODR) or that have received a negative recommendation from one of these expert advisory committees.

SA coverage will normally only be approved for the treatment of indications and in dosages listed in the official product monograph approved by Health Canada and published in the most recent edition of the Compendium of Pharmaceuticals and Specialities (CPS).

See Appendix A for further detail regarding the SA process.

## **"NO-SUBSTITUTION" PRESCRIPTIONS**

Both generic and brand name products are manufactured under the same standards of good manufacturing practice, and only those brands which meet accepted standards of equivalence are accepted in Prince Edward Island.

Unless special authorization is granted, patients must pay the pharmacy the standard co-pay, plus any cost difference between the brand name requested and the price paid by government for the least expensive generic product.

In cases where a patient experiences problems with a specific brand of medication (e.g. a documented allergy) and has tried all other eligible generic products, a prescriber may apply to PEI Pharmacare for exemption from the cost of the higher cost brand by submitting a completed Special Authorization Request form.

## EXTEMPORANEOUS PREPARATIONS

Extemporaneous preparations are defined as a drug or mixture of drugs prepared or compounded in a pharmacy according to the orders of a prescriber.

To be eligible as a benefit, extemporaneous preparations must:

1. Be specifically tailored to a prescription;
2. Contain one or more medications presently listed as a benefit under the Program for which the person is eligible and all of which are considered a therapeutic benefit in the concentrations and manner used (subject to the review procedure for SA coverage, if deemed appropriate); and
3. Not duplicate the formulation of a manufactured drug product, dilute or alter its formulation, as to result in a product of equivalent therapeutic advantage or one which offers no clear therapeutic advantage relative to a listed benefit.

Claims for extemporaneous preparations are to be submitted electronically using the major ingredient DIN and the appropriate CPhA compound type code.

## EXCLUSIONS

The following are excluded as benefits under PEI Pharmacare:

- All benefits a person is entitled to under any other provincial or federal program (e.g. Workers Compensation, Department of Veterans Affairs, Non-Insured Health Benefits, etc.) or legislation.
- Drugs not authorized for sale and use in Canada (e.g. drugs obtained through Health Canada's Special Access Program, experimental or investigational drugs).
- The following classes of products, except for those specifically listed in the Formulary:
  - Over-the-counter (OTC) or non-prescription medications (some programs)
  - Dietary and nutritional supplements (e.g. Ensure, Boost)
  - Weight loss products
  - Cannabis and cannabis products
  - Soaps, cleansers, and shampoos
  - Oral ergoloid mesylates (i.e. Hydergine)
  - Peripheral vasodilators (e.g. Arlidin)
  - Combination anti-spasmodic/sedative products (e.g. Donnatal, Librax, Stelabid)
  - Combination sedative/analgesic products (e.g. Fiorinal, Tecnal)
  - Allergy serums
  - Products for the treatment of impotence or infertility.
  - Diagnostic agents (except diabetes)
  - Prostheses, medical devices and appliances, and medical supplies, including first aid supplies and syringes

**PRESCRIPTION QUANTITIES**

Maximum days’ supply is based on the negotiated Pharmacy Service Contract between the Province and the PEI Pharmacists’ Association. Coverage is limited to a maximum 90-day supply unless otherwise noted.

Pharmacare limits coverage to a maximum 30-day supply for:

- Narcotic medications, controlled medications, targeted medications, antibiotics, antifungals, sedatives and sleeping pills;
- Non-oral medications;
- Any high cost medication. For this purpose, a high cost medication is defined as a medication eligible for coverage under the High Cost Drug Program;
- Medications requiring a special authorization request for coverage.

Some medications may differ from this day’s supply standard. In this case, the maximum day supply will be noted within the formulary drug listing and, if applicable, the special authorization criteria.

<b>Program</b>	<b>Maximum Allowable Days’ Supply</b>
HIV Drug Program	60
Cystic Fibrosis Drug Program	60 30 - drugs under SA coverage
Diabetes Drug Program	25 – test strips
Diabetes Glucose Sensor Program	28 – Libre sensors, 30 - Dexcom sensors 35 - Medtronic sensors 90 - Dexcom transmitter 365 – Medtronic transmitter
Growth Hormone Drug Program	30
Hepatitis Drug Program	30
High Cost Drug Program	30, unless otherwise specified in criteria for drug(s).
Substance Use Harm Reduction Drug Program	Up to 30 days
Smoking Cessation Drug Program	28 days – OTC Drugs ; 28 days – Prescription drugs
Transplant Drugs Program	60
Tuberculosis Drug Program	60

**COORDINATION OF BENEFITS**

Coordination of benefits allows one to claim under a private insurance plan and PEI Pharmacare for up to the reasonable and customary amount of the covered expense. A claim must be submitted to private insurance before PEI Pharmacare. If the claim is not submitted electronically to the private insurance plan, it should not be adjudicated through PEI Pharmacare electronically. The client may submit a request for reimbursement to Pharmacare after payment through the private insurance plan.

When benefits are coordinated, the eligible cost reimbursed by Pharmacare is based on the initial cost of the prescription, not the cost remaining after payment through the private insurance. If one's private insurance reimburses more than the amount eligible through Pharmacare, the remaining prescription costs are not eligible for coverage through Pharmacare.

**STANDARDIZATION OF PACKAGE SIZES**

In order to ensure claims are paid correctly, please use the following guidelines when calculating quantities for each claim and ensure your cost per unit is correct in your system.

<b>FORM</b>	<b>QUANTITY</b>	<b>FORM</b>	<b>QUANTITY</b>
Aerosols	Per dose	Nasal sprays	Per dose
Capsules	Per capsule	Nebules	Per mL
Creams	Per gram	Ointments	Per gram
Enemas	Per gm/per mL	Oral Contraceptives	Per tablet
Gels	Per gram	Patches	Per patch
Inhalers	Per dose	Powders	Per gram
Insulins (vials, pens, cartridges)	Per mL	Powder injectables	Per vial
Kits	Per kit	Sensor (glucose)	Per
Liquid Injectables	Per mL	Suppositories	Per supp
Liquids	Per mL	Tablets	Per tablet
		Test Strips	Per strip

LEGEND

**08:12.16 ANTIBIOTICS PENICILLINS 1**

**AMOXICILLIN 2**

250MG CAPSULE 3

00406724 4	NOVAMOXIN 5	TEV 6	CFGNQSW 7
00628115	APO-AMOXI	APX	CFGNQSW
02352710	AMOXICILLIN	SNS	CFGNQSW
02388073	AURO-AMOXICILLIN	ARO	CFGNQSW
02401495	AMOXICILLIN	SIV	CFGNQSW
02433060	JAMP-AMOXICILLIN	JPC	CFGNQSW

**CEFPROZIL**

[SEE APPENDIX A](#) FOR SA CRITERIA 8

250MG TABLET

02293528	TARO-CEFPROZIL (SA) 8	RAN	FGNQSW
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**LATANOPROST**

50UG/ML OPHTHALMIC SOLUTION

02231493	XALATAN	UJC	FNQSW
02254786	TEVA-LATANOPROST	TEV	FGNQSW
02296527	APO-LATANOPROST	APX	FGNQSW
02341085	RIVA-LATANOPROST	RIV	FGNQSW
02367335	SANDOZ-LATANOPROST	SDZ	FGNQSW
02373041	GD-LATANOPROST	GMD	FGNQSW
02426935	MED-LATANOPROST	GMP	FGNQSW
02453355	JAMP-LATANOPROST	JPC	FGNQSW
02489570	LATANOPROST	TLG	FGNQSW

Note: The provincial drug programs will only pay for one 2.5 mL bottle of Latanoprost per patient every 30 days. Patients are responsible for the entire prescription cost of any Latanoprost required beyond this. 9

**Legend Key:**

1. Pharmacological-Therapeutic sub-classification
2. Non-proprietary or generic name of the drug.
3. Drug strength and dosage form.
4. Drug Identification Number (DIN) assigned by Health Canada, or a Pseudo-Identification Number (PDIN) assigned by PEI Pharmacare for billing purposes only.
5. Brand name of the drug
6. Three letter identification codes are assigned to each manufacturer. The codes are listed in the Formulary.
7. Drug programs for which the product is considered to be a benefit:

A	HIV Drug Program	N	Nursing Home/Institutional
B	Community Mental Health Drug Program	P	Phenylkentonuria (PKU) Program
C	Cystic Fibrosis Drug Program	Q	Catastrophic Drug Program
D	Diabetes Drug Program	S	Seniors Drug Program
E	Erythropoietin Program	T	Transplant Drug Program
F	Family Health Benefit Drug Program	V	Sexually Transmitted Diseases Program
G	Generic Drug Program	W	Financial Assistance Program / Children-In-Care Program
H	Hepatitis Drug Program	X	Tuberculosis (TB) Drug Program
L	Substance Use Harm Reduction Program	Y	Growth Hormone Program
M	High Cost Drug Program	Z	Smoking Cessation Drug Program

8. This product requires Special Authorization Status (SA) approval (see Appendix A for SA criteria).
9. Special note regarding the product(s) listed in this section.



## **04:00.00 ANTIHISTAMINES**

### **CETIRIZINE**

#### 10MG TABLET

02223554	REACTINE	MCL	NW
02231603	APO-CETIRIZINE	APX	NW
02451778	JAMP-CETIRIZINE	JPC	NW
02517566	CETIRIZINE EXTRA STRENGTH	JPC	NW

#### 20MG TABLET

02315963	PMS-CETIRIZINE	PMS	FGNQSW
02427141	MAR-CETIRIZINE	MAR	FGNQSW
02453363	APO-CETIRIZINE	APX	FGNQSW
02512025	M-CETIRIZINE	MRA	FGNQSW
02515695	CETIRIZINE	SNS	FGNQSW
02517353	JAMP-CETIRIZINE	JPC	FGNQSW
02528681	TEVA-CETIRIZINE	TEV	FGNQSW
02534126	CETIRIZINE	SIV	FGNQSW

### **DIPHENHYDRAMINE HCL**

#### 25MG CAPSULE

00757683	PDP-DIPHENHYDRAMINE	PEN	NW
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#### 50MG CAPSULE

00757691	PDP-DIPHENHYDRAMINE	PEN	NW
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#### 12.5MG/5ML ELIXIR

02019736	BENADRYL	MCL	NW
02298503	DIPHENHYDRAMINE HCL	JPC	NW

#### 50MG/ML INTRAMUSCULAR INJECTION

00596612	DIPHENHYDRAMINE	SDZ	NW
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### **LORATADINE**

#### 10MG TABLET

00782696	CLARITIN	BAY	W
02243880	APO-LORATADINE	APX	W

## **04:04.16 PIPERAZINE DERIVATIVES**

### **FLUNARIZINE HCL**

#### 5MG CAPSULE

02246082	FLUNARIZINE	AAA	FGNQSW
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## **08:08.00 ANTHELMINTICS**

### **MEBENDAZOLE**

100MG TABLET  
00556734 VERMOX JAN FNQW

### **PRAZIQUANTEL**

600MG TABLET  
02230897 BILTRICIDE BAY FNQSW

### **PYRANTEL PAMOATE**

125MG TABLET  
01944363 COMBANTRIN MCL NW

## **08:12.02 ANTIBIOTICS AMINOGLYCOSIDES**

### **GENTAMICIN SULFATE**

80MG/2ML INJECTION SOLUTION (2ML)  
02242652 GENTAMICIN SDZ FGNQSW

### **TOBRAMYCIN**

80MG/2ML INJECTION SOLUTION  
02241210 TOBRAMYCIN INJECTION USP SDZ CFGNQSW  
02502372 TOBRAMYCIN SULFATE STE CFGNQSW  
02533103 TOBRAMYCIN JPC CFGNQSW

## **08:12.04 ANTIBIOTICS ANTIFUNGALS**

### **FLUCONAZOLE**

50MG TABLET  
02236978 TEVA-FLUCONAZOLE TEV AFGNQSW  
02237370 APO-FLUCONAZOLE APX AFGNQSW  
02245292 MYLAN-FLUCONAZOLE MYL AFGNQSW  
02245643 PMS-FLUCONAZOLE PMS AFGNQSW  
02281260 ACT-FLUCONAZOLE TEV AFGNQSW  
02517396 FLUCONAZOLE SNS AFGNQSW  
02534886 FLUCONAZOLE SIV AFGNQSW

100MG TABLET

02236979	TEV-FLUCONAZOLE	TEV	<b>AFGNQSW</b>
02237371	APO-FLUCONAZOLE	APX	<b>AFGNQSW</b>
02245293	MYLAN-FLUCONAZOLE	MYL	<b>AFGNQSW</b>
02245644	PMS-FLUCONAZOLE	PMS	<b>AFGNQSW</b>
02281279	ACT-FLUCONAZOLE	TEV	<b>AFGNQSW</b>
02517418	FLUCONAZOLE	SNS	<b>AFGNQSW</b>
02534894	FLUCONAZOLE	SIV	<b>AFGNQSW</b>

150MG TABLET

02141442	DIFLUCAN ONE	PFI	<b>AFNQSW</b>
02241895	APO-FLUCONAZOLE	APX	<b>AFGNQSW</b>
02428792	MAR-FLUCONAZOLE	MAR	<b>AFGNQSW</b>
02432471	JAMP-FLUCONAZOLE	JPC	<b>AFGNQSW</b>
02521229	FLUCONAZOLE-150	SNS	<b>AFGNQSW</b>

**ISAVUCONAZOLE**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE

02483971	CRESEMBA (SA)	AVI	<b>NMQW</b>
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**ITRACONAZOLE**

100MG CAPSULE

02047454	SPORANOX	JAN	<b>FNQSW</b>
02462559	MINT-ITRACONAZOLE	MNT	<b>FGNQSW</b>

10MG/ML ORAL SOLUTION

02495988	ODAN ITRACONAZOLE (SA)	ODN	<b>FGNQSW</b>
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**KETOCONAZOLE**

200MG TABLET

02231061	TEVA-KETOCONAZOLE	TEV	<b>AFGNQSW</b>
02237235	APO-KETOCONAZOLE	APX	<b>AFGNQSW</b>

**08:12.06 ANTIBIOTICS CEPHALOSPORINS**

**CEFADROXIL**

500MG CAPSULE

02235134	TEVA-CEFADROXIL	TEV	<b>FGNQSW</b>
02240774	APO-CEFADROXIL	APX	<b>FGNQSW</b>
02544792	JAMP-CEFADROXIL	JPC	<b>FGNQSW</b>

**CEFIXIME**

400MG TABLET

00868981 SUPRAX  
02432773 AURO-CEFIXIMEODN **FNQSVW**  
ARO **FGNQSVW**

100MG/5ML ORAL SUSPENSION

00868965 SUPRAX  
02468689 AURO-CEFIXIMEODN **FNQSW**  
ARO **FGNQSW****CEFPROZIL**

250MG TABLET

02293528 TARO-CEFPROZIL

TAR **FGNQSW**

500MG TABLET

02293536 TARO-CEFPROZIL  
02347253 AURO-CEFPROZILTAR **FGNQSW**  
ARO **FGNQSW**

25MG/ML ORAL SUSPENSION

02329204 TARO-CEFPROZIL  
02347261 AURO-CEFPROZILTAR **FGNQSW**  
ARO **FGNQSW**

50MG/ML ORAL SUSPENSION

02293579 TARO-CEFPROZIL  
02347288 AURO-CEFPROZILTAR **FGNQSW**  
ARO **FGNQSW****CEFTRIAZONE**

1.0G/VIAL INTRAMUSCULAR INJECTION

02287633 CEFTRIAZONE FOR SODIUM  
02292270 CEFTRIAZONE  
02325616 CEFTRIAZONE SODIUMTEV **NQ**  
SDZ **NQ**  
STE **NQ****CEFUROXIME AXETIL**

250MG TABLET

02244393 APO-CEFUROXIME  
02344823 AURO-CEFUROXIMEAPX **CFGNQSW**  
ARO **CFGNQSW**

500MG TABLET

02244394 APO-CEFUROXIME  
02344831 AURO-CEFUROXIMEAPX **CFGNQSW**  
ARO **CFGNQSW**

25MG/ML ORAL SUSPENSION

02212307 CEFTIN

SDZ **CFNQSW****CEPHALEXIN MONOHYDRATE**

250MG CAPSULE

00342084 TEVA-CEPHALEXIN

TEV **CFGNQSW**

500MG CAPSULE				
00342114	TEVA-CEPHALEXIN	TEV	CFGNQSW	
250MG TABLET				
00583413	TEVA-CEPHALEXIN	TEV	CFGNQSW	
00768723	APO-CEPHALEX	APX	CFGNQSW	
02470578	AURO-CEPHALEXIN	ARO	CFGNQSW	
02494698	JAMP-CEPHALEXIN	JPC	CFGNQSW	
02521253	CEPHALEXIN	SNS	CFGNQSW	
500MG TABLET				
00583421	TEVA-CEPHALEXIN	TEV	CFGNQSW	
00768715	APO-CEPHALEX	APX	CFGNQSW	
02470586	AURO-CEPHALEXIN	ARO	CFGNQSW	
02494701	JAMP-CEPHALEXIN	JPC	CFGNQSW	
02495651	CEPHALEXIN	SIV	CFGNQSW	
02521261	CEPHALEXIN	SNS	CFGNQSW	
25MG/ML ORAL SUSPENSION				
00342106	TEVA-CEPHALEXIN	TEV	CFGNQSW	
02469170	LUPIN-CEPHALEXIN	LUP	CFGNQSW	
02497743	AURO-CEPHALEXIN	ARO	CFGNQSW	
02528436	JAMP-CEPHALEXIN	JPC	CFGNQSW	
50MG/ML ORAL SUSPENSION				
00342092	TEVA-CEPHALEXIN	TEV	CFGNQSW	
02469189	LUPIN-CEPHALEXIN	LUP	CFGNQSW	
02497751	AURO-CEPHALEXIN	ARO	CFGNQSW	
02528444	JAMP-CEPHALEXIN	JPC	CFGNQSW	

### **08:12.07 MONOBACTAMS**

#### **AZTREONAM**

[SEE APPENDIX A](#) FOR SA CRITERIA

75MG/ML INHALATION VIAL

02329840	CAYSTON (SA)	GIL	NMQW
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### **08:12.12 ANTIBIOTICS ERYTHROMYCINS**

## **AZITHROMYCIN**

[SEE APPENDIX A](#) FOR SA CRITERIA (HIV, CYSTIC FIBROSIS, SEXUALLY TRANSMITTED DISEASES AND TUBERCULOSIS DO NOT REQUIRE A SA REQUEST)

### **250MG TABLET**

02212021	ZITHROMAX (SA)	PFI	<b>ACFNQSWVX</b>
02261634	PMS-AZITHROMYCIN (SA)	PMS	<b>ACFGNQSWVX</b>
02265826	SANDOZ AZITHROMYCIN (SA)	SDZ	<b>ACFGNQSWVX</b>
02267845	TEVA-AZITHROMYCIN (SA)	TEV	<b>ACFGNQSWVX</b>
02275309	RIVA-AZITHROMYCIN (SA)	RIV	<b>ACFGNQSWVX</b>
02330881	AZITHROMYCIN (SA)	SNS	<b>ACFGNQSWVX</b>
02415542	APO-AZITHROMYCIN Z (SA)	APX	<b>ACFGNQSWVX</b>
02442434	AZITHROMYCIN (SA)	SIV	<b>ACFGNQSWVX</b>
02452308	JAMP-AZITHROMYCIN (SA)	JPC	<b>ACFGNQSWVX</b>
02452324	MAR-AZITHROMYCIN (SA)	MAR	<b>ACFGNQSWVX</b>
02479680	NRA-AZITHROMYCIN (SA)	NRA	<b>ACFGNQSWVX</b>
02480700	AG-AZITHROMYCIN (SA)	AGP	<b>ACFGNQSWVX</b>
02502038	M-AZITHROMYCIN (SA)	MRA	<b>ACFGNQSWVX</b>

### **600MG TABLET**

02261642	PMS-AZITHROMYCIN (SA)	PMS	<b>ACFGNQSWX</b>
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### **20MG/ML ORAL SUSPENSION**

02223716	ZITHROMAX (SA)	PFI	<b>ACFNQSWX</b>
02332388	SANDOZ-AZITHROMYCIN (SA)	SDZ	<b>ACFGNQSWX</b>
02482363	AURO-AZITHROMYCIN (SA)	ARO	<b>ACFGNQSWX</b>

### **40MG/ML ORAL SUSPENSION**

02223724	ZITHROMAX (SA)	PFI	<b>ACFNQSWX</b>
02332396	SANDOZ-AZITHROMYCIN (SA)	SDZ	<b>ACFGNQSWX</b>
02482371	AURO-AZITHROMYCIN (SA)	ARO	<b>ACFGNQSWX</b>

## **CLARITHROMYCIN**

### **250MG TABLET**

01984853	BIAXIN	BGP	<b>AFCNQSWX</b>
02247573	PMS-CLARITHROMYCIN	PMS	<b>ACFGNQSWX</b>
02266539	SANDOZ CLARITHROMYCIN	SDZ	<b>ACFGNQSWX</b>
02274744	APO-CLARITHROMYCIN	APX	<b>ACFGNQSWX</b>
02361426	RAN-CLARITHROMYCIN	RAN	<b>ACFGNQSWX</b>
02442469	CLARITHROMYCIN	SIV	<b>ACFGNQSWX</b>
02466120	CLARITHROMYCIN	SNS	<b>ACFGNQSWX</b>
02471388	M-CLARITHROMYCIN	MRA	<b>ACFGNQSWX</b>

### **500MG TABLET**

02126710	BIAXIN	BGP	<b>ACFNQSWX</b>
02247574	PMS-CLARITHROMYCIN	PMS	<b>ACFGNQSWX</b>
02266547	SANDOZ-CLARITHROMYCIN	SDZ	<b>ACFGNQSWX</b>

02274752	APO-CLARITHROMYCIN	APX	ACFGNQSWX
02361434	RAN-CLARITHROMYCIN	RAN	ACFGNQSWX
02442485	CLARITHROMYCIN	SIV	ACFGNQSWX
02466139	CLARITHROMYCIN	SNS	ACFGNQSWX
02471396	M-CLARITHROMYCIN	MRA	ACFGNQSWX
500MG EXTENDED-RELEASE TABLET			
02403196	ACT-CLARITHROMYCIN XL	TEV	CFGNQSW
02413345	APO-CLARITHROMYCIN XL	APX	CFGNQSW
25MG/ML ORAL SUSPENSION			
02146908	BIAXIN	BGP	CFNQSWX
02390442	TARO-CLARITHROMYCIN	TAR	CFGNQSWX
50MG/ML ORAL SUSPENSION			
02244641	BIAXIN	BGP	CFNQSWX
02390450	TARO-CLARITHROMYCIN	TAR	CFGNQSWX
<b>ERYTHROMYCIN BASE</b>			
250MG TABLET			
00682020	ERYTHRO-BASE	AAA	CFGNQSVW
<b>FIDAXOMICIN</b>			
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
200MG TABLET			
02387174	DIFICID (SA)	MSD	FNQSW

## **08:12.16 ANTIBIOTICS PENICILLINS**

### **AMOXICILLIN**

#### 250MG CAPSULE

00406724	NOVAMOXIN	TEV	CFGNQSW
00628115	APO-AMOXI	APX	CFGNQSW
02388073	AURO-AMOXICILLIN	ARO	CFGNQSW
02433060	JAMP-AMOXICILLIN	JPC	CFGNQSW
02525348	AMOXICILLIN	SNS	CFGNQSW
02532042	PRZ-AMOXICILLIN	PRZ	CFGNQSW

#### 500MG CAPSULE

00406716	NOVAMOXIN	TEV	CFGNQSVW
00628123	APO-AMOXI	APX	CFGNQSVW
02388081	AURO-AMOXICILLIN	ARO	CFGNQSVW
02401509	AMOXICILLIN	SIV	CFGNQSVW
02433079	JAMP-AMOXICILLIN	JPC	CFGNQSVW

02477726	AG-AMOXICILLIN	ANG	CFGNQSVW
02525356	AMOXICILLIN	SNS	CFGNQSVW
02532050	PRZ-AMOXICILLIN	PRZ	CFGNQSW

25MG/ML ORAL SUSPENSION

00628131	APO-AMOXI	APX	CFGNQSW
02458586	AURO-AMOXICILLIN	ARO	CFGNQSW
02535793	JAMP-AMOXICILLIN	JPC	CFGNQSW

50MG/ML ORAL SUSPENSION

00452130	NOVAMOXIN	TEV	CFGNQSW
00628158	APO-AMOXI	APX	CFGNQSW
01934163	NOVAMOXIN	TEV	CFGNQSW
02352788	AMOXICILLIN SUGAR REDUCED	SNS	CFGNQSW
02352753	AMOXICILLIN	SNS	CFGNQSW
02401541	AMOXICILLIN	SIV	CFGNQSW
02458594	AURO-AMOXICILLIN	ARO	CFGNQSW
02495864	SANDOZ-AMOXICILLIN	SDZ	CFGNQSW
02535815	JAMP-AMOXICILLIN	JPC	CFGNQSW

**AMOXICILLIN & CLAVULANIC ACID**

250MG & 125MG TABLET

02243350	APO-AMOXI CLAV	APX	CFGNQSW
02471671	AURO-AMOXICLAV	ARO	CFGNQSW
02508249	JAMP-AMOXI CLAV	JPC	CFGNQSW

500MG & 125MG TABLET

01916858	CLAVULIN	GSK	CFNQSW
02243351	APO-AMOXI CLAV	APX	CFGNQSW
02471698	AURO-AMOXICLAV	ARO	CFGNQSW
02482576	SANDOZ-AMOXI CLAV	SDZ	CFGNQSW
02508257	JAMP-AMOXI CLAV	JPC	CFGNQSW
02536021	AMOXICILLIN-CLAV	SNS	CFGNQSW

875MG & 125MG TABLET

02238829	CLAVULIN	GSK	CFNQSW
02245623	APO-AMOXI CLAV	APX	CFGNQSW
02471701	AURO-AMOXICLAV	ARO	CFGNQSW
02482584	SANDOZ-AMOXI CLAV	SDZ	CFGNQSW
02508265	JAMP-AMOXI CLAV	JPC	CFGNQSW
02536048	AMOXICILLIN-CLAV	SNS	CFGNQSW

25MG & 6.25MG/ML ORAL SUSPENSION

01916882	CLAVULIN	GSK	CFNQSW
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50MG & 12.5MG/ML ORAL SUSPENSION



01916874	CLAVULIN	GSK	CFNQSW
02542226	M-AMOXI CLAV	MRA	CFGNQSW
02539438	JAMP-AMOXI CLAV	JPC	CFGNQSW

80MG & 11.4MG/ML ORAL SUSPENSION

02238830	CLAVULIN	GSK	CFNQSW
02530694	M-AMOXI CLAV	MRA	CFGNQSW
02539446	JAMP-AMOXI CLAV	JPC	CFGNQSW

**AMPICILLIN**

250MG CAPSULE			
00020877	TEVA-AMPICILLIN	TEV	CFGNQSW

500MG CAPSULE			
00020885	TEVA-AMPICILLIN	TEV	CFGNQSW

500MG INJECTION POWDER			
00872652	AMPICILLIN SODIUM	TEV	NQ

**CLOXACILLIN**

250MG CAPSULE			
00337765	TEVA-CLOXACILLIN	TEV	CFGNQSW
02510731	JAMP-CLOXACILLIN	JPC	CFGNQSW

500MG CAPSULE			
00337773	TEVA-CLOXACILLIN	TEV	CFGNQSW
02510758	JAMP-CLOXACILLIN	JPC	CFGNQSW

25MG/ML ORAL LIQUID			
00337757	TEVA-CLOXACILLIN	TEV	CFGNQSW

**PENICILLIN V (POTASSIUM)**

300MG TABLET			
00642215	PEN-VK	AAA	CFGNQSW

**8:12.18 QUINOLONES**

**CIPROFLOXACIN**

[SEE APPENDIX A](#) FOR SA CRITERIA (CYSTIC FIBROSIS, NURSING HOME, AND TUBERCULOSIS PROGRAMS DO NOT REQUIRE AN SA REQUEST)

250MG TABLET			
02247339	ACT-CIPROFLOXACIN (SA)	TEV	CFGNQSWX
02248437	PMS-CIPROFLOXACIN (SA)	PMS	CFGNQSWX

02248756	SANDOZ CIPROFLOXACIN (SA)	SDZ	CFGNQSWX
02303728	RAN-CIPROFLOXACIN (SA)	RAN	CFGNQSWX
02353318	CIPROFLOXACIN (SA)	SNS	CFGNQSWX
02379686	MAR-CIPROFLOXACIN (SA)	MAR	CFGNQSWX
02380358	JAMP-CIPROFLOXACIN (SA)	JPC	CFGNQSWX
02381907	AURO-CIPROFLOXACIN (SA)	ARO	CFGNQSWX
02386119	CIPROFLOXACIN (SA)	SIV	CFGNQSWX

**500MG TABLET**

02247340	ACT-CIPROFLOXACIN (SA)	TEV	CFGNQSWX
02248438	PMS-CIPROFLOXACIN (SA)	PMS	CFGNQSWX
02248757	SANDOZ CIPROFLOXACIN (SA)	SDZ	CFGNQSWX
02303736	RAN-CIPROFLOXACIN (SA)	RAN	CFGNQSWX
02353326	CIPROFLOXACIN (SA)	SNS	CFGNQSWX
02379694	MAR-CIPROFLOXACIN (SA)	MAR	CFGNQSWX
02380366	JAMP-CIPROFLOXACIN (SA)	JPC	CFGNQSWX
02381923	AURO-CIPROFLOXACIN (SA)	ARO	CFGNQSWX
02386127	CIPROFLOXACIN (SA)	SIV	CFGNQSWX
02423561	MINT-CIPROFLOX (SA)	MNT	CFGNQSWX
02492008	NRA-CIPROFLOXACIN (SA)	NRA	CFGNQSWX

**750MG TABLET**

02247341	ACT-CIPROFLOXACIN (SA)	TEV	CFGNQSW
02248439	PMS-CIPROFLOXACIN (SA)	PMS	CFGNQSW
02248758	SANDOZ-CIPROFLOXACIN (SA)	SDZ	CFGNQSW
02303744	RAN-CIPROFLOXACIN (SA)	RAN	CFGNQSW
02379708	MAR-CIPROFLOXACIN (SA)	MAR	CFGNQSW
02380374	JAMP-CIPROFLOXACIN (SA)	JPC	CFGNQSW

**100MG/ML ORAL SUSPENSION**

02237514	CIPRO (SA)	BAY	CFNQSW
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**LEVOFLOXACIN**

[SEE APPENDIX A](#) FOR SA CRITERIA (CYSTIC FIBROSIS AND NURSING HOME PROGRAMS DO NOT REQUIRE AN SA REQUEST)

**250MG TABLET**

02284707	APO-LEVOFLOX (SA)	APX	CFGNQSW
02298635	SANDOZ-LEVOFLOXACIN (SA)	SDZ	CFGNQSW
02315424	ACT-LEVOFLOXACIN (SA)	TEV	CFGNQSW
02505797	MINT-LEVOFLOXACIN (SA)	MNT	CFGNQSW
02508443	JAMP-LEVOFLOXACIN (SA)	JPC	CFGNQSW

**500MG TABLET**

02284715	APO-LEVOFLOX (SA)	APX	CFGNQSW
02298643	SANDOZ-LEVOFLOXACIN (SA)	SDZ	CFGNQSW
02315432	ACT-LEVOFLOXACIN (SA)	TEV	CFGNQSW

02505819	MINT-LEVOFLOXACIN (SA)	MNT	CFGNQSW
02508451	JAMP-LEVOFLOXACIN (SA)	JPC	CFGNQSW
750MG TABLET			
02315440	ACT-LEVOFLOXACIN (SA)	TEV	CFGNQSW
02325942	APO-LEVOFLOXACIN (SA)	APX	CFGNQSW
02508478	JAMP-LEVOFLOXACIN (SA)	JPC	CFGNQSW

### MOXIFLOXACIN HCL

[SEE APPENDIX A](#) FOR SA CRITERIA (CYSTIC FIBROSIS PROGRAM DOES NOT REQUIRE AN SA REQUEST)

400MG TABLET

02375702	TEVA-MOXIFLOXACIN (SA)	TEV	CFGNQSW
02383381	SANDOZ-MOXIFLOXACIN (SA)	SDZ	CFGNQSW
02404923	APO-MOXIFLOXACIN (SA)	APX	CFGNQSW
02432242	AURO-MOXIFLOXACIN (SA)	ARO	CFGNQSW
02443929	JAMP-MOXIFLOXACIN (SA)	JPC	CFGNQSW
02447053	MAR-MOXIFLOXACIN (SA)	MAR	CFGNQSW
02447061	JAMP-MOXIFLOXACIN (SA)	JPC	CFGNQSW
02472791	M-MOXIFLOXACIN (SA)	MRA	CFGNQSW
02478137	AG-MOXIFLOXACIN (SA)	ANG	CFGNQSW
02520710	MOXIFLOXACIN (SA)	SNS	CFGNQSW

### NORFLOXACIN

[SEE APPENDIX A](#) FOR SA CRITERIA (NURSING HOME PROGRAM DOES NOT REQUIRE AN SA REQUEST)

400MG TABLET

02229524	NORFLOXACIN (SA)	AAA	FGNQSW
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## 08:12.20 SULFONAMIDES

### SULFAMETHOXAZOLE & TRIMETHOPRIM

100MG & 20MG TABLET

00445266	SULFATRIM PEDIATRIC	AAA	ACFGNQSWX
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400MG & 80MG TABLET

00445274	SULFATRIM	AAA	ACFGNQSWX
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800MG & 160MG TABLET

00445282	SULFATRIM DS	AAA	ACFGNQSWX
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40MG & 8MG/ML ORAL SUSPENSION

00726540	TEVA-TRIMEL	TEV	ACFGNQSWX
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**08:12.24 ANTIBIOTICS TETRACYCLINES**

**DOXYCYCLINE**

100MG CAPSULE

00725250	TEVA-DOXYCYCLINE	TEV	CFGNQSVWX
00740713	APO-DOXY	APX	CFGNQSVWX
02351234	DOXYCYCLINE	SNS	CFGNQSVWX
02528940	JAMP-DOXYCYCLINE	JPC	CFGNQSVWX

100MG TABLET

00860751	DOXYCIN	RIV	CFGNQSVWX
00874256	APO-DOXY	APX	CFGNQSVWX
02158574	TEVA-DOXYCYCLINE	TEV	CFGNQSVWX
02351242	DOXYCYCLINE	SNS	CFGNQSVWX
02536250	PRZ-DOXYCYCLINE	PRZ	CFGNQSVWX
02543478	M-DOXYCYCLINE	MRA	CFGNQSVWX

**MINOCYCLINE HCL**

50MG CAPSULE

02084090	MINOCYCLINE	AAA	FGNQSW
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100 MG CAPSULE

02084104	MINOCYCLINE	AAA	FGNQSW
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**TETRACYCLINE**

250MG CAPSULE

00580929	TETRACYCLINE	AAA	CFGNQSW
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**8:12.28 ANTIBIOTICS OTHER ANTIBIOTICS**

**CLINDAMYCIN HCL**

150MG CAPSULE

02241709	TEVA-CLINDAMYCIN	TEV	CFGNQSW
02245232	APO-CLINDAMYCIN	APX	CFGNQSW
02400529	CLINDAMYCIN	SNS	CFGNQSW
02436906	AURO-CLINDAMYCIN	ARO	CFGNQSW
02462656	MED-CLINDAMYCIN	GMP	CFGNQSW
02468476	RIVA-CLINDAMYCIN	RIV	CFGNQSW
02479923	M-CLINDAMYCIN	MRA	CFGNQSW
02483734	JAMP-CLINDAMYCIN	JPC	CFGNQSW
02493748	NRA-CLINDAMYCIN	NRA	CFGNQSW

**300MG CAPSULE**

02182866	DALACIN C	PFI	CFNQSW
02241710	TEVA-CLINDAMYCIN	TEV	CFGNQSW
02245233	APO-CLINDAMYCIN	APX	CFGNQSW
02400537	CLINDAMYCIN	SNS	CFGNQSW
02436914	AURO-CLINDAMYCIN	ARO	CFGNQSW
02462664	MED-CLINDAMYCIN	GMP	CFGNQSW
02468484	RIVA-CLINDAMYCIN	RIV	CFGNQSW
02479931	M-CLINDAMYCIN	MRA	CFGNQSW
02483742	JAMP-CLINDAMYCIN	JPC	CFGNQSW
02493756	NRA-CLINDAMYCIN	NRA	CFGNQSW

**CLINDAMYCIN PALMITATE HCL**

15MG/ML ORAL SOLUTION

00225851	DALACIN C	PFI	FNQSW
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**LINEZOLID**

[SEE APPENDIX A](#) FOR SA CRITERIA

600MG TABLET

02422689	SANDOZ-LINEZOLID (SA)	SDZ	FGNQSW
02426552	APO-LINEZOLID (SA)	APX	FGNQSW
02520354	JAMP-LINEZOLID (SA)	JPC	FGNQSW

**RIFAXIMIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

550MG TABLET

02410702	ZAXINE (SA)	LUP	FNQSW
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**VANCOMYCIN HCL**

125MG CAPSULE

00800430	VANCOCIN	MRS	FNQSW
02407744	JAMP-VANCOMYCIN	JPC	FGNQSW

**08:14.00 ANTIFUNGALS**

**NYSTATIN**

100,000U/ML ORAL SUSPENSION

00792667	PMS-NYSTATIN	PMS	AFGNQSW
02194201	RATIO-NYSTATIN	TEV	AFGNQSW
02433443	JAMP-NYSTATIN	JPC	AFGNQSW

**TERBINAFINE**

250MG TABLET

02239893	APO-TERBINAFINE	APX	<b>AFGNQSW</b>
02254727	ACT-TERBINAFINE	TEV	<b>AFGNQSW</b>
02294273	PMS-TERBINAFINE	PMS	<b>AFGNQSW</b>
02320134	AURO-TERBINAFINE	ARO	<b>AFGNQSW</b>
02353121	TERBINAFINE	SNS	<b>AFGNQSW</b>
02385279	TERBINAFINE	SIV	<b>AFGNQSW</b>

**VORICONAZOLE**[SEE APPENDIX A](#) FOR SA CRITERIA

50MG TABLET

02256460	VFEND (SA)	PFI	<b>FNQSW</b>
02396866	TEVA-VORICONAZOLE (SA)	TEV	<b>FGNQSW</b>
02399245	SANDOZ-VORICONAZOLE (SA)	SDZ	<b>FGNQSW</b>
02525771	JAMP-VORICONAZOLE (SA)	JPC	<b>FGNQSW</b>

200MG TABLET

02256479	VFEND (SA)	PFI	<b>FNQSW</b>
02396874	TEVA-VORICONAZOLE (SA)	TEV	<b>FGNQSW</b>
02399253	SANDOZ-VORICONAZOLE (SA)	SDZ	<b>FGNQSW</b>
02525798	JAMP-VORICONAZOLE (SA)	JPC	<b>FGNQSW</b>

**08:16.00 ANTITUBERCULOSIS AGENTS****ETHAMBUTOL**

100MG TABLET

00247960	ETIBI	VAL	<b>AX</b>
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400MG TABLET

00247979	ETIBI	VAL	<b>AX</b>
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**ISONIAZID**

300MG TABLET

00577804	PDP-ISONIAZID	PEN	<b>AX</b>
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**PYRAZINAMIDE**

500MG TABLET

00618810	PDP-PYRAZINAMIDE	PEN	<b>X</b>
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**RIFABUTIN**

150MG CAPSULE

02063786	MYCOBUTIN	PFI	<b>AX</b>
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**RIFAMPIN**

150MG CAPSULE

00393444 ROFACT

VAL **AQX**

300MG CAPSULE

00343617 ROFACT

VAL **AQX****8:16.92 MISCELLANEOUS ANTIMYCOBACTERIALS****DAPSONE**

100MG TABLET

02041510 DAPSONE  
02481227 MAR-DAPSONE  
02489058 RIVA-DAPSONEJAC **AFGNQSW**  
MAR **AFGNQSW**  
RIV **AFGNQSW****8:18.00 ANTIVIRALS****ACYCLOVIR**

200MG TABLET

02207621 APO-ACYCLOVIR  
02242784 MYLAN-ACYCLOVIR  
02285959 TEVA-ACYCLOVIR  
02524708 MINT-ACYCLOVIRAPX **AFGNQSW**  
MYL **AFGNQSW**  
TEV **AFGNQSW**  
MNT **AFGNQSW**

400MG TABLET

02207648 APO-ACYCLOVIR  
02242463 MYLAN-ACYCLOVIR  
02285967 TEVA-ACYCLOVIR  
02524716 MINT-ACYCLOVIRAPX **AFGNQSW**  
MYL **AFGNQSW**  
TEV **AFGNQSW**  
MNT **AFGNQSW**

800MG TABLET

02207656 APO-ACYCLOVIR  
02242464 MYLAN-ACYCLOVIR  
02285975 TEVA-ACYCLOVIR  
02524724 MINT-ACYCLOVIRAPX **AFGNQSW**  
MYL **AFGNQSW**  
TEV **AFGNQSW**  
MNT **AFGNQSW****FAMCICLOVIR**

125MG TABLET

02229110 FAMVIR  
02292025 APO-FAMCICLOVIR  
02305682 ACT-FAMCICLOVIRATN **AFNQSW**  
APX **AFGNQSW**  
TEV **AFGNQSW**

250MG TABLET

02229129	FAMVIR	ATN	<b>AFNQSW</b>
02292041	APO-FAMCICLOVIR	APX	<b>AFGNQSW</b>
02305690	ACT-FAMCICLOVIR	TEV	<b>AFGNQSW</b>

500MG TABLET

02177102	FAMVIR	ATN	<b>AFNQSW</b>
02292068	APO-FAMCICLOVIR	APX	<b>AFGNQSW</b>
02305704	ACT-FAMCICLOVIR	TEV	<b>AFGNQSW</b>

**VALACYCLOVIR**

500MG TABLET

02219492	VALTREX	GSK	<b>AFNQSW</b>
02295822	APO-VALACYCLOVIR	APX	<b>AFGNQSW</b>
02298457	PMS-VALACYCLOVIR	PMS	<b>AFGNQSW</b>
02347091	SANDOZ-VALACYCLOVIR	SDZ	<b>AFGNQSW</b>
02351579	MYLAN-VALACYCLOVIR	MYL	<b>AFGNQSW</b>
02357534	TEVA-VALACYCLOVIR	TEV	<b>AFGNQSW</b>
02405040	AURO-VALACYCLOVIR	ARO	<b>AFGNQSW</b>
02440598	JAMP-VALACYCLOVIR	JPC	<b>AFGNQSW</b>
02441454	JAMP-VALACYCLOVIR	JPC	<b>AFGNQSW</b>
02442000	VALACYCLOVIR	SIV	<b>AFGNQSW</b>
02454645	VALACYCLOVIR	SNS	<b>AFGNQSW</b>

1000MG TABLET

02351560	MYLAN-VALACYCLOVIR	MYL	<b>AFGNQSW</b>
02354705	APO-VALACYCLOVIR	APX	<b>AFGNQSW</b>
02381230	PMS-VALACYCLOVIR	PMS	<b>AFGNQSW</b>
02405059	AURO-VALACYCLOVIR	ARO	<b>AFGNQSW</b>
02519585	VALACYCLOVIR	SNS	<b>AFGNQSW</b>

**VALGANCICLOVIR**

[SEE APPENDIX A](#) FOR SA CRITERIA

450MG TABLET

02245777	VALCYTE (SA)	XPI	<b>AT</b>
02413825	TEVA-VALGANCICLOVIR (SA)	TEV	<b>AT</b>
02435179	AURO-VALGANCICLOVIR (SA)	ARO	<b>AT</b>
02495457	MINT-VALGANCICLOVIR (SA)	MNT	<b>AT</b>

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG/ML ORAL SOLUTION

02306085	VALCYTE (SA)	XPI	<b>AT</b>
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## **8:18.04 ADAMANTANES**

### **AMANTADINE HCL**

10MG/ML SYRUP

02022826	PDP-AMANTADINE	PEN	FGNQSW
02538601	ODAN-AMANTADINE	ODN	FGNQSW

100MG CAPSULE

01990403	PDP-AMANTADINE	PEN	FGNQSW
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## **08:18.08.04 ANTIRETROVIRAL AGENTS (HIV ENTRY AND FUSION INHIBITORS)**

### **ENFUVRTIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

90MG/ML INJECTION KIT

02247725	FUZEON (SA)	HLR	A
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### **MARAVIROX**

150MG TABLET

02299844	CELSENTRI	VII	A
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300MG TABLET

02299852	CELSENTRI	VII	A
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## **08:18.08.08 ANTIRETROVIRAL AGENTS (PROTEASE INHIBITORS)**

### **ATAZANAVIR**

150MG CAPSULE

02248610	REYATAZ	BMS	A
02443791	TEVA-ATAZANAVIR	TEV	A
02456877	MYLAN-ATAZANAVIR	MYL	A
02513102	JAMP-ATAZANAVIR	JPC	A

200MG CAPSULE

02248611	REYATAZ	BMS	A
02443813	TEVA-ATAZANAVIR	TEV	A
02456885	MYLAN-ATAZANAVIR	MYL	A
02513110	JAMP-ATAZANAVIR	JPC	A

300MG CAPSULE

02294176	REYATAZ	BMS	A
02443821	TEVA-ATAZANAVIR	TEV	A
02456893	MYLAN-ATAZANAVIR	MYL	A
02513129	JAMP-ATAZANAVIR	JPC	A

**DARUNAVIR**

75MG TABLET			
02338432	PREZISTA	JAN	A

150MG TABLET			
02369753	PREZISTA	JAN	A

600MG TABLET			
02324024	PREZISTA	JAN	A
02486121	AURO-DARUNAVIR	ARO	A
02487241	APO-DARUNAVIR	APX	A
02521342	DARUNAVIR	JPC	A

800MG TABLET			
02393050	PREZISTA	JAN	A
02486148	AURO-DARUNAVIR	ARO	A
02487268	APO-DARUNAVIR	APX	A
02521350	DARUNAVIR	JPC	A

**DARUNAVIR/COBICISTAT**

800MG/150MG TABLET			
02426501	PREZCOBIX	JAN	A

**ELVITEGRAVIR/COBICISTAT/EMTRICITABINE/TENOFOVIR  
DISOPROXIL FUMARATE**

150MG/150MG/200MG/300MG TABLET			
02397137	STRIBILD	GIL	A

**FOSAMPRENAVIR**

700MG TABLET			
02261545	TELZIR	VII	A

**LOPINAVIR & RITONAVIR**

200MG & 50MG TABLET			
02285533	KALETRA	ABV	A

**NELFINAVIR MESYLATE**

250MG TABLET			
02238617	VIRACEPT	PFI	A

**RITONAVIR**

100MG FILM COATED TABLET  
02357593 NORVIR ABV A

**TIPRANAVIR**

[SEE APPENDIX A](#) FOR SA CRITERIA  
250MG CAPSULE  
02273322 APTIVUS (SA) BOE A

**8:18.08.12 ANTIRETROVIRAL AGENTS (INTEGRASE INHIBITORS)**

**ABACAVIR & DOLUTEGRAVIR & LAMIVUDINE**

600MG & 50MG & 300MG TABLET  
02430932 TRIUMEQ VII A

**BICTEGRAVIR & EMTRICITABINE & TENOFOVIR ALAFENAMIDE**

50MG & 200MG & 25MG  
02478579 BIKTARVY GIL A

**CABOTEGRAVIR**

[SEE APPENDIX A](#) FOR SA CRITERIA  
30MG TABLET  
02497204 VOCABRIA (SA) VII A

**CABOTEGRAVIR & RILPIVIRINE**

[SEE APPENDIX A](#) FOR SA CRITERIA  
400MG/600MG VIAL  
02497220 CABENUVA (SA) VII A

600MG/900MG VIAL

02497247 CABENUVA (SA) VII A

**DOLUTEGRAVIR SODIUM**

50MG TABLET  
02414945 TIVICAY VII A

**DOLUTEGRAVIR SODIUM & LAMIVUDINE**

50MG & 300MG TABLET  
02491753 DOVATO VII A

**DOLUTEGRAVIR/ & RILPIVIRINE**

50MG & 25MG TABLET  
02475774 JULUCA VII A

**ELVITEGRAVIR & COBICISTAT & EMTRICITABINE & TENOFOVIR ALAFENAMIDE**

150MG & 150MG & 200MG & 10MG TABLET  
02449498 GENVOYA GIL A

**RALTEGRAVIR**

400MG TABLET  
02301881 ISENTRESS MSD A

**8:18.08.16 ANTIRETROVIRAL AGENTS (NON-NUCLEOSIDE REVERSE TRANSCRIPTASE INHIBITORS)**

**DORAVIRINE**

100MG TABLET  
02481545 PIFELTRO MER A

**DORAVIRINE & LAMIVUDINE & TENOFIVIR**

100MG & 300MG & 300MG TABLET  
02482592 DELSTRIGO MER A

**EFAVIRENZ**

600MG TABLET  
02381524 MYLAN-EFAVIRENZ MYL A  
02389762 TEVA-EFAVIRENZ TEV A  
02418428 AURO-EFAVIRENZ ARO A  
02458233 JAMP-EFAVIRENZ JPC A

**EMTRICITABINE & RILPIVIRINE & TENOFOVIR**

200MG & 25MG & 300MG TABLET  
02374129 COMPLERA GIL A

**EMTRICITABINE & RILPIVIRINE & TENOFOVIR ALAFENAMIDE**

200MG & 25MG & 25MG TABLET  
02461463 ODEFSEY GIL A

**ETRAVIRINE**

100MG TABLET  
02306778 INTELENCE JAN A

**NEVIRAPINE**

200MG TABLET  
02318601 AURO-NEVIRAPINE ARO A  
02387727 MYLAN-NEVIRAPINE MYL A  
02405776 JAMP-NEVIRAPINE JPC A

**NEVIRAPINE**

400MG EXTENDED RELEASE TABLET

02427931 APO-NEVIRAPINE XR APX A

**RILPIVIRINE**

25MG TABLET

02370603 EDURANT JAN A

**08:18.08.20 ANTIRETROVIRAL AGENTS (NUCLEOSIDE REVERSE TRANSCRIPTASE INHIBITORS)****ABACAVIR SULFATE**

300MG TABLET

02240357 ZIAGEN VII A

02396769 APO-ABACAVIR APX A

02480956 MINT-ABACAVIR MNT A

**ABACAVIR & LAMIVUDINE**

600MG &amp; 300MG TABLET

02269341 KIVEXA VII A

02399539 APO-ABACAVIR/LAMUVIDINE APX A

02416662 TEVA-ABACAVIR/LAMUVIDINE TEV A

02450682 MYLAN-ABACAVIR/LAMUVIDINE MYL A

02454513 AURO-ABACAVIR/LAMUVIDINE ARO A

02458381 PMS-ABACAVIR/LAMIVUDINE PMS A

02497654 JAMP-ABACAVIR/LAMIVUDINE JPC A

**EFAVIRENZ & EMBRICITABINE & TENOFOVIR**

600MG &amp; 200MG &amp; 300MG TABLET

02393549 TEVA-EFAVIRENZ-EMTRICITABINE-TENOFOVIR TEV A

02461412 MYLAN-EFAVIRENZ-EMTRICITABINE-TENOFOVIR MYL A

02468247 APO-EFAVIRENZ-EMTRICITABINE-TENOFOVIR APX A

02478404 AURO-EFAVIRENZ-EMTRICITABINE-TENOFOVIR ARO A

02484676 SANDOZ-EFAVIRENZ-EMTRICITABINE-TENOFOV MYL A

02487284 PMS-EFAVIRENZ-EMTRICITABINE-TENOFOVIR PMS A

02519461 JAMP-EFAVIRENZ/EMTRICITABINE/TENOFOVIR JPC A

**EMTRICITABINE & TENOFOVIR**

200MG &amp; 300MG TABLET

02274906 TRUVADA GIL A

02399059 TEVA-EMTRICITABINE-TENOFOVIR TEV A

02443902 MYLAN-EMTRICITABINE-TENOFOVIR MYL A

02452006	APO-EMTRICITABINE-TENOFOVIR	APX	A
02461110	PMS-EMTRICITABINE-TENOFOVIR	PMS	A
02487012	JAMP-EMTRICITABINE-TENOFOVIR	JPC	A
02490684	AURO-EMTRICITABINE-TENOFOVIR	ARO	A
02496356	AG-EMTRICITABINE-TENOFOVIR	ANG	A
02521547	MINT-EMTRICITABINE-TENOFOVIR	MNT	A

### LAMIVUDINE

#### 100MG TABLET

02239193	HEPTOVIR	GSK	H
02393239	APO-LAMIVUDINE HBV	APX	H
02512467	JAMP-LAMIVUDINE HBV	JPC	H

#### 150MG TABLET

02192683	3TC	VII	AH
02369052	APO-LAMIVUDINE	APX	AH
02507110	JAMP-LAMIVUDINE	JPC	AH

#### 300MG TABLET

02247825	3TC	VII	AH
02369060	APO-LAMIVUDINE	APX	AH
02507129	JAMP-LAMIVUDINE	JPC	AH

### LAMIVUDINE & ZIDOVUDINE

#### 150MG & 300MG TABLET

02239213	COMBIVIR	VII	A
02375540	APO-LAMIVUDINE/ZIDOVUDINE	APX	A
02414414	AURO-LAMIVUDINE/ZIDOVUDINE	ARO	A
02502801	JAMP-LAMIVUDINE/ZIDOVUDINE	JPC	A

### TENOFOVIR

#### 300MG TABLET

02247128	VIREAD	GIL	AH
02403889	TEVA-TENOFOVIR	TEV	AH
02451980	APO-TENOFOVIR	APX	AH
02452634	MYLAN-TENOFOVIR	MYL	AH
02453940	PMS-TENOFOVIR	PMS	AH
02460173	AURO-TENOFOVIR	ARO	AH
02472511	NAT-TENOFOVIR	NAT	AH
02479087	JAMP-TENOFOVIR	JPC	AH
02512327	TENOFOVIR DISOPROXIL FUMARATE	SNS	AH
02512939	MINT-TENOFOVIR	MNT	AH
02523922	TENOFOVIR	SIV	AH

### ZIDOVUDINE (AZT)

#### 100MG CAPSULE

01946323

APO-ZIDOVUDINE

APX A

**08:18.32 NUCLEOSIDES AND NUCLEOTIDES**

**ENTECAVIR**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.5MG TABLET

02282224	BARACLUDE (SA)	BMS	H
02396955	APO-ENTECAVIR (SA)	APX	H
02430576	PMS-ENTECAVIR (SA)	PMS	H
02448777	AURO-ENTECAVIR (SA)	ARO	H
02453797	ENTECAVIR (SA)	STR	H
02467232	JAMP-ENTECAVIR (SA)	JPC	H
02485907	MINT-ENTECAVIR (SA)	MNT	H
02527154	ENTECAVIR (SA)	SNS	H

**08:18.40 HCV PROTEASE INHIBITORS**

**GLECAPREVIR & PIBRENTASVIR**

100MG & 40MG TABLET

02467550	MAVIRET	ABV	H
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**08:18.92 MISCELLANEOUS ANTIVIRALS**

**LETERMOVIR**

[SEE APPENDIX A](#) FOR SA CRITERIA

240MG TABLET

02469375	PREVYMIS (SA)	MER	NMQW
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480MG TABLET

02469383	PREVYMIS (SA)	MER	NMQW
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**NIRMATRELVIR & RITONAVIR**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG (2) & 100MG TABLET

02524031	PAXLOVID (SA)	PFI	FNQSW
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150MG & 100MG TABLET

02527804	PAXLOVID (SA)	PFI	FNQSW
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## **08:30.08 ANTIMALARIALS**

### **HYDROXYCHLOROQUINE SULFATE**

200MG TABLET

02017709	PLAQUENIL	AVN	<b>FNQSW</b>
02246691	APO-HYDROXYQUINE	APX	<b>FGNQSW</b>
02424991	MINT-HYDROXYCHLOROQUINE	MNT	<b>FGNQSW</b>
02491427	JAMP-HYDROXYCHLOROQUINE SULFATE	JPC	<b>FGNQSW</b>
02511886	NRA-HYDROXYCHLOROQUINE	NRA	<b>FGNQSW</b>
02519348	HYDROXYCHLOROQUINE	SNS	<b>FGNQSW</b>

## **08:30.92 MISCELLANEOUS ANTIPROTOZOALS**

### **METRONIDAZOLE**

250MG TABLET

00545066	APO-METRONIDAZOLE	APX	<b>CFGNQSW</b>
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## **08:36.00 URINARY ANTI INFECTIVES**

### **FOSFOMYCIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

3G SACHET

02240335	MONUROL (*)	PAL	<b>FNQSW</b>
02473801	JAMP-FOSFOMYCIN (*)	JPC	<b>FGNQSW</b>

\*quantity limit of 3 doses per 12 month period (one sachet per dispense). The prescriber can submit a request for consideration should beneficiaries require more than 3 doses per 12 month period.

### **NITROFURANTOIN**

50MG CAPSULE (MACROCRYSTALS)

02231015	TEVA-NITROFURANTOIN	TEV	<b>FGNQSW</b>
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100MG CAPSULE (MACROCRYSTALS)

02231016	TEVA-NITROFURANTOIN	TEV	<b>FGNQSW</b>
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50MG TABLET

00319511	NITROFURANTOIN	AAA	<b>FGNQSW</b>
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100MG TABLET			
00312738	NITROFURANTOIN	AAA	<b>FGNQSW</b>

**NITROFURANTOIN MONOHYDRATE/MACROCRYSTALS**

100MG CAPSULE			
02455676	PMS-NITROFURANTOIN	PMS	<b>FGNQSW</b>
02466392	AURO-NITROFURANTOIN	ARO	<b>FGNQSW</b>

**TRIMETHOPRIM**

100MG TABLET			
02243116	TRIMETHOPRIM	AAA	<b>FGNQSW</b>

200MG TABLET			
02243117	TRIMETHOPRIM	AAA	<b>FGNQSW</b>

**10:00.00 ANTINEOPLASTIC AGENTS**

**ABEMACICLIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG TABLET			
02487098	VERZENIO (SA)	LIL	<b>NMQW</b>

100MG TABLET			
02487101	VERZENIO (SA)	LIL	<b>NMQW</b>

150MG TABLET			
02487128	VERZENIO (SA)	LIL	<b>NMQW</b>

**ABIRATERONE ACETATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

250MG TABLET			
02371065	ZYTIGA (SA)	JAN	<b>NMQW</b>
02477114	REDDY-ABIRATERONE (SA)	RCH	<b>NMQW</b>
02486393	SANDOZ-ABIRATERONE (SA)	SDZ	<b>NMQW</b>
02491397	APO-ABIRATERONE (SA)	APX	<b>NMQW</b>
02492601	PMS-ABIRATERONE (SA)	PMS	<b>NMQW</b>
02494132	NAT-ABIRATERONE (SA)	NAT	<b>NMQW</b>
02502305	JAMP-ABIRATERONE (SA)	JPC	<b>NMQW</b>
02503980	MAR-ABIRATERONE (SA)	MAR	<b>NMQW</b>

500MG TABLET			
02457113	ZYTIGA (SA)	JAN	<b>NMQW</b>

02491400	APO-ABIRATERONE (SA)	APX	NMQW
02501503	PMS-ABIRATERONE (SA)	PMS	NMQW
02503999	MAR-ABIRATERONE (SA)	MAR	NMQW
02521644	SANDOZ-ABIRATERONE (SA)	SDZ	NMQW
02525380	ABIRATERONE (SA)	JPC	NMQW
02529629	JAMP ABIRATERONE (SA)	JPC	NMQW
02533251	REDDY-ABIRATERONE (SA)	RCH	NMQW

### ACALABRUTINIB

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE

02491788	CALQUENCE (SA)	AZE	NMQW
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100MG TABLET

02535696	CALQUENCE (SA)	AZE	NMQW
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### AFATINIB

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG TABLET

02415666	GIOTRIF (SA)	BOE	NMQW
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30MG TABLET

02415674	GIOTRIF (SA)	BOE	NMQW
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40MG TABLET

02415682	GIOTRIF (SA)	BOE	NMQW
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### ALECTINIB

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG CAPSULE

02458136	ALECENSARO (SA)	HLR	NMQW
00904400	ALECENSARO (SA)*		NMQW

\*use when drug cost in excess of CPHA maximum

### ANASTROZOLE

1MG TABLET

02224135	ARIMIDEX	AZE	FNQSW
02320738	PMS-ANASTROZOLE	PMS	FGNQSW
02338467	SANDOZ-ANASTROZOLE	SDZ	FGNQSW
02339080	JAMP-ANASTROZOLE	JPC	FGNQSW
02351218	ACH-ANASTROZOLE	ACH	FGNQSW
02365650	TARO-ANASTROZOLE	TAR	FGNQSW
02374420	APO-ANASTROZOLE	APX	FGNQSW
02379562	MAR-ANASTROZOLE	MAR	FGNQSW
02392259	RIVA-ANASTROZOLE	RIV	FGNQSW
02393573	MINT-ANASTROZOLE	MNT	FGNQSW

02394898	TEVA-ANASTROZOLE	TEV	FGNQSW
02417855	NAT-ANASTROZOLE	NAT	FGNQSW
02442736	ANASTROZOLE	SNS	FGNQSW
02529904	ANASTROZOLE	SIV	FGNQSW

**APALUTAMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

60MG TABLET

02478374	ERLEADA (SA)	JAN	NMQW
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240MG TABLET

02540185	ERLEADA (SA)	JAN	NMQW
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**ASCIMINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG TABLET

02528320	SCEMBLIX (SA)	NVR	NMQW
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40MG TABLET

02528339	SCEMBLIX (SA)	NVR	NMQW
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**AXITINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

1MG TABLET

02389630	INLYTA (SA)	PFI	NMQW
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5MG TABLET

02389649	INLYTA (SA)	PFI	NMQW
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**AZACITIDINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

200MG TABLET

02510197	ONUREG (SA)	CEL	NMQW
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300MG TABLET

02510200	ONUREG (SA)	CEL	NMQW
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**BICALUTAMIDE**

50MG TABLET

02184478	CASODEX	AZE	FNQSW
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02270226	TEVA-BICALUTAMIDE	TEV	FGNQSW
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02275589	PMS-BICALUTAMIDE	PMS	FGNQSW
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02296063	APO-BICALUTAMIDE	APX	FGNQSW
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02325985	BICALUTAMIDE	ACH	FGNQSW
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02357216	JAMP-BICALUTAMIDE	JPC	FGNQSW
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02519178	BICALUTAMIDE	SNS	FGNQSW
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**BINIMETINIB**[SEE APPENDIX A](#) FOR SA CRITERIA

15MG TABLET

02513080 MEKTOVI (SA) PFI **NMQW****BOSUTINIB**[SEE APPENDIX A](#) FOR SA CRITERIA

100MG TABLET

02419149 BOSULIF (SA) PFI **NMQW**

500MG TABLET

02419157 BOSULIF (SA) PFI **NMQW****BRIGATINIB**[SEE APPENDIX A](#) FOR SA CRITERIA

30MG TABLET

02479206 ALUNBRIG (SA) TAK **NMQW**00904758 ALUNBRIG (SA)\* **NMQW**

\*use when drug cost in excess of CPHA maximum

90MG TABLET

02479214 ALUNBRIG (SA) TAK **NMQW**00904759 ALUNBRIG (SA)\* **NMQW**

\*use when drug cost in excess of CPHA maximum

180MG TABLET

02479222 ALUNBRIG (SA) TAK **NMQW**00904760 ALUNBRIG (SA)\* **NMQW**

\*use when drug cost in excess of CPHA maximum

90MG (7) &amp; 180MG (21) INITIATION PACK

02479230 ALUNBRIG (SA) TAK **NMQW**00904761 ALUNBRIG (SA)\* **NMQW**

\*use when drug cost in excess of CPHA maximum

**BUSERELIN ACETATE**

6.3MG IMPLANT

02228955 SUPREFACT DEPOT XPI **FNQSW**

9.45MG IMPLANT

02240749 SUPREFACT DEPOT XPI **FNQSW****BUSULFAN**

2MG TABLET

00004618 MYLERAN ASN **FNQSW**

**CABOZANTINIB**[SEE APPENDIX A](#) FOR SA CRITERIA

20MG TABLET

02480824 CABOMETYX (SA) IPS **NMQW**

40MG TABLET

02480832 CABOMETYX (SA) IPS **NMQW**

60MG TABLET

02480840 CABOMETYX (SA) IPX **NMQW****CAPECITABINE**

150MG TABLET

02400022 TEVA-CAPECITABINE TEV **FGNQSW**02421917 SANDOZ-CAPECITABINE SDZ **FGNQSW**02426757 ACH-CAPECITABINE ACH **FGNQSW**02457490 TARO-CAPECITABINE TAR **FGNQSW**02514982 CAPECITABINE SNS **FGNQSW**02519879 CAPECITABINE JPC **FGNQSW**

500MG TABLET

02421925 SANDOZ-CAPECITABINE SDZ **FGNQSW**02426765 ACH-CAPECITABINE ACH **FGNQSW**02457504 TARO-CAPECITABINE TAR **FGNQSW**02508028 MINT-CAPECITABINE MNT **FGNQSW**02514990 CAPECITABINE SNS **FGNQSW**02519887 CAPECITABINE JPC **FGNQSW****CERITINIIB**[SEE APPENDIX A](#) FOR SA CRITERIA

150MG CAPSULE

02436779 ZYKADIA (SA) NVR **NMQW****CHLORAMBUCIL**

2MG TABLET

00004626 LEUKERAN ASN **FNQSW****COBIMETINIB**[SEE APPENDIX A](#) FOR SA CRITERIA

20MG TABLET

02452340 COTELLIC (SA) HLR **NMQW****CRIZOTINIB**[SEE APPENDIX A](#) FOR SA CRITERIA

200MG CAPSULE

02384256 XALKORI (SA) PFI NMQW

250MG CAPSULE

02384264 XALKORI (SA) PFI NMQW

### CYCLOPHOSPHAMIDE

25MG TABLET

02241795 PROCYTOX BAX FNQSW

50MG TABLET

02241796 PROCYTOX BAX FNQSW

### CYPROTERONE ACETATE

50MG TABLET

00704431 ANDROCUR BAY FNQSW  
02245898 APO-CYPROTERONE AAA FGNQSW  
02390760 MED-CYPROTERONE GMP FGNQSW

### DABRAFENIB

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG TABLET

02409607 TAFINLAR (SA) NVR NMQW

75MG TABLET

02409615 TAFINLAR (SA) NVR NMQW

### DAROLUTAMIDE

[SEE APPENDIX A](#) FOR SA CRITERIA

300MG TABLET

02496348 NUBEQA (SA) BAY NMQW

### DASATINIB

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG TABLET

02293129 SPRYCEL (SA) BMS NMQW  
02470705 APO-DASATINIB (SA) APX NMQW  
02478307 TEVA-DASATINIB (SA) TEV NMQW  
02499282 TARO-DASATINIB (SA) TAR NMQW  
02514737 REDDY-DASATINIB (SA) RCH NMQW

50MG TABLET

02293137 SPRYCEL (SA) BMS NMQW  
02470713 APO-DASATINIB (SA) APX NMQW  
02478315 TEVA-DASATINIB (SA) TEV NMQW  
02499304 TARO-DASATINIB (SA) TAR NMQW  
02514745 REDDY-DASATINIB (SA) RCH NMQW

**70MG TABLET**

02293145	SPRYCEL (SA)	BMS	<b>NMQW</b>
02478323	TEVA-DASATINIB (SA)	TEV	<b>NMQW</b>
02481499	APO-DASATINIB (SA)	APX	<b>NMQW</b>
02499312	TARO-DASATINIB (SA)	TAR	<b>NMQW</b>
02514753	REDDY-DASATINIB (SA)	RCH	<b>NMQW</b>

**80MG TABLET**

02360810	SPRYCEL (SA)	BMS	<b>NMQW</b>
02478331	TEVA-DASATINIB (SA)	TEV	<b>NMQW</b>
02481502	APO-DASATINIB (SA)	APX	<b>NMQW</b>
02499320	TARO-DASATINIB (SA)	TAR	<b>NMQW</b>
02514761	REDDY-DASATINIB (SA)	RCH	<b>NMQW</b>

**100MG TABLET**

02320193	SPRYCEL (SA)	BMS	<b>NMQW</b>
02470721	APO-DASATINIB (SA)	APX	<b>NMQW</b>
02478358	TEVA-DASATINIB (SA)	TEV	<b>NMQW</b>
02499339	TARO-DASATINIB (SA)	TAR	<b>NMQW</b>
02514788	REDDY-DASATINIB (SA)	RCH	<b>NMQW</b>

**140MG TABLET**

02360829	SPRYCEL (SA)	BMS	<b>NMQW</b>
02499347	TARO-DASATINIB (SA)	TAR	<b>NMQW</b>
02514796	REDDY-DASATINIB (SA)	RCH	<b>NMQW</b>

**DECITABINE & CEDAZURIDINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**35MG & 100MG TABLET**

02501600	INQOVI (SA)	TAI	<b>NMQW</b>
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**DEGARELIX**

**80MG/VIAL POWDER FOR INJECTION**

02337029	FIRMAGON	FEI	<b>FNQSW</b>
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**120MG/VIAL POWDER FOR INJECTION**

02337037	FIRMAGON	FEI	<b>FNQSW</b>
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**ENCORAFENIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

**75MG CAPSULE**

02513099	BRAFTOVI (SA)	PFI	<b>NMQW</b>
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**ENTRECTINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE  
02495007 ROZLYTREK (SA) HLR NMQW

200MG CAPSULE  
02495015 ROZLYTREK (SA) HLR NMQW

**ENZALUTAMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

40MG CAPSULE  
02407329 XTANDI (SA) AST NMQW

**ERLOTINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

25MG TABLET

02269007 TARCEVA (SA) HLR FMNQSW  
02377691 TEVA-ERLOTINIB (SA) TEV FGMNQSW  
02461862 APO-ERLOTINIB (SA) APX FGMNQSW  
02483912 NAT-ERLOTINIB (SA) NAT FGMNQSW

100MG TABLET

02269015 TARCEVA (SA) HLR FMNQSW  
02377705 TEVA-ERLOTINIB (SA) TEV FGMNQSW  
02454386 PMS-ERLOTINIB (SA) PMS FGMNQSW  
02461870 APO-ERLOTINIB (SA) APX FGMNQSW  
02483920 NAT-ERLOTIBIN (SA) NAT FGMNQSW

150MG TABLET

02269023 TARCEVA (SA) HLR FMNQSW  
02377713 TEVA-ERLOTINIB (SA) TEV FGMNQSW  
02454394 PMS-ERLOTINIB (SA) PMS FGMNQSW  
02461889 APO-ERLOTINIB (SA) APX FGMNQSW  
02483939 NAT-ERLOTINIB (SA) NAT FGMNQSW

**ETOPOSIDE**

50MG CAPSULE

00616192 VEPESID XPI MNQW

**EVEROLIMUS**

[SEE APPENDIX A](#) FOR SA CRITERIA

2.5MG TABLET

02463229 TEVA-EVEROLIMUS (SA) TEV NMQW  
02492911 SANDOZ EVEROLIMUS (SA) SDZ NMQW  
02504677 PMS-EVEROLIMUS (SA) PMS NMQW  
02530090 NAT-EVEROLIMUS (SA) NAT NMQW

5MG TABLET



02463237	TEVA-EVEROLIMUS (SA)	TEV	<b>NMQW</b>
02492938	SANDOZ EVEROLIMUS (SA)	SDZ	<b>NMQW</b>
02504685	PMS-EVEROLIMUS (SA)	PMS	<b>NMQW</b>
02530104	NAT-EVEROLIMUS (SA)	NAT	<b>NMQW</b>

10MG TABLET

02463253	TEVA-EVEROLIMUS (SA)	TEV	<b>NMQW</b>
02492946	SANDOZ EVEROLIMUS (SA)	SDZ	<b>NMQW</b>
02504693	PMS-EVEROLIMUS (SA)	PMS	<b>NMQW</b>
02530120	NAT-EVEROLIMUS (SA)	NAT	<b>NMQW</b>

**EXEMESTANE**

25MG TABLET

02242705	AROMASIN	PFI	<b>FNQSW</b>
02390183	ACT-EXEMESTANE	TEV	<b>FGNQSW</b>
02407841	MED-EXEMESTANE	GMP	<b>FGNQSW</b>
02408473	TEVA-EXEMESTANE	TEV	<b>FGNQSW</b>

**FEDRATINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE

02502445	INREBIC (SA)	CEL	<b>NMQW</b>
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**FLUDARABINE PHOSPHATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

10 MG TABLET

02246226	FLUDARA (SA)	AVN	<b>NMQW</b>
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**FLUOROURACIL /SALICYLIC ACID**

0.5%-10% TOPICAL SOLUTION

02428946	ACTIKERALL	CIP	<b>FNQSW</b>
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**FLUTAMIDE**

250MG TABLET

02238560	FLUTAMIDE	AAA	<b>FGNQSW</b>
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**FULVESTRANT**

[SEE APPENDIX A](#) FOR SA CRITERIA

250MG/5ML SYRINGE

02460130	TEVA-FULVESTRANT (SA)	TEV	<b>FGNQSW</b>
02483610	FULVESTRANT (SA)	SDZ	<b>FGNQSW</b>

**GILTERITINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

40MG TABLET

02495058	XOSPATA (SA)	AST	<b>NMQW</b>
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00904658 XOSPATA (SA)\*  
 00904659 XOSPATA (SA)\*  
 \*use when drug cost in excess of CPHA maximum

**HYDROXYUREA**

500MG CAPSULE

00465283	HYDREA	XPI	<b>FNQSW</b>
02242920	MYLAN-HYDROXYUREA	MYL	<b>FGNQSW</b>
02247937	APO-HYDROXYUREA	APX	<b>FGNQSW</b>

**IBRUTINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

140MG CAPSULE

02434407	IMBRUVICA (SA)	JAN	<b>NMQW</b>
00904337	IMBRUVICA (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**IDELALISIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG TABLET

02438798	ZYDELIG (SA)	GIL	<b>NMQW</b>
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150MG TABLET

02438801	ZYDELIG (SA)	GIL	<b>NMQW</b>
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**IMATINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG TABLET

02253275	GLEEVEC (SA)	NVR	<b>FMNQSW</b>
02355337	APO-IMATINIB (SA)	APX	<b>FGMNQSW</b>
02397285	NAT-IMATINIB (SA)	NAT	<b>FGMNQSW</b>
02399806	TEVA-IMATINIB (SA)	TEV	<b>FGMNQSW</b>
02431114	PMS-IMATINIB (SA)	PMS	<b>FGMNQSW</b>
02490986	ACH-IMATINIB (SA)	ACH	<b>FGMNQSW</b>
02492334	MINT-IMATINIB (SA)	MNT	<b>FGMNQSW</b>
02495066	JAMP-IMATINIB (SA)	JPC	<b>FGMNQSW</b>
02504596	IMATINIB (SA)	SNS	<b>FGMNQSW</b>
02521202	IMATINIB (SA)	SIV	<b>FGMNQSW</b>

400MG TABLET

02253283	GLEEVEC (SA)	NVR	<b>FMNQSW</b>
02355345	APO-IMATINIB (SA)	APX	<b>FGMNQSW</b>
02397293	NAT-IMATINIB (SA)	NAT	<b>FGMNQSW</b>
02399814	TEVA-IMATINIB (SA)	TEV	<b>FGMNQSW</b>
02431122	PMS-IMATINIB (SA)	PMS	<b>FGMNQSW</b>
02490994	ACH-IMATINIB (SA)	ACH	<b>FGMNQSW</b>

02492342	MINT-IMATINIB (SA)	MNT	<b>FGMNQSW</b>
02495074	JAMP-IMATINIB (SA)	JPC	<b>FGMNQSW</b>
02504618	IMATINIB (SA)	SNS	<b>FGMNQSW</b>
02521210	IMATINIB (SA)	SIV	<b>FGMNQSW</b>

**LAROTRECTINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

**25MG CAPSULE**

02490315	VITRAKVI (SA)	BAY	<b>NMQW</b>
00900012	VITRAKVI (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**100MG CAPSULE**

02490323	VITRAKVI (SA)	BAY	<b>NMQW</b>
00900013	VITRAKVI (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**20MG/ML ORAL LIQUID**

02490331	VITRAKVI (SA)	BAY	<b>NMQW</b>
00900014	VITRAKVI (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**LENALIDOMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**2.5MG CAPSULE**

02484714	REDDY-LENALIDOMIDE (SA)	RCH	<b>NMQW</b>
02493837	NAT-LENALIDOMIDE (SA)	NAT	<b>NMQW</b>
02506130	JAMP LENALIDOMIDE (SA)	JPC	<b>NMQW</b>
02507862	TARO-LENALIDOMIDE (SA)	TAR	<b>NMQW</b>
02507927	APO-LENALIDOMIDE (SA)	APZ	<b>NMQW</b>
02518562	SANDOZ-LENALIDOMIDE (SA)	SDZ	<b>NMQW</b>

**5MG CAPSULE**

02304899	REVLIMID (SA)	CEL	<b>NMQW</b>
02483017	REDDY-LENALIDOMIDE (SA)	RCH	<b>NMQW</b>
02493845	NAT-LENALIDOMIDE (SA)	NAT	<b>NMQW</b>
02506149	JAMP-LENALIDOMIDE (SA)	JPC	<b>NMQW</b>
02507870	TARO-LENALIDOMIDE (SA)	TAR	<b>NMQW</b>
02507935	APO-LENALIDOMIDE (SA)	APX	<b>NMQW</b>
02518570	SANDOZ-LENALIDOMIDE (SA)	SDZ	<b>NMQW</b>

**10MG CAPSULE**

02304902	REVLIMID (SA)	CEL	<b>NMQW</b>
02483025	REDDY-LENALIDOMIDE (SA)	RCH	<b>NMQW</b>
02493861	NAT-LENALIDOMIDE (SA)	NAT	<b>NMQW</b>
02506157	JAMP-LENALIDOMIDE (SA)	JPC	<b>NMQW</b>

02507889	TARO-LENALIDOMIDE (SA)	TAR	<b>NMQW</b>
02507943	APO-LENALIDOMIDE (SA)	APX	<b>NMQW</b>
02518589	SANDOZ-LENALIDOMIDE (SA)	SDZ	<b>NMQW</b>

15MG CAPSULE

02317699	REVLIMID (SA)	CEL	<b>NMQW</b>
02483033	REDDY-LENALIDOMIDE (SA)	RCH	<b>NMQW</b>
02493888	NAT-LENALIDOMIDE (SA)	NAT	<b>NMQW</b>
02506165	JAMP-LENALIDOMIDE (SA)	JPC	<b>NMQW</b>
02507897	TARO-LENALIDOMIDE (SA)	TAR	<b>NMQW</b>
02507951	APO-LENALIDOMIDE (SA)	APX	<b>NMQW</b>
02518597	SANDOZ-LENALIDOMIDE (SA)	SDZ	<b>NMQW</b>

20MG CAPSULE

02483041	REDDY-LENALIDOMIDE (SA)	RCH	<b>NMQW</b>
02493896	NAT-LENALIDOMIDE (SA)	NAT	<b>NMQW</b>
02506173	JAMP LLENALIDOMIDE (SA)	JPC	<b>NMQW</b>
02507900	TARO-LENALIDOMIDE (SA)	TAR	<b>NMQW</b>
02507978	APO-LENALIDOMIDE (SA)	APX	<b>NMQW</b>
02518600	SANDOZ-LENALIDOMIDE (SA)	SDZ	<b>NMQW</b>

25MG CAPSULE

02317710	REVLIMID (SA)	CEL	<b>NMQW</b>
02483068	REDDY-LENALIDOMIDE (SA)	RCH	<b>NMQW</b>
02493918	NAT-LENALIDOMIDE (SA)	NAT	<b>NMQW</b>
02506181	JAMP-LENALIDOMIDE (SA)	JPC	<b>NMQW</b>
02507919	TARO-LENALIDOMIDE (SA)	TAR	<b>NMQW</b>
02507986	APO-LENALIDOMIDE (SA)	APX	<b>NMQW</b>
02518619	SANDOZ-LENALIDOMIDE (SA)	SDZ	<b>NMQW</b>

**LENVATINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

4MG CAPSULE

02484056	LENVIMA (SA)	EIS	<b>NMQW</b>
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8MG (2X4MG)

02468220	LENVIMA (SA)	EIS	<b>NMQW</b>
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10MG CAPSULE

02450321	LENVIMA (SA)	EIS	<b>NMQW</b>
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12MG (3X4MG)

02484129	LENVIMA (SA)	EIS	<b>NMQW</b>
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14MG (1X10MG AND 1X4MG)

02450313	LENVIMA (SA)	EIS	<b>NMQW</b>
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22.5MG DEPOT VIAL  
02462699 ZEULIDE DEPOT VER **FNQSW**

30MG DEPOT SYRINGE  
02239833 LUPRON DEPOT ABV **FNQSW**

45MG DEPOT VIAL  
02268892 ELIGARD TOL **FNQSW**

**LORLATINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

25MG TABLET  
02485966 LORBRENA (SA) PFI **NMQW**

100MG TABLET  
02485974 LORBRENA (SA) PFI **NMQW**  
00900025 LORBRENA (SA)\* PFI **NMQW**

\*use when drug cost in excess of CPHA maximum

**MEDROXYPROGESTERONE ACETATE**

100MG TABLET  
02267640 APO-MEDROXY APX **FGNQSW**

**MEGESTROL ACETATE**

40MG TABLET  
02195917 MEGESTROL AAA **AFGNQSW**

160MG TABLET  
02195925 MEGESTROL AAA **AFGNQSW**

**MELPHALAN**

2MG TABLET  
00004715 ALKERAN ASN **FNQSW**

**MERCAPTOPURINE**

50MG TABLET  
00004723 PURINETHOL TEV **FGNQSW**  
02415275 MERCAPTOPURINE STE **FGNQSW**

**METHOTREXATE**

2.5MG TABLET  
02170698 PMS-METHOTREXATE PMS **FGNQSW**  
02182963 APO-METHOTREXATE APX **FGNQSW**  
02509067 ACH-METHOTREXATE ACH **FGNQSW**  
02524023 AURO-METHOTREXATE ARO **FGNQSW**  
02534916 M-METHOTREXATE MRA **FGNQSW**

10MG TABLET			
02182750	METHOTREXATE	PFI	<b>FGNQSW</b>
7.5MG & 0.3ML PREFILLED SYRINGE			
02422166	METHOTREXATE	PMS	<b>FGNQSW</b>
10MG & 0.2ML PREFILLED SYRINGE			
02454831	METOJECT	MED	<b>FNQSW</b>
02539608	PMS-METHOTREXATE	PMS	<b>FGNQSW</b>
10MG & 0.4ML PREFILLED SYRINGE			
02422174	METHOTREXATE	PMS	<b>FGNQSW</b>
12.5MG & 0.25ML PREFILLED SYRINGE			
02454750	METOJECT	MED	<b>FNQSW</b>
02539616	PMS-METHOTREXATE	PMS	<b>FGNQSW</b>
15MG & 0.3ML PREFILLED SYRINGE			
02454858	METOJECT	MED	<b>FNQSW</b>
02491311	METHOTREXATE	ACH	<b>FGNQSW</b>
02539624	PMS-METHOTREXATE	PMS	<b>FGNQSW</b>
15MG & 0.6ML PREFILLED SYRINGE			
02422182	METHOTREXATE	PMS	<b>FGNQSW</b>
17.5MG & 0.35ML PREFILLED SYRINGE			
02454769	METOJECT	MED	<b>FNQSW</b>
02491338	METHOTREXATE	ACH	<b>FGNQSW</b>
02539632	PMS-METHOTREXATE	PMS	<b>FGNQSW</b>
20MG & 0.4ML PREFILLED SYRINGE			
02454866	METOJECT	MED	<b>FNQSW</b>
02491346	METHOTREXATE	ACH	<b>FGNQSW</b>
02539640	PMS-METHOTREXATE	PMS	<b>FGNQSW</b>
20MG & 0.7ML PREFILLED SYRINGE			
02422190	METHOTREXATE	PMS	<b>FGNQSW</b>
22.5MG & 0.45ML PREFILLED SYRINGE			
02454777	METOJECT	MED	<b>FNQSW</b>
02491354	METHOTREXATE	ACH	<b>FGNQSW</b>
02539659	PMS-METHOTREXATE	PMS	<b>FGNQSW</b>
25MG & 0.5ML PREFILLED SYRINGE			
02454874	METOJECT	MED	<b>FNQSW</b>

02491362	METHOTREXATE	ACH	<b>FGNQSW</b>
02539667	PMS-METHOTREXATE	PMS	<b>FGNQSW</b>

25MG/ML PREFILLED SYRINGE			
02422204	METHOTREXATE	PMS	<b>FGNQSW</b>

25MG/ML INJECTION SOLUTION (WITH PRESERVATIVE)			
02182777	METHOTREXATE	PFI	<b>FNQSW</b>
02464365	METHOTREXATE	ACH	<b>FGNQSW</b>

25MG/ML INJECTION SOLUTION			
02099705	METHOTREXATE SODIUM	TEV	<b>FGNQSW</b>
02182955	METHOTREXATE/PF	PFI	<b>FNQSW</b>
02417626	METHOTREXATE	MYL	<b>FGNQSW</b>
02419173	JAMP-METHOTREXATE	JPC	<b>FGNQSW</b>

**MIDOSTAURIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

25MG CAPSULE

02466236	RYDAPT (SA)	NVR	<b>NMQW</b>
00904390	RYDAPT (SA)		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**NILOTINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG CAPSULE

02368250	TASIGNA (SA)	NVR	<b>NMQW</b>
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200MG CAPSULE

02315874	TASIGNA (SA)	NVR	<b>NMQW</b>
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**NIRAPARIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE

02489783	ZEJULA (SA)	GSK	<b>NMQW</b>
00904719	ZEJULA (SA)*		<b>NMQW</b>

100MG TABLET

02530031	ZEJULA (SA)	GSK	<b>NMQW</b>
00904985	ZEJULA (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**OLAPARIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG TABLET



02475200 LYNPARZA (SA) AZE NMQW

150MG TABLET

02475219 LYNPARZA (SA) AZE NMQW

**OSIMERTINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

40MG TABLET

02456214 TAGRISSO (SA) AZE NMQW

80MG TABLET

02456222 TAGRISSO (SA) AZE NMQW

**PALBOCICLIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

75MG CAPSULE

02453150 IBRANCE (SA) PFI NMQW

75MG TABLET

02493535 IBRANCE (SA) PFI NMQW

100MG CAPSULE

02453169 IBRANCE (SA) PFI NMQW

100MG TABLET

02493543 IBRANCE (SA) PFI NMQW

125MG CAPSULE

02453177 IBRANCE (SA) PFI NMQW

125MG TABLET

02493551 IBRANCE (SA) PFI NMQW

**PAZOPANIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

200MG TABLET

02352303 VOTRIENT (SA) NVR NMQW

02525666 PMS-PAZOPANIB (SA) PMS NMQW

**POMALIDOMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

1MG CAPSULE

02419580 POMALYST (SA) CEL NMQW

02504073 REDDY-POMALIDOMIDE (SA) RCH NMQW

02506394 NAT-POMALIDOMIDE (SA) NAT NMQW

02520427 APO-POMALIDOMIDE (SA) APX NMQW

02523973	SANDOZ-POMALIDOMIDE (SA)	SDZ	NMQW
02538059	JAMP-POMALIDOMIDE (SA)	JPC	NMQW

**2MG CAPSULE**

02419599	POMALYST (SA)	CEL	NMQW
02504081	REDDY-POMALIDOMIDE (SA)	RCH	NMQW
02506408	NAT-POMALIDOMIDE (SA)	NAT	NMQW
02520435	APO-POMALIDOMIDE (SA)	APX	NMQW
02523981	SANDOZ-POMALIDOMIDE (SA)	SDZ	NMQW
02538075	JAMP-POMALIDOMIDE (SA)	JPC	NMQW

**3MG CAPSULE**

02419602	POMALYST (SA)	CEL	NMQW
02504103	REDDY-POMALIDOMIDE (SA)	RCH	NMQW
02506416	NAT-POMALIDOMIDE (SA)	NAT	NMQW
02520443	APO-POMALIDOMIDE (SA)	APX	NMQW
02524007	SANDOZ-POMALIDOMIDE (SA)	SDZ	NMQW
02538083	JAMP-POMALIDOMIDE (SA)	JPC	NMQW

**4MG CAPSULE**

02419610	POMALYST (SA)	CEL	NMQW
02504111	REDDY-POMALIDOMIDE (SA)	RCH	NMQW
02506424	NAT-POMALIDOMIDE (SA)	NAT	NMQW
02520451	APO-POMALIDOMIDE (SA)	APX	NMQW
02524015	SANDOZ-POMALIDOMIDE (SA)	SDZ	NMQW
02538091	JAMP-POMALIDOMIDE (SA)	JPC	NMQW

**PONATINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

**15MG TABLET**

02437333	ICLUSIG (SA)	ARI	NMQW
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**REGORAFENIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

**40MG TABLET**

02403390	STIVARGA (SA)	BAY	NMQW
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**RIBOCICLIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

**200MG TABLET**

02473569	KISQALI (SA)	NVR	NMQW
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**RIPRETINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

**50MG TABLET**

02500833	QINLOCK (SA)	MDP	NMQW
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00900026	QINLOCK (SA)*		<b>NMQW</b>
00900027	QINLOCK (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**RITUXIMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG/ML VIAL

02478382	TRUXIMA (SA)	TEV	<b>NMQW</b>
00904561	TRUXIMA (SA)*	TEV	<b>NMQW</b>
02478390	TRUXIMA (SA)	TEV	<b>NMQW</b>
00904560	TRUXIMA (SA)*	TEV	<b>NMQW</b>
02495724	RUXIENCE (SA)	PFI	<b>NMQW</b>
00904559	RUXIENCE (SA)*	PFI	<b>NMQW</b>
02498316	RIXIMYO (SA)	SDZ	<b>NMQW</b>
00904590	RIXIMYO (SA)*	SDZ	<b>NMQW</b>
02513447	RIABNI (SA)	AMG	<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**RUXOLITINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG TABLET

02388006	JAKAVI (SA)	NVR	<b>NMQW</b>
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10MG TABLET

02434814	JAKAVI (SA)	NVR	<b>NMQW</b>
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15MG TABLET

02388014	JAKAVI (SA)	NVR	<b>NMQW</b>
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20MG TABLET

02388022	JAKAVI (SA)	NVR	<b>NMQW</b>
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**SELINEXOR**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG TABLET

02527677	XPOVIO (SA)	FTI	<b>NMQW</b>
00900031	XPOVIO (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**SELPERCATINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

40MG CAPSULE

02516918	RETEVMO (SA)	LIL	<b>NMQW</b>
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80MG CAPSULE

02516926	RETEVMO (SA)	LIL	<b>NMQW</b>
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**SUNITINIB MALATE**[SEE APPENDIX A](#) FOR SA CRITERIA**12.5MG CAPSULE**

02280795	SUTENT (SA)	PFI	<b>NMQW</b>
02524058	TARO-SUNITINIB (SA)	TAR	<b>NMQW</b>
02526204	TEVA-SUNITINIB (SA)	TEV	<b>NMQW</b>
02532840	SANDOZ-SUNITINIB (SA)	SDZ	<b>NMQW</b>

**25MG CAPSULE**

02280809	SUTENT (SA)	PFI	<b>NMQW</b>
02524066	TARO-SUNITINIB (SA)	TAR	<b>NMQW</b>
02526212	TEVA-SUNITINIB (SA)	TEV	<b>NMQW</b>
02532867	SANDOZ-SUNITINIB (SA)	SDZ	<b>NMQW</b>

**50MG CAPSULE**

02280817	SUTENT (SA)	PFI	<b>NMQW</b>
02524082	TARO-SUNITINIB (SA)	TAR	<b>NMQW</b>
02526220	TEVA-SUNITINIB (SA)	TEV	<b>NMQW</b>
02532883	SANDOZ-SUNITINIB (SA)	SDZ	<b>NMQW</b>

**TAMOXIFEN CITRATE****10MG TABLET**

00812404	APO-TAMOX	APX	<b>FGNQSW</b>
00851965	TEVA-TAMOXIFEN	TEV	<b>FGNQSW</b>

**20MG TABLET**

00812390	APO-TAMOX	APX	<b>FGNQSW</b>
00851973	TEVA-TAMOXIFEN	TEV	<b>FGNQSW</b>
02048485	NOLVADEX D	AZE	<b>FGNQSW</b>

**TEMOZOLOMIDE**[SEE APPENDIX A](#) FOR HIGH COST DRUG PROGRAM CRITERIA**5MG CAPSULE**

02241093	TEMODAL	MSD	<b>FMNQSW</b>
02441160	TEVA-TEMOZOLOMIDE	TEV	<b>FGMNQSW</b>
02443473	TARO-TEMOZOLOMIDE	TAR	<b>FGMNQSW</b>
02516799	JAMP-TEMOZOLOMIDE	JPC	<b>FGMNQSW</b>

**20MG CAPSULE**

02241094	TEMODAL	MSD	<b>FMNQSW</b>
02395274	TEVA-TEMOZOLOMIDE	TEV	<b>FGMNQSW</b>
02443481	TARO-TEMOZOLOMIDE	TAR	<b>FGMNQSW</b>
02516802	JAMP-TEMOZOLOMIDE	JPC	<b>FGMNQSW</b>

**100MG CAPSULE**

02241095	TEMODAL	MSD	<b>FMNQSW</b>
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02395282	TEVA-TEMOZOLOMIDE	TEV	<b>FGMNQSW</b>
02443511	TARO-TEMOZOLOMIDE	TAR	<b>FGMNQSW</b>
02516810	JAMP-TEMOZOLOMIDE	JPC	<b>FGMNQSW</b>

**140MG CAPSULE**

02312794	TEMODAL	MSD	<b>FMNQSW</b>
02395290	TEVA-TEMOZOLOMIDE	TEV	<b>FGMNQSW</b>
02443538	TARO-TEMOZOLOMIDE	TAR	<b>FGMNQSW</b>
02516829	JAMP-TEMOZOLOMIDE	JPC	<b>FGMNQSW</b>

**250MG CAPSULE**

02241096	TEMODAL	MSD	<b>FMNQSW</b>
02395312	TEVA-TEMOZOLOMIDE	TEV	<b>FGMNQSW</b>
02443554	TARO-TEMOZOLOMIDE	TAR	<b>FGMNQSW</b>
02516845	JAMP-TEMOZOLOMIDE	JPC	<b>FGMNQSW</b>

**THIOGUANINE**

40MG TABLET			
00282081	LANVIS	ASN	<b>FNQSW</b>

**TRAMETINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

**0.5MG TABLET**

02409623	MEKINIST (SA)	NVR	<b>NMQW</b>
00904170	MEKINIST (SA)*		<b>NMQW</b>

**2MG TABLET**

02409658	MEKINIST (SA)	NVR	<b>NMQW</b>
00904171	MEKINIST (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**TRETINOIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

**10MG CAPSULE**

02145839	VESANOID (SA)	XPI	<b>NMQW</b>
02520036	JAMP-TRETINOIN (SA)	JPC	<b>NMQW</b>

**TRIFLURIDINE & TIPIRACIL HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

**15MG & 6.14MG TABLET**

02472104	LONSURF (SA)	TAI	<b>NMQW</b>
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**20MG & 8.19MG TABLET**

02472112	LONSURF (SA)	TAI	<b>NMQW</b>
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**TRIPTORELIN**

3.75MG INTRAMUSCULAR INJECTION  
02240000 TRELSTAR

KNI **FNQSW**

11.25MG INTRAMUSCULAR INJECTION  
02243856 TRELSTAR LA

KNI **FNQSW**

**TUCATINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG TABLET  
02499827 TUKYSA (SA)

SGC **NMQW**

150MG TABLET  
02499835 TUKYSA (SA)  
00904820 TUKYSA (SA)\*

SGC **NMQW**

SGC **NMQW**

\*use when drug cost in excess of CPHA maximum

**VANDETANIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG TABLET  
02378582 CAPRELSA (SA)

GZY **NMQW**

300MG TABLET  
02378590 CAPRELSA (SA)

GZY **NMQW**

**VEMURAFENIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

240MG TABLET  
02380242 ZELBORAF (SA)

HLR **NMQW**

**VENETOCLAX**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG (14), 50MG (7), 100MG (7), 100MG (14) STARTER PACK  
02458063 VENCLEXTA (SA)

ABV **NMQW**

10MG TABLET  
02458039 VENCLEXTA (SA)

ABV **NMQW**

50MG TABLET  
02458047 VENCLEXTA (SA)

ABV **NMQW**

100MG TABLET  
02458055 VENCLEXTA (SA)

ABV **NMQW**

**VISMODEGIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG CAPSULE			
02409267	ERIVEDGE (SA)	HLR	NMQW

**ZANUBRUTINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

80MG CAPSULE			
02512963	BRUKINSA (SA)	BGN	NMQW

**12:04.00 PARASYMPATHOMIMETIC (CHOLINERGIC) AGENTS**

**BETHANECHOL CHLORIDE**

10MG TABLET			
01947958	DUVOID	PAL	FNQSW
25MG TABLET			
01947931	DUVOID	PAL	FNQSW
50MG TABLET			
01947923	DUVOID	PAL	FNQSW

**DONEPEZIL**

5MG TABLET			
02232043	ARICEPT	PFI	FNQSW
02322331	PMS-DONEPEZIL	PMS	FGNQSW
02328666	SANDOZ-DONEPEZIL	SDZ	FGNQSW
02340607	TEVA-DONEPEZIL	TEV	FGNQSW
02362260	APO-DONEPEZIL	APX	FGNQSW
02381508	RAN-DONEPEZIL	RAN	FGNQSW
02400561	AURO-DONEPEZIL	ARO	FGNQSW
02402092	MAR-DONEPEZIL	MAR	FGNQSW
02402645	DONEPEZIL	SIV	FGNQSW
02408600	MINT-DONEPEZIL	MNT	FGNQSW
02416948	JAMP-DONEPEZIL	JPC	FGNQSW
02420597	DONEPEZIL	SIV	FGNQSW
02426846	DONEPEZIL	SNS	FGNQSW
02432684	AG-DONEPEZIL	ANG	FGNQSW
02439557	NAT-DONEPEZIL	NAT	FGNQSW
02467453	M-DONEPEZIL	MRA	FGNQSW
02475278	DONEPEZIL	RIV	FGNQSW

10MG TABLET			
02232044	ARICEPT	PFI	FNQSW
02322358	PMS-DONEPEZIL	PMS	FGNQSW
02328682	SANDOZ-DONEPEZIL	SDZ	FGNQSW

02340615	TEVA-DONEPEZIL	TEV	FGNQSW
02362279	APO-DONEPEZIL	APX	FGNQSW
02381516	RAN-DONEPEZIL	RAN	FGNQSW
02400588	AURO-DONEPEZIL	ARO	FGNQSW
02402106	MAR-DONEPEZIL	MAR	FGNQSW
02402653	DONEPEZIL	SIV	FGNQSW
02408619	MINT-DONEPEZIL	MNT	FGNQSW
02416956	JAMP-DONEPEZIL	JPC	FGNQSW
02420600	DONEPEZIL	SIV	FGNQSW
02426854	DONEPEZIL	SNS	FGNQSW
02432692	AG-DONEPEZIL	ANG	FGNQSW
02439565	NAT-DONEPEZIL	NAT	FGNQSW
02467461	M-DONEPEZIL	MRA	FGNQSW
02475286	DONEPEZIL	RIV	FGNQSW

### **GALANTAMINE**

SEE CHOLINESTERASE INHIBITORS IN APPENDIX A FOR SA CRITERIA

[SEE APPENDIX A](#) FOR SA CRITERIA

8MG EXTENDED RELEASE CAPSULE

02316943	PAT-GALANTAMINE ER (SA)	PAT	FGNQSW
02339439	MYLAN-GALANTAMINE (SA)	MYL	FGNQSW
02425157	AURO-GALANTAMINE ER (SA)	ARO	FGNQSW
02443015	GALANTAMINE ER (SA)	SNS	FGNQSW

16MG EXTENDED RELEASE CAPSULE

02316951	PAT-GALANTAMINE ER (SA)	PAT	FGNQSW
02339447	MYLAN-GALANTAMINE (SA)	MYL	FGNQSW
02425165	AURO-GALANTAMINE ER (SA)	ARO	FGNQSW
02443023	GALANTAMINE ER (SA)	SNS	FGNQSW

24MG EXTENDED RELEASE CAPSULE

02316978	PAT-GALANTAMINE ER (SA)	PAT	FGNQSW
02339455	MYLAN-GALANTAMINE (SA)	MYL	FGNQSW
02425173	AURO-GALANTAMINE ER (SA)	ARO	FGNQSW
02443031	GALANTAMINE ER (SA)	SNS	FGNQSW

### **PILOCARPINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG TABLET

02216345	SALAGEN (SA)	AMD	FNQSW
02496119	M-PILOCARPINE (SA)	MRA	FGNQSW
02509571	JAMP-PILOCARPINE (SA)	JPC	FGNQSW

### **PYRIDOSTIGMINE BROMIDE**

60MG TABLET

00869961	MESTINON	VAL	FNQSW
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02495643	RIVA-PYRIDOSTIGMINE	RIV	<b>FGNQSW</b>
02508362	JAMP PYRIDOSTIGMINE BROMIDE	JPC	<b>FGNQSW</b>

180MG LONG ACTING TABLET			
00869953	MESTINON	VAL	<b>FNQSW</b>

**RIVASTIGMINE**

SEE CHOLINESTERASE INHIBITORS FOR SA CRITERIA

[SEE APPENDIX A](#) FOR SA CRITERIA

**1.5MG CAPSULE**

02242115	EXELON (SA)	KNI	<b>FNQSW</b>
02324563	SANDOZ-RIVASTIGMINE (SA)	SDZ	<b>FGNQSW</b>
02336715	APO-RIVASTIGMINE (SA)	APX	<b>FGNQSW</b>
02401614	MED-RIVASTIGMINE (SA)	GMP	<b>FGNQSW</b>
02485362	JAMP-RIVASTIGMINE (SA)	JPC	<b>FGNQSW</b>

**3MG CAPSULE**

02242116	EXELON (SA)	KNI	<b>FNQSW</b>
02324571	SANDOZ-RIVASTIGMINE (SA)	SDZ	<b>FGNQSW</b>
02336723	APO-RIVASTIGMINE (SA)	APX	<b>FGNQSW</b>
02401622	MED-RIVASTIGMINE (SA)	GMP	<b>FGNQSW</b>
02485370	JAMP-RIVASTIGMINE (SA)	JPC	<b>FGNQSW</b>

**4.5MG CAPSULE**

02242117	EXELON (SA)	KNI	<b>FNQSW</b>
02324598	SANDOZ-RIVASTIGMINE (SA)	SDZ	<b>FGNQSW</b>
02336731	APO-RIVASTIGMINE (SA)	APX	<b>FGNQSW</b>
02401630	MED-RIVASTIGMINE (SA)	GMP	<b>FGNQSW</b>
02485389	JAMP-RIVASTIGMINE (SA)	JPC	<b>FGNQSW</b>

**6MG CAPSULE**

02242118	EXELON (SA)	KNI	<b>FNQSW</b>
02324601	SANDOZ-RIVASTIGMINE (SA)	SDZ	<b>FGNQSW</b>
02336758	APO-RIVASTIGMINE (SA)	APX	<b>FGNQSW</b>
02401649	MED-RIVASTIGMINE (SA)	GMP	<b>FGNQSW</b>
02485397	JAMP-RIVASTIGMINE (SA)	JPC	<b>FGNQSW</b>

**12:08.08 ANTIMUSCARINICS/ANTISPASMODICS**

**ACLIDINIUM BROMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

400MCG/ACTUATION AEROSOL POWDER			
02409720	TUDORZA GENUAIR (SA)	AZE	<b>FNQSW</b>

**ACLIDINIUM BROMIDE & FORMOTEROL FUMARATE DIHYDRATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

400MCG & 12MCG/ACTUATION AEROSOL POWDER  
02439530 DUAKLIR GENUAIR (SA)

AZE **FNQSW**

**ATROPINE SULFATE**

0.6MG/ML INJECTION SOLUTION (1ML)  
00392693 ATROPINE SULFATE

SDZ **N**

**BUDESONIDE & GLYCOPYRRONIUM & FORMOTEROL**

[SEE APPENDIX A](#) FOR SA CRITERIA

160MCG & 7.2MCG & 5MCG METERED DOSE INHALER  
02518058 BREZTRI AEROSPHERE (SA)

AZE **FNQSW**

**FLUTICASONE & UMECLIDIINIUM & VILANTEROL**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MCG & 62.5MCG & 25MCG DRY POWDER FOR INHALATION  
02474522 TRELEGY ELLIPTA (SA)

GSK **FNQSW**

**GLYCOPYRRONIUM BROMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MCG INHALATION CAPSULE  
02394936 SEEBRI BREEZHALER (SA)

NVR **FNQSW**

**HYOSCINE BUTYLBROMIDE**

10MG TABLET  
00363812 BUSCOPAN  
02512335 ACCEL-HYOSCINE

SNC **FNQSW**  
ACC **FGNQSW**

20MG/ML VIAL  
02229868 HYOSCINE BUTYLBROMIDE

SDZ **N**

**INDACATEROL & GLYCOPYRRONIUM**

[SEE APPENDIX A](#) FOR SA CRITERIA

110MCG & 50MCG INHALATION CAPSULE  
02418282 ULTIBRO BREEZHALER (SA)

NVR **FNQSW**

**INDACATEROL & GLYCOPYRRONIUM & MOMETASONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MCG & 50MCG & 160MCG INHALATION CAPSULE  
02501244 ENERZAIR BREEZHALER (SA)

NVR **FNQSW**

**INDACATEROL & MOMETASONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MCG & 80MCG INHALATION CAPSULE

02498685	ATECTURA BREEZHALER (SA)	NVR	<b>FNQSW</b>
150MCG & 160MCG INHALATION CAPSULE			
02498707	ATECTURA BREEZHALER (SA)	NVR	<b>FNQSW</b>
150MCG & 320MCG INHALATION CAPSULE			
02498693	ATECTURA BREEZHALER (SA)	NVR	<b>FNQSW</b>
<b>IPRATROPIUM BROMIDE</b>			
200UG/DOSE INHALER AEROSOL (200 DOSE)			
02247686	ATROVENT HFA	BOE	<b>CFNQSW</b>
02542587	JAMP-IPRATROPIUM HFA	JPC	<b>CFGNQSW</b>
0.25MG/ML INHALATION SOLUTION (20ML)			
02126222	AA-IPRAVENT	AAA	<b>CFGNQSW</b>
02231136	PMS-IPRATROPIUM	PMS	<b>CFGNQSW</b>
0.0125% INHALATION SOLUTION NEBULE (2ML)			
02231135	PMS-IPRATROPIUM	PMS	<b>FGNQSW</b>
0.025% INHALATION SOLUTION NEBULE (2ML)			
02216221	TEVA-IPRATROPIUM	TEV	<b>FGNQSW</b>
02231245	PMS-IPRATROPIUM	PMS	<b>FGNQSW</b>
0.03% NASAL SPRAY - 345 DOSES			
02239627	PMS-IPRATROPIUM	PMS	<b>CFGNQSW</b>
<b>IPRATROPIUM &amp; SALBUTAMOL</b>			
1.0MG & 0.2MG PER ML INHALATION SOLUTION NEBULE (2.5ML)			
02272695	TEVA-COMBO STERINEBS	TEV	<b>FGNQSW</b>
02483394	IPRATROPIUM-SALBUTAMOL	MDN	<b>FGNQSW</b>
<b>IPRATROPIUM BROMIDE &amp; SALBUTAMOL SULPHATE</b>			
20MCG-100MCG/ACTUATION MIST INHALER			
02419106	COMBIVENT RESPIMAT	BOE	<b>FNQSW</b>
<b>PINAVERIUM BROMIDE</b>			
50MG TABLET			
01950592	DICETEL	BGP	<b>FNQSW</b>
02469677	PINAVERIUM	AAA	<b>FGNQSW</b>
100MG TABLET			
02230684	DICETEL	BGP	<b>FNQSW</b>
02469685	PINAVERIUM	AAA	<b>FGNQSW</b>

**SCOPOLAMINE HYDROBROMIDE**

0.4MG/ML VIAL INJECTION

02242810 SCOPOLAMINE HYDROBROMIDE

OMG NQ

**TIOTROPIUM BROMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

18UG CAPSULE WITH INHALATION DEVICE

02246793 SPIRIVA (SA)

BOE FNQSW

02537850 LUPIN-TIOTROPIUM (SA)

LUP FGNQSW

2.5UG/ACTUATION MIST INHALER

02435381 SPIRIVA RESPIMAT (SA)

BOE FNQSW

**TIOTROPIUM & OLODATEROL**

[SEE APPENDIX A](#) FOR SA CRITERIA

2.5MCG & 2.5MCG/ACTUATION MIST INHALER

02441888 INSPIOLTO RESPIMAT (SA)

BOE FNQSW

**UNMECLIDINIUM BROMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

62.5MCG/ACTUATION BLISTER WITH INHALATION DEVICE

02423596 INCRUSE ELLIPTA (SA)

GKS FNQSW

**UMECLIDINIUM BROMIDE & VILANTEROL TRIFENATATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

62.5MCG & 25MCG/ACTUATION BLISTER WITH INHALATION DEVICE

02418401 ANORO ELLIPTA (SA)

GSK FNQSW

**12:12.00 SYMPATHOMIMETIC (ADRENERGIC) AGENTS**

**EPINEPHRINE HCL**

1MG/ML INJECTION SOLUTION (1ML)

00721891 EPINEPHRINE INJECTION USP

HOS NQ

00155357 ADRENALINE CHLORIDE

ERF NQ

02435810 EPINEPHRINE

TLG NQ

[SEE APPENDIX A](#) FOR SA CRITERIA

0.15MG PER DOSE AUTO-INJECTOR

00578657 EPIPEN JR. (\*)

PFI FQW

0.3MG PER DOSE AUTO-INJECTOR

00509558 EPIPEN (\*)

PFI FQW

\*quantity limit of two (2) injections per 12 month period (one unit per dispense). The prescriber can submit a request for consideration should beneficiaries require more than two (2) injections per 12 month period.

**EPINEPHRINE BITARTRATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.15MG PER DOSE PRE-FILLED PEN

02458438 EMERADE (\*) BAU FQW

0.3MG PER DOSE PRE-FILLED PEN

02458446 EMERADE (\*) BAU FQW

0.5MG PER DOSE PRE-FILLED PEN

02458454 EMERADE (\*) BAU FQW

\*quantity limit of two (2) injections per 12 month period (one unit per dispense). The prescriber can submit a request for consideration should beneficiaries require more than two (2) injections per 12 month period.

**FLUTICASONE FUROATE/VILANTEROL**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MCG-25MCG/DOSE

02408872 BREO ELLIPTA (SA) GSK FNQSW

200MCG-25MCG/DOSE

02444186 BREO ELLIPTA (SA) GSK FNQSW

**FORMOTEROL FUMARATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

6UG/DOSE INHALER POWDER

02237225 OXEZE TURBUHALER (SA) AZE FNQSW

12UG/DOSE INHALER POWDER

02237224 OXEZE TURBUHALER (SA) AZE FNQSW

**FORMOTEROL & BUDESONIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

6UG & 100UG PER DOSE INHALER POWDER

02245385 SYMBICORT TURBUHALER (SA) AZE FNQSW

6UG & 200UG PER DOSE INHALER POWDER

02245386 SYMBICORT TURBUHALER (SA) AZE FNQSW

**INDACATEROL**

[SEE APPENDIX A](#) FOR SA CRITERIA

75MCG INHALATION POWDER CAPSULE

02376938 ONBREZ (SA) NVR **FNQSW**

**⑤MIDODRINE HCL**

**2.5MG TABLET**

02278677	APO-MIDODRINE	APX	<b>FGNQSW</b>
02473984	MAR-MIDODRINE	MAR	<b>FGNQSW</b>
02517701	JAMP-MIDODRINE	JPC	<b>FGNQSW</b>
02533200	MIDODRINE	SNS	<b>FGNQSW</b>

**5MG TABLET**

02278685	APO-MIDODRINE	APX	<b>FGNQSW</b>
02473992	MAR-MIDODRINE	MAR	<b>FGNQSW</b>
02517728	JAMP-MIDODRINE	JPC	<b>FGNQSW</b>
02533219	MIDODRINE	SNS	<b>FGNQSW</b>

**MOMETASONE FUROATE/FORMOTEROL FUMARATE DIHYDRATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**100MCG/5MCG INHALER**

02361752 ZENHALE (SA) MSD **FNQSW**

**200MCG/5MCG INHALER**

02361760 ZENHALE (SA) MSD **FNQSW**

**SALBUTAMOL**

**100UG/DOSE INHALER AEROSOL HYDROFLUOROALKANE (HFA) (200 DOSE)**

02232570	AIROMIR HFA	VAL	<b>CFNQSW</b>
02241497	VENTOLIN HFA	GSK	<b>CFNQSW</b>
02245669	APO-SALVENT CFC FREE	APX	<b>CFGNQSW</b>
02326450	NOVO-SALBUTAMOL HFA	TEV	<b>CFGNQSW</b>
02419858	SALBUTAMOL HFA	SNS	<b>CFGNQSW</b>

**200UG/DOSE INHALER POWDER**

02243115 VENTOLIN DISKUS GSK **CFNQSW**

**5MG/ML INHALATION SOLUTION (10ML)**

02213486 VENTOLIN GSK **CFNQSW**

**0.5MG/ML INHALATION SOLUTION PRESERVATIVE FREE NEBULE (2.5ML)**

02208245 PMS-SALBUTAMOL PMS **CFGNQSW**

**1MG/ML INHALATION SOLUTION PRESERVATIVE FREE NEBULE (2.5ML)**

01926934	TEVA-SALBUTAMOL STERINEB	TEV	<b>CFGNQSW</b>
02208229	PMS-SALBUTAMOL	PMS	<b>CFGNQSW</b>
02213419	VENTOLIN NEBULES P.F.	GSK	<b>CFNQSW</b>

**2MG/ML INHALATION SOLUTION PRESERVATIVE FREE NEBULE (2.5ML)**

02213427	VENTOLIN NEBULES P.F.	GSK	CFNQSW
02173360	TEVA-SALBUTAMOL STERINEB	TEV	CFGNQSW
02208237	PMS-SALBUTAMOL	PMS	CFGNQSW

**SALMETEROL XINAFOATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MCG/DOSE INHALED POWDER DISK (60)

02231129	SEREVENT DISKUS (SA)	GSK	FNQSW
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**SALMETEROL & FLUTICASONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

25MCG & 125MCG/DOSE INHALER AEROSOL

02245126	ADVAIR (SA)	GSK	FNQSW
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25MCG & 250MCG/DOSE INHALER AEROSOL

02245127	ADVAIR (SA)	GSK	FNQSW
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50MCG & 100MCG/DOSE INHALER POWDER DISK

02240835	ADVAIR DISKUS (SA)	GSK	FNQSW
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02494507	PMS-FLUTICASONE/SALMETEROL (SA)	PMS	FGNQSW
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02495597	WIXELA INHUB (SA)	MYL	FGNQSW
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50MCG & 250MCG/DOSE INHALER POWDER DISK

02240836	ADVAIR DISKUS (SA)	GSK	FNQSW
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02494515	PMS-FLUTICASONE/SALMETEROL (SA)	PMS	FGNQSW
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02495600	WIXELA INHUB (SA)	MYL	FGNQSW
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50MCG & 500MCG/DOSE INHALER POWDER DISK

02240837	ADVAIR DISKUS (SA)	GSK	FNQSW
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02494523	PMS-FLUTICASONE/SALMETEROL (SA)	PMS	FGNQSW
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02495619	WIXELA INHUB (SA)	MYL	FGNQSW
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**TERBUTALINE SULFATE**

0.5MG/DOSE INHALER POWDER

00786616	BRICANYL TURBUHALER	AZE	CFNQSW
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**12:16.00 SYMPATHOLYTIC AGENTS (ANTIMIGRAINE DRUGS)**

**DIHYDROERGOTAMINE MESYLATE**

4MG/ML NASAL SPRAY

02228947	MIGRANAL	STE	FNQSW
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Note: Coverage is limited to 6 bottles per 30 day period.

## **12:16.04 SELECTIVE ALPHA-1-ADRENERGIC BLOCKING AGENTS**

### **ALFUZOSIN**

10MG EXTENDED RELEASE TABLET

02304678	SANDOZ-ALFUZOSIN	SDZ	FGNQSW
02315866	APO-ALFUZOSIN	APX	FGNQSW
02443201	AURO-ALFUZOSIN	ARO	FGNQSW
02447576	ALFUZOSIN	SIV	FGNQSW
02519844	ALFUZOSIN	SNS	FGNQSW

### **TAMSULOSIN**

0.4MG CONTROL RELEASE TABLET

02270102	FLOMAX CR	BOE	FNQSW
02340208	SANDOZ-TAMSULOSIN	SDZ	FGNQSW
02362406	APO-TAMULOSIN	APX	FGNQSW
02368242	TEVA-TAMSULOSIN CR	TEV	FGNQSW
02427117	TAMSULOSIN CR	SNS	FGNQSW
02429667	TAMSULOSIN CR	SIV	FGNQSW
02545179	AURO-TAMSULOSIN CR	ARO	FGNQSW

## **12:20.00 SKELETAL MUSCLE RELAXANTS**

### **BACLOFEN**

10MG TABLET

02063735	PMS-BACLOFEN	PMS	FGNQSW
02088398	MYLAN BACLOFEN	MYL	FGNQSW
02139332	APO-BACLOFEN	APX	FGNQSW
02287021	BACLOFEN	SNS	FGNQSW
02544660	BACLOFEN	SIV	FGNQSW

20MG TABLET

02063743	PMS-BACLOFEN	PMS	FGNQSW
02088401	MYLAN BACLOFEN	MYL	FGNQSW
02139391	APO-BACLOFEN	APX	FGNQSW
02287048	BACLOFEN	SNS	FGNQSW
02544679	BACLOFEN	SIV	FGNQSW

### **CYCLOBENZAPRINE HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET

02080052	TEVA- CYCLOBENZAPRINE (SA)	TEV	FGNQSW
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02177145	APO-CYCLOBENZAPRINE (SA)	APX	FGNQSW
02212048	PMS-CYCLOBENZAPRINE (SA)	PMS	FGNQSW
02287064	CYCLOBENZAPRINE (SA)	SNS	FGNQSW
02348853	AURO-CYCLOBENZAPRINE (SA)	ARO	FGNQSW
02357127	JAMP-CYCLOBENZAPRINE (SA)	JPC	FGNQSW
02424584	CYCLOBENZAPRINE (SA)	SIV	FGNQSW
02485419	AG-CYCLOBENZAPRINE (SA)	ANG	FGNQSW
02495422	FLEXERIL (SA)	ORI	FGNQSW

**DANTROLENE SODIUM**

25MG CAPSULE

01997602	DANTRIUM	PAL	FNQSW
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**METHOCARBAMOL & ACETAMINOPHEN**

400MG & 325MG CAPLET

02026805	ROBAXACET	PFI	W
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**METHOCARBAMOL & ACETYLSALICYLIC ACID**

400MG & 325MG CAPLET

00868868	METHOXISAL	ROG	W
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**METHOCARBAMOL & ACETYLSALICYLIC ACID & CODEINE**

400MG & 325MG & 16.2MG CAPLET

01934783	ROBAXISAL C-1/4	PFI	FQW
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400MG & 325MG & 32.4MG CAPLET

01934791	ROBAXISAL C-1/2	PFI	FQW
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**TIZANIDINE HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

4MG TABLET

02259893	APO-TIZANIDINE (SA)	APX	FGNQSW
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02536765	MINT-TIZANIDINE (SA)	MNT	FGNQSW
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**12:92.00 MISCELLANEOUS AUTONOMIC DRUGS**

**BUPROPION**

150 MG SUSTAINED RELEASE TABLET

02238441	ZYBAN	VAL	Z
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**NICOTINE**

7MG/24HOUR TRANSDERMAL PATCH

00999973	NICOTINE PATCH (DIN for billing purposes only)		Z
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14MG/24HOUR TRANSDERMAL PATCH  
00999974 NICOTINE PATCH (DIN for billing purposes only) Z

21MG/24HOUR TRANSDERMAL PATCH  
00999975 NICOTINE PATCH (DIN for billing purposes only) Z

10MG INHALATION CARTRIDGE  
02241742 NICORETTE INHALER Z

**NICOTINE BITARTRATE**

1MG LOZENGE  
80007461 THRIVE LOZENGE Z

2MG LOZENGE  
80007464 THRIVE LOZENGE Z

**NICOTINE POLACRILEX**

2MG GUM  
00999976 NICOTINE GUM (DIN for billing purposes only) Z

4MG GUM  
00999980 NICOTINE GUM (DIN for billing purposes only) Z

2MG LOZENGE  
02247347 NICORETTE LOZENGE Z

4MG LOZENGE  
02247348 NICORETTE LOZENGE Z

**VARENICLINE TARTRATE**

0.5MG TABLET  
02419882 APO-VARENICLINE APX Z  
02426226 TEVA-VARENICLINE TEV Z  
02542951 NRA-VARENICLINE NRA Z

1MG TABLET  
02419890 APO-VARENICLINE APX Z  
02426234 TEVA-VARENICLINE TEV Z  
02542978 NRA-VARENICLINE NRA Z

0.5MG-1MG TABLET DOSE PACK  
02426781 TEVA-VARENICLINE TEV Z  
02435675 APO-VARENICLINE APX Z  
02542986 NRA-VARENICLINE NRA Z

## **20:04.04 IRON PREPARATIONS**

### **FERROUS GLUCONATE**

300MG (35MG IRON) TABLET			
80000435	NOVO-FERROGLUC	TEV	CNW
00031097	JAMP-FERROUS GLUCONATE	JPC	CNW

### **FERROUS SULFATE**

30MG (6MG IRON)/ML ORAL LIQUID			
80008295	JAMP-FERROUS SULFATE	JPC	CNW

75MG (15MG IRON)/ML ORAL DROPS			
02237385	FERODAN INFANT	ODN	W
80008309	JAMP-FERROUS SULFATE	JPC	W

300MG (60MG IRON) TABLET			
00031100	JAMP-FERROUS SULFATE	JPC	CNW

## **20:12.04 ANTI COAGULANTS**

### **⑤APIXABAN**

#### **2.5MG TABLET**

02377233	ELIQUIS	BMS	FNQSW
02484994	TEVA-APIXABAN	TEV	FGNQSW
02486806	AURO-APIXABAN	ARO	FGNQSW
02487381	APO-APIXABAN	APX	FGNQSW
02487713	ACH-APIXABAN	ACH	FGNQSW
02489228	SANDOZ-APIXABAN	SDZ	FGNQSW
02492369	MAR-APIXABAN	MAR	FGNQSW
02492814	NAT-APIXABAN	NAT	FGNQSW
02495430	MINT-APIXABAN	MNT	FGNQSW
02510464	TARO-APIXABAN	TAR	FGNQSW
02528924	JAMP-APIXABAN	JPC	FGNQSW
02529009	M-APIXABAN	MRA	FGNQSW
02530708	APIXABAN	SIV	FGNQSW

#### **5MG TABLET**

02397714	ELIQUIS	BMS	FNQSW
02485001	TEVA-APIXABAN	TEV	FGNQSW
02486814	AURO-APIXABAN	ARO	FGNQSW
02487403	APO-APIXABAN	APX	FGNQSW

02487721	ACH-APIXABAN	ACH	<b>FGNQSW</b>
02489236	SANDOZ-APIXABAN	SDZ	<b>FGNQSW</b>
02492377	MAR-APIXABAN	MAR	<b>FGNQSW</b>
02492822	NAT-APIXABAN	NAT	<b>FGNQSW</b>
02495449	MINT-APIXABAN	MNT	<b>FGNQSW</b>
02510472	TARO-APIXABAN	TAR	<b>FGNQSW</b>
02528932	JAMP-APIXABAN	JPC	<b>FGNQSW</b>
02529017	M-APIXABAN	MRA	<b>FGNQSW</b>
02530716	APIXABAN	SIV	<b>FGNQSW</b>

**⑤ DABIGATRAN**

[SEE APPENDIX A](#) FOR SA CRITERIA  
110MG CAPSULE

02312441	PRADAXA (SA)	BOE	<b>FNQSW</b>
02468905	APO-DABIGATRAN (SA)	APX	<b>FGNQSW</b>

150MG CAPSULE

02358808	PRADAXA (SA)	BOE	<b>FNQSW</b>
02468913	APO-DABIGATRAN (SA)	APX	<b>FGNQSW</b>

**⑤ DALTEPARIN**

[SEE APPENDIX A](#) FOR SA CRITERIA (NURSING HOME PROGRAM DOES NOT REQUIRE AN SA REQUEST)

PRE-FILLED SYRINGE 2,500 IU/0.2ML

02132621	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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PRE-FILLED SYRINGE 5,000 IU/0.2ML

02132648	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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PRE-FILLED SYRINGE 7500 IU/0.3ML

02352648	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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PRE-FILLED SYRINGE 10,000 IU/0.4ML

02352656	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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PRE-FILLED SYRINGE 12,500 IU/0.5ML

02352664	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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PRE-FILLED SYRINGE 15,000 IU/0.6ML

02352672	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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PRE-FILLED SYRINGE 18,000 UNITS/0.72ML

02352680	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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MULTIDOSE VIAL 25,000 IU/ML (3.8ML)

02231171	FRAGMIN (SA)	PFI	<b>FNQSW</b>
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**⑤ EDOXABAN**

15MG TABLET

02458640 LIXIANA SER **FNQSW**

30MG TABLET

02458659 LIXIANA SER **FNQSW**

60MG TABLET

02458667 LIXIANA SER **FNQSW**

**⑤ ENOXAPARIN**

PRE-FILLED SYRINGE 20MG/.02ML

02506440 NOROMBY JUN **FNQSW**

PRE-FILLED SYRINGE 30MG/0.3ML

02506459 NOROMBY JUN **FNQSW**

02507501 INCLUNOX SDZ **FNQSW**

02509075 REDESCA VAL **FNQSW**

02532247 ELONOX FKB **FNQSW**

PRE-FILLED SYRINGE 40MG/0.4ML

02506467 NOROMBY JUN **FNQSW**

02507528 INCLUNOX SDZ **FNQSW**

02509083 REDESCA VAL **FNQSW**

02532255 ELONOX FKB **FNQSW**

PRE-FILLED SYRINGE 60MG/0.6ML

02506475 NOROMBY JUN **FNQSW**

02507536 INCLUNOX SDZ **FNQSW**

02509091 REDESCA VAL **FNQSW**

02532263 ELONOX FKB **FNQSW**

PRE-FILLED SYRINGE 80MG/0.8ML

02506483 NOROMBY JUN **FNQSW**

02507544 INCLUNOX SDZ **FNQSW**

02509105 REDESCA VAL **FNQSW**

02532271 ELONOX FKB **FNQSW**

PRE-FILLED SYRINGE 100MG/1.0ML

02506491 NOROMBY JUN **FNQSW**

02507552	INCLUNOX	SDZ	<b>FNQSW</b>
02509113	REDESCA	VAL	<b>FNQSW</b>
02532298	ELONOX	FKB	<b>FNQSW</b>

PRE-FILLED SYRINGE 120MG/0.8ML

02506505	NOROMBY	JUN	<b>FNQSW</b>
02507560	INCLUNOX	SDZ	<b>FNQSW</b>
02509148	REDESCA	VAL	<b>FNQSW</b>
02532301	ELONOX	FKB	<b>FNQSW</b>

PRE-FILLED SYRINGE 150MG/1.0ML

02506513	NOROMBY	JUN	<b>FNQSW</b>
02507579	INCLUNOX	SDZ	<b>FNQSW</b>
02509156	REDESCA	VAL	<b>FNQSW</b>
02532328	ELONOX	FKB	<b>FNQSW</b>

MULIDOSE VIAL

02509121	REDESCA	VAL	<b>FNQSW</b>
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**HEPARIN**

100 IU/ML LOCK FLUSH SOLUTION

00727520	HEPARIN	LEO	<b>NQ</b>
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1000 IU/ML VIAL

00453811	HEPARIN	LEO	<b>NQ</b>
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**⑤ TINZAPARIN**

[SEE APPENDIX A](#) FOR SA CRITERIA (NURSING HOME PROGRAM DOES NOT REQUIRE AN SA REQUEST)

2500 IU/0.25ML SYRINGE

02229755	INNOHEP (SA)	LEO	<b>FNQSW</b>
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3500 IU/0.35ML SYRINGE

02358158	INNOHEP (SA)	LEO	<b>FNQSW</b>
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4500 IU/0.45ML SYRINGE

02358166	INNOHEP (SA)	LEO	<b>FNQSW</b>
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8000 IU/0.4ML SYRINGE

02429462	INNOHEP (SA)	LEO	<b>FNQSW</b>
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10000 IU/0.5ML SYRINGE

02231478	INNOHEP (SA)	LEO	<b>FNQSW</b>
12000 IU/0.6ML SYRINGE			
02429470	INNOHEP (SA)	LEO	<b>FNQSW</b>
14000 IU/0.7ML SYRINGE			
02358174	INNOHEP (SA)	LEO	<b>FNQSW</b>
16000 IU/0.8ML SYRINGE			
02429489	INNOHEP (SA)	LEO	<b>FNQSW</b>
18000 IU/0.9ML SYRINGE			
02358182	INNOHEP (SA)	LEO	<b>FNQSW</b>
10000 IU/ML MULTIDOSE VIAL			
02167840	INNOHEP (SA)	LEO	<b>FNQSW</b>
20000 IU/ML MULTIDOSE VIAL			
02229515	INNOHEP (SA)	LEO	<b>FNQSW</b>
<b>⑤WARFARIN</b>			
1MG TABLET			
02242680	TARO-WARFARIN	TAR	<b>FGNQSW</b>
02242924	APO-WARFARIN	APX	<b>FGNQSW</b>
2MG TABLET			
02242681	TARO-WARFARIN	TAR	<b>FGNQSW</b>
02242925	APO-WARFARIN	APX	<b>FGNQSW</b>
2.5MG TABLET			
02242682	TARO-WARFARIN	TAR	<b>FGNQSW</b>
02242926	APO-WARFARIN	APX	<b>FGNQSW</b>
3MG TABLET			
02242683	TARO-WARFARIN	TAR	<b>FGNQSW</b>
02245618	APO-WARFARIN	APX	<b>FGNQSW</b>
4MG TABLET			
02242684	TARO-WARFARIN	TAR	<b>FGNQSW</b>
02242927	APO-WARFARIN	APX	<b>FGNQSW</b>
5MG TABLET			
02242685	TARO-WARFARIN	TAR	<b>FGNQSW</b>
02242928	APO-WARFARIN	APX	<b>FGNQSW</b>
10MG TABLET			

02242687	TARO-WARFARIN	TAR	<b>FGNQSW</b>
02242929	APO-WARFARIN	APX	<b>FGNQSW</b>

**Ⓢ RIVAROXABAN**

**2.5MG TABLET**

02480808	XARELTO	BAY	<b>FNQSW</b>
02524503	REDDY-RIVAROXABAN	RCH	<b>FGNQSW</b>
02526786	TARO-RIVAROXABAN	TAR	<b>FGNQSW</b>
02527537	PMS-RIVAROXABAN	PMS	<b>FGNQSW</b>
02537877	SANDOZ-RIVOROXABAN	SDZ	<b>FGNQSW</b>
02541467	RIVAROXABAN	SIV	<b>FGNQSW</b>
02541734	APO-RIVAROXABAN	APX	<b>FGNQSW</b>

**10MG TABLET**

02316986	XARELTO	BAY	<b>FNQSW</b>
02470497	APO-RIVAROXABAN	APX	<b>FGNQSW</b>
02472414	REDDY-RIVAROXABAN	RCH	<b>FGNQSW</b>
02482223	SANDOZ-RIVAROXABAN	SDZ	<b>FGNQSW</b>
02483807	TARO-RIVAROXABAN	TAR	<b>FGNQSW</b>
02507196	TEVA-RIVAROXABAN	TEV	<b>FGNQSW</b>
02512041	PMS-RIVAROXABAN	PMS	<b>FGNQSW</b>
02516292	JAMP-RIVAROXABAN	JPC	<b>FGNQSW</b>
02541475	RIVAROXABAN	SIV	<b>FGNQSW</b>

**15MG TABLET**

02378604	XARELTO	BAY	<b>FNQSW</b>
02470500	APO-RIVAROXABAN	APX	<b>FGNQSW</b>
02472430	REDDY-RIVAROXABAN	RCH	<b>FGNQSW</b>
02482231	SANDOZ-RIVAROXABAN	SDZ	<b>FGNQSW</b>
02483815	TARO-RIVAROXABAN	TAR	<b>FGNQSW</b>
02507218	TEVA-RIVAROXABAN	TEV	<b>FGNQSW</b>
02512068	PMS-RIVAROXABAN	PMS	<b>FGNQSW</b>
02516306	JAMP-RIVAROXABAN	JPC	<b>FGNQSW</b>
02541483	RIVAROXABAN	SIV	<b>FGNQSW</b>

**20MG TABLET**

02378612	XARELTO	BAY	<b>FNQSW</b>
02470519	APO-RIVAROXABAN	APX	<b>FGNQSW</b>
02472422	REDDY-RIVAROXABAN	RCH	<b>FGNQSW</b>
02482258	SANDOZ-RIVAROXABAN	SDZ	<b>FGNQSW</b>
02483823	TARO-RIVAROXABAN	TAR	<b>FGNQSW</b>
02507226	TEVA-RIVAROXABAN	TEV	<b>FGNQSW</b>
02512076	PMS-RIVAROXABAN	PMS	<b>FGNQSW</b>
02516314	JAMP-RIVAROXABAN	JPC	<b>FGNQSW</b>
02541491	RIVAROXABAN	SIV	<b>FGNQSW</b>



## **20:12.14 PLATELET REDUCING AGENTS**

### **ANAGRELIDE HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.5MG CAPSULE

02236859	AGRYLIN (SA)	SHR	<b>FNQSW</b>
02274949	PMS-ANAGRELIDE (SA)	PMS	<b>FGNQSW</b>

## **20:12.18 PLATELET AGGREGATION INHIBITORS**

### **⑤ CLOPIDOGREL BISULFATE**

75MG TABLET

02238682	PLAVIX	AVN	<b>FNQSW</b>
02293161	TEVA-CLOPIDOGREL	TEV	<b>FGNQSW</b>
02252767	APO-CLOPIDOGREL	APX	<b>FGNQSW</b>
02303027	ACT-CLOPIDOGREL	TEV	<b>FGNQSW</b>
02348004	PMS-CLOPIDOGREL	PMS	<b>FGNQSW</b>
02379813	RAN-CLOPIDOGREL	RAN	<b>FGNQSW</b>
02385813	CLOPIDOGREL	SIV	<b>FGNQSW</b>
02400553	CLOPIDOGREL	SNS	<b>FGNQSW</b>
02408910	MINT-CLOPIDOGREL	MNT	<b>FGNQSW</b>
02415550	JAMP-CLOPIDOGREL	JPC	<b>FGNQSW</b>
02416387	AURO-CLOPIDOGREL	ARO	<b>FGNQSW</b>
02422255	MAR-CLOPIDOGREL	MAR	<b>FGNQSW</b>
02482037	NRA-CLOPIDOGREL	NRA	<b>FGNQSW</b>
02502283	M-CLOPIDOGREL	MRA	<b>FGNQSW</b>

### **PRASUGREL**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET

02502429	JAMP-PRASUGREL (SA)	JPC	<b>FGNQSW</b>
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### **⑤ TICAGRELOR**

[SEE APPENDIX A](#) FOR SA CRITERIA

90MG TABLET

02368544	BRILINTA (SA)	AZE	<b>FNQSW</b>
02482630	APO-TICAGRELOR (SA)	APX	<b>FGNQSW</b>
02492598	TARO-TICAGRELOR (SA)	TAR	<b>FGNQSW</b>
02529769	M-TICAGRELOR (SA)	MRA	<b>FGNQSW</b>
02531801	JAMP-TICAGRELOR (SA)	JPC	<b>FGNQSW</b>

**TICLOPIDINE HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA  
250MG TABLET  
02237701 TICLOPIDINE (SA)

AAA FGNQSW

**20:16.00 HEMATOPOIETIC AGENTS**

**DARBEPOETIN ALFA**

[SEE APPENDIX A](#) FOR SA CRITERIA  
10MCG/ML PRE-FILLED SYRINGE  
02392313 ARANESP (SA)

AMG E

20MCG/ML PRE-FILLED SYRINGE  
02392321 ARANESP (SA)

AMG E

30MCG/ML PRE-FILLED SYRINGE  
02392348 ARANESP (SA)

AMG E

40MCG/ML PRE-FILLED SYRINGE  
02391740 ARANESP (SA)

AMG E

50MCG/ML PRE-FILLED SYRINGE  
02391759 ARANESP (SA)

AMG E

60MCG/ML PRE-FILLED SYRINGE  
02392356 ARANESP (SA)

AMG E

80MCG/ML PRE-FILLED SYRINGE  
02391767 ARANESP (SA)

AMG E

100MCG/ML PRE-FILLED SYRINGE  
02391775 ARANESP (SA)

AMG E

150MCG/ML PRE-FILLED SYRINGE  
02391791 ARANESP (SA)

AMG E

200MCG/ML PRE-FILLED SYRINGE  
02391805 ARANESP (SA)

AMG E

500MCG/ML PRE-FILLED SYRINGE  
02392364 ARANESP (SA)

AMG E

**EPOETIN ALFA**[SEE APPENDIX A](#) FOR SA CRITERIA

4000IU/0.4ML PRE-FILLED SYRINGE

02231586 EPREX (SA) JAN E

6000IU/0.6ML PRE-FILLED SYRINGE

02243401 EPREX (SA) JAN E

8000IU/0.8ML PRE-FILLED SYRINGE

02243403 EPREX (SA) JAN E

10,000IU/1.0ML PRE-FILLED SYRINGE

02231587 EPREX (SA) JAN E

**FILGRASTIM**[SEE APPENDIX A](#) FOR SA CRITERIA

300 MCG/0.5ML PREFILLED SYRINGE

02441489 GRASTOFIL (SA) APX **NMQW**02485575 NIVESTYM (SA) PFI **NMQW**02520990 NYPOZI (SA) TAV **NMQW**

300 MCG/ML VIAL

02485591 NIVESTYM (SA) PFI **NMQW**

480 MCG/0.8ML PREFILLED SYRINGE

02454548 GRASTOFIL (SA) APX **NMQW**02485583 NIVESTYM (SA) PFI **NMQW**02521008 NYPOZI (SA) TAV **NMQW**

480 MCG/1.6ML VIAL

02485656 NIVESTYM (SA) PFI **NMQW****LUSPATERCEPT**[SEE APPENDIX A](#) FOR SA CRITERIA

25 MG VIAL

02505541 REBLOZYL (SA) CEL **NMQW**00904728 REBLOZYL (SA)\* **NMQW**

75 MG VIAL

02505568 REBLOZYL (SA) CEL **NMQW**00904729 REBLOZYL (SA)\* **NMQW**

\*use when drug cost in excess of CPHA maximum

**PEGFILGRASTIM**

[SEE APPENDIX A](#) FOR SA CRITERIA

6 MG/0.6ML PREFILLED SYRINGE

02474565	LAPELGA (SA)	APX	NMQW
02484153	FULPHILA (SA)	BGP	NMQW
02497395	ZIEXTENZO (SA)	SDZ	NMQW
02506238	NYVEPRIA (SA)	PFI	NMQW

6 MG/0.6ML AUTOINJECTOR

02529343	LAPELGA (SA)	APX	NMQW
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**20:24.00 HEMORRHEOLOGIC AGENTS**

**⑤PENTOXIFYLLINE**

400 MG SUSTAINED RELEASE TABLET

02230090	PENTOXIFYLLINE SR	AAA	FGNQSW
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**20:28.00 HEMOSTATIC AGENTS**

TRANEXAMIC ACID

500 MG TABLET

02401231	TRANEXAMIC ACID	STE	FGNQSW
02496232	MAR-TRANEXAMIC ACID	MAR	FGNQSW
02519194	TRAMEXAMIC ACID	JPC	FGNQSW

**24:00.00 CARDIAC DRUGS**

**⑤ACEBUTOLOL HCL**

100 MG TABLET

02147602	APO-ACEBUTOLOL	APX	FGNQSW
02204517	TEVA-ACEBUTOLOL	TEV	FGNQSW

200 MG TABLET

02147610	APO-ACEBUTOLOL	APX	FGNQSW
02204525	TEVA-ACEBUTOLOL	TEV	FGNQSW

400 MG TABLET

02147629	APO-ACEBUTOLOL	APX	FGNQSW
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02204533 TEVA-ACEBUTOLOL TEV FGNQSW

**⑤ AMIODARONE**

100 MG TABLET

02292173 PMS-AMIODARONE PMS FGNQSW

200 MG TABLET

02239835 TEVA-AMIODARONE TEV FGNQSW

02242472 PMS-AMIODARONE PMS FGNQSW

02243836 SANDOZ-AMIODARONE SDZ FGNQSW

02246194 APO-AMIODARONE APX FGNQSW

02364336 SANIS-AMIODARONE SNS FGNQSW

02385465 AMIODARONE SIV FGNQSW

02531844 JAMP-AMIODARONE JPC FGNQSW

**⑤ AMLODIPINE BESYLATE**

2.5 MG TABLET

02297477 ACT-AMLODIPINE TEV FGNQSW

02295148 PMS-AMLODIPINE PMS FGNQSW

02330474 SANDOZ-AMLODIPINE SDZ FGNQSW

02357186 JAMP-AMLODIPINE JPC FGNQSW

02371707 MAR-AMLODIPINE MAR FGNQSW

02385783 AMLODIPINE SIV FGNQSW

02419556 AMLODIPINE BESYLATE ACH FGNQSW

02468018 M-AMLODIPINE MRA FGNQSW

02469022 PHARMA-AMLODIPINE PMS FGNQSW

02476452 NRA-AMLODIPINE NRA FGNQSW

02478587 AMLODIPINE SNS FGNQSW

02492199 AMLODIPINE JPC FGNQSW

02522500 PRZ-AMLODIPINE PRZ FGNQSW

5 MG TABLET

00878928 NORVASC UJC FNQSW

02272113 MYLAN-AMLODIPINE MYL FGNQSW

02273373 APO-AMLODIPINE APX FGNQSW

02284065 PMS-AMLODIPINE PMS FGNQSW

02284383 SANDOZ-AMLODIPINE SDZ FGNQSW

02297485 ACT-AMLODIPINE TEV FGNQSW

02321858 RAN-AMLODIPINE RAN FGNQSW

02331284 SANIS-AMLODIPINE SNS FGNQSW

02357194 JAMP-AMLODIPINE JPC FGNQSW

02385791 AMLODIPINE SIV FGNQSW

02362651 MINT-AMLODIPINE MNT FGNQSW

02371715 MAR-AMLODIPINE MAR FGNQSW

02397072 AURO- AMLODIPINE ARO FGNQSW

02419564 AMLODIPINE BESYLATE ACH FGNQSW

02429217	AMLODIPINE	JPC	FGNQSW
02468026	M-AMLODIPINE	MRA	FGNQSW
02476460	NRA-AMLODIPINE	NRA	FGNQSW
02522519	PRZ-AMLODIPINE	PRZ	FGNQSW

10 MG TABLET

00878936	NORVASC	UJC	FNQSW
02272121	MYLAN-AMLODIPINE	MYL	FGNQSW
02273381	APO-AMLODIPINE	APX	FGNQSW
02284073	PMS-AMLODIPINE	PMS	FGNQSW
02284391	SANDOZ-AMLODIPINE	SDZ	FGNQSW
02297493	ACT-AMLODIPINE	TEV	FGNQSW
02321866	RAN-AMLODIPINE	RAN	FGNQSW
02331292	SANIS-AMLODIPINE	SNS	FGNQSW
02357208	JAMP-AMLODIPINE	JPC	FGNQSW
02362678	MINT-AMLODIPINE	MNT	FGNQSW
02371723	MAR-AMLODIPINE	MAR	FGNQSW
02385805	AMLODIPINE	SIV	FGNQSW
02397080	AURO-AMLODIPINE	ARO	FGNQSW
02419572	AMLODIPINE BESYLATE	ACH	FGNQSW
02429225	AMLODIPINE	JPC	FGNQSW
02468034	M-AMLODIPINE	MRA	FGNQSW
02476479	NRA-AMLODIPINE	NRA	FGNQSW
02522527	PRZ-AMLODIPINE	PRZ	FGNQSW

[SEE APPENDIX A](#) FOR SA CRITERIA

1 MG/ML ORAL SOLUTION

02484706	PDP-AMLODIPINE	PEN	FGNQSW
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⑤ **ATENOLOL**

25 MG TABLET

02246581	PMS-ATENOLOL	PMS	FGNQSW
02266660	TEVA-ATENOL	TEV	FGNQSW
02367556	JAMP-ATENOLOL	JPC	FGNQSW
02371979	MAR-ATENOLOL	MAR	FGNQSW
02373963	RAN-ATENOLOL	RAN	FGNQSW
02541564	ATENOLOL	SIV	FGNQSW

50 MG TABLET

00773689	APO-ATENOL	APX	FGNQSW
02039532	TENORMIN	AZE	FNQSW
02171791	TEVA-ATENOLOL	TEV	FGNQSW
02237600	PMS-ATENOLOL	PMS	FGNQSW
02238316	ATENOLOL	SIV	FGNQSW
02267985	RAN-ATENOLOL	RAN	FGNQSW
02367564	JAMP-ATENOLOL	JPC	FGNQSW

02368021	MINT-ATENOL	MNT	FGNQSW
02369184	AG-ATENOLOL	ANG	FGNQSW
02371987	MAR-ATENOLOL	MAR	FGNQSW
02466465	ATENOLOL	SNS	FGNQSW

100 MG TABLET

00773697	APO-ATENOL	APX	FGNQSW
02039540	TENORMIN	AZE	FNQSW
02171805	TEVA-ATENOLOL	TEV	FGNQSW
02237601	PMS-ATENOLOL	PMS	FGNQSW
02238318	ATENOLOL	SIV	FGNQSW
02267993	RAN-ATENOLOL	RAN	FGNQSW
02367572	JAMP-ATENOLOL	JPC	FGNQSW
02368048	MINT-ATENOL	MNT	FGNQSW
02369192	AG-ATENOLOL	ANG	FGNQSW
02371995	MAR-ATENOL	MAR	FGNQSW
02466473	ATENOLOL	SNS	FGNQSW

**⑤ ATENOLOL & CHLORTHALIDONE**

50 MG & 25MG TABLET

02248763	AA-ATENIDONE	AAA	FGNQSW
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100MG & 25MG TABLET

02248764	AA-ATENIDONE	AAA	FGNQSW
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**⑤ BISOPROLOL**

5MG TABLET

02256134	APO-BISOPROLOL	APX	FGNQSW
02267470	TEVA-BISOPROLOL	TEV	FGNQSW
02391589	BISOPROLOL	SNS	FGNQSW
02465612	MINT-BISOPROLOL	MNT	FGNQSW
02494035	SANDOZ-BISOPROLOLS	SDZ	FGNQSW
02495562	BISOPROLOL	SIV	FGNQSW
02518805	JAMP-BISOPROLOL	JPC	FGNQSW

10MG TABLET

02256177	APO-BISOPROLOL	APX	FGNQSW
02267489	TEVA-BISOPROLOL	TEV	FGNQSW
02391597	BISOPROLOL	SNS	FGNQSW
02465620	MINT-BISOPROLOL	MNT	FGNQSW
02494043	SANDOZ-BISOPROLOL	SDZ	FGNQSW
02495570	BISOPROLOL	SIV	FGNQSW
02518791	JAMP-BISOPROLOL	JPC	FGNQSW

**⑤ CARVEDILOL**

3.125MG TABLET

02245914	PMS-CARVEDILOL	PMS	<b>FGNQSW</b>
02247933	APO-CARVEDILOL	APX	<b>FGNQSW</b>
02248752	CARVEDILOL	SIV	<b>FGNQSW</b>
02252309	TEVA-CARVEDILOL	TEV	<b>FGNQSW</b>
02364913	CARVEDILOL	SNS	<b>FGNQSW</b>
02368897	JAMP-CARVEDILOL	JPC	<b>FGNQSW</b>
02418495	AURO-CARVEDILOL	ARO	<b>FGNQSW</b>

**6.25MG TABLET**

02245915	PMS-CARVEDILOL	PMS	<b>FGNQSW</b>
02247934	APO-CARVEDILOL	APX	<b>FGNQSW</b>
02248753	CARVEDILOL	SIV	<b>FGNQSW</b>
02252317	TEVA-CARVEDILOL	TEV	<b>FGNQSW</b>
02364921	CARVEDILOL	SNS	<b>FGNQSW</b>
02368900	JAMP-CARVEDILOL	JPC	<b>FGNQSW</b>
02418509	AURO-CARVEDILOL	ARO	<b>FGNQSW</b>

**12.5MG TABLET**

02245916	PMS-CARVEDILOL	PMS	<b>FGNQSW</b>
02247935	APO-CARVEDILOL	APX	<b>FGNQSW</b>
02248754	CARVEDILOL	SIV	<b>FGNQSW</b>
02252325	TEVA-CARVEDILOL	TEV	<b>FGNQSW</b>
02364948	CARVEDILOL	SNS	<b>FGNQSW</b>
02368919	JAMP-CARVEDILOL	JPC	<b>FGNQSW</b>
02418517	AURO-CARVEDILOL	ARO	<b>FGNQSW</b>

**25MG TABLET**

02245917	PMS-CARVEDILOL	PMS	<b>FGNQSW</b>
02247936	APO-CARVEDILOL	APX	<b>FGNQSW</b>
02248755	CARVEDILOL	SIV	<b>FGNQSW</b>
02252333	TEVA-CARVEDILOL	TEV	<b>FGNQSW</b>
02364956	CARVEDILOL	SNS	<b>FGNQSW</b>
02368927	JAMP-CARVEDILOL	JPC	<b>FGNQSW</b>
02418525	AURO-CARVEDILOL	ARO	<b>FGNQSW</b>

**⑤DIGOXIN**

**0.05MG/ML ELIXIR**

02242320	PMS-DIGOXIN	PMS	<b>FGNQSW</b>
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**0.0625MG TABLET**

02335700	PMS-DIGOXIN	PMS	<b>FGNQSW</b>
02498502	JAMP-DIGOXIN	JPC	<b>FGNQSW</b>

**0.125MG TABLET**

02335719	PMS-DIGOXIN	PMS	<b>FGNQSW</b>
02498510	JAMP-DIGOXIN	JPC	<b>FGNQSW</b>



0.25 MG/ML INJECTION SOLUTION  
02048264 DIGOXIN

SDZ NQ

**⑤DILTIAZEM**

120MG EXTENDED RELEASE CAPSULE

02231150	TIAZAC	VAL	<b>FNQSW</b>
02245918	SANDOZ-DILTIAZEM T	SDZ	<b>FGNQSW</b>
02271605	TEVA-DILTIAZEM ER	TEV	<b>FGNQSW</b>
02370441	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02465353	MAR-DILTIAZEM T	MAR	<b>FGNQSW</b>
02495376	JAMP-DILTIAZEM T	JPC	<b>FGNQSW</b>
02516101	DILTIAZEM T	SNS	<b>FGNQSW</b>

180MG EXTENDED RELEASE CAPSULE

02231151	TIAZAC	VAL	<b>FNQSW</b>
02245919	SANDOZ-DILTIAZEM T	SDZ	<b>FGNQSW</b>
02271613	TEVA-DILTIAZEM ER	TEV	<b>FGNQSW</b>
02370492	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02465361	MAR-DILTIAZEM T	MAR	<b>FGNQSW</b>
02495384	JAMP-DILTIAZEM T	JPC	<b>FGNQSW</b>
02516128	DILTIAZEM T	SNS	<b>FGNQSW</b>

240MG EXTENDED RELEASE CAPSULE

02231152	TIAZAC	VAL	<b>FNQSW</b>
02271621	TEVA-DILTIAZEM ER	TEV	<b>FGNQSW</b>
02370506	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02465388	MAR-DILTIAZEM T	MAR	<b>FGNQSW</b>
02495392	JAMP-DILTIAZEM T	JPC	<b>FGNQSW</b>
02516136	DILTIAZEM T	SNS	<b>FGNQSW</b>

300MG EXTENDED RELEASE CAPSULE

02231154	TIAZAC	VAL	<b>FNQSW</b>
02271648	TEVA-DILTIAZEM ER	TEV	<b>FGNQSW</b>
02370514	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02465396	MAR-DILTIAZEM T	MAR	<b>FGNQSW</b>
02495406	JAMP-DILTIAZEM T	JPC	<b>FGNQSW</b>
02516144	DILTIAZEM T	SNS	<b>FGNQSW</b>

360MG EXTENDED RELEASE CAPSULE

02231155	TIAZAC	VAL	<b>FNQSW</b>
02271656	TEVA-DILTIAZEM ER	TEV	<b>FGNQSW</b>
02370522	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02465418	MAR-DILTIAZEM T	MAR	<b>FGNQSW</b>
02495414	JAMP-DILTIAZEM T	JPC	<b>FGNQSW</b>
02516152	DILTIAZEM T	SNS	<b>FGNQSW</b>

120MG EXTENDED RELEASE TABLET			
02256738	TIAZAC XC	VAL	<b>FNQSW</b>
180MG EXTENDED RELEASE TABLET			
02256746	TIAZAC XC	VAL	<b>FNQSW</b>
02429322	TEVA-DILTIAZEM XC	TEV	<b>FGNQSW</b>
240MG EXTENDED RELEASE TABLET			
02256754	TIAZAC XC	VAL	<b>FNQSW</b>
02429330	TEVA-DILTIAZEM XC	TEV	<b>FGNQSW</b>
300MG EXTENDED RELEASE TABLET			
02256762	TIAZAC XC	VAL	<b>FNQSW</b>
02429349	TEVA-DILTIAZEM XC	TEV	<b>FGNQSW</b>
360MG EXTENDED RELEASE TABLET			
02256770	TIAZAC XC	BVL	<b>FNQSW</b>
02429357	TEVA-DILTIAZEM XC	TEV	<b>FGNQSW</b>
120MG CONTROLLED DELIVERY CAPSULE			
02230997	APO-DILTIAZ CD	APX	<b>FGNQSW</b>
02242538	TEVA-DILTIAZEM CD	TEV	<b>FGNQSW</b>
02243338	SANDOZ-DILTIAZEM CD	SDZ	<b>FGNQSW</b>
02370611	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02400421	DILTIAZEM CD	SNS	<b>FGNQSW</b>
02445999	DILTIAZEM CD	SIV	<b>FGNQSW</b>
02484064	MAR-DILTIAZEM CD	MAR	<b>FGNQSW</b>
02528037	JAMP-DILTIAZEM CD	JPC	<b>FGNQSW</b>
180MG CONTROLLED DELIVERY CAPSULE			
02230998	APO-DILTIAZ CD	APX	<b>FGNQSW</b>
02242539	TEVA-DILTIAZEM CD	TEV	<b>FGNQSW</b>
02243339	SANDOZ-DILTIAZEM CD	SDZ	<b>FGNQSW</b>
02370638	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02400448	DILTIAZEM CD	SNS	<b>FGNQSW</b>
02446006	DILTIAZEM CD	SIV	<b>FGNQSW</b>
02484072	MAR-DILTIAZEM CD	MAR	<b>FGNQSW</b>
02528045	JAMP-DILTIAZEM CD	JPC	<b>FGNQSW</b>
240MG CONTROLLED DELIVERY CAPSULE			
02230999	APO-DILTIAZ CD	APX	<b>FGNQSW</b>
02242540	TEVA-DILTIAZEM CD	TEV	<b>FGNQSW</b>
02243340	SANDOZ-DILTIAZEM CD	SDZ	<b>FGNQSW</b>
02400456	DILTIAZEM CD	SNS	<b>FGNQSW</b>
02446014	DILTIAZEM CD	SIV	<b>FGNQSW</b>

02484080	MAR-DILTIAZEM CD	MAR	<b>FGNQSW</b>
02528053	JAMP-DILTIAZEM CD	JPC	<b>FGNQSW</b>

300MG CONTROLLED DELIVERY CAPSULE

02229526	APO-DILTIAZ CD	APX	<b>FGNQSW</b>
02242541	TEVA-DILTAZEM CD	TEV	<b>FGNQSW</b>
02243341	SANDOZ-DILTIAZEM CD	SDZ	<b>FGNQSW</b>
02370654	ACT-DILTIAZEM	TEV	<b>FGNQSW</b>
02400464	DILTIAZEM CD	SNS	<b>FGNQSW</b>
02446022	DILTIAZEM CD	SIV	<b>FGNQSW</b>
02484099	MAR-DILTIAZEM CD	MAR	<b>FGNQSW</b>
02528061	JAMP-DILTIAZEM CD	JPC	<b>FGNQSW</b>

30MG TABLET

00771376	AA-DILTIAZ	AAA	<b>FGNQSW</b>
00862924	TEVA-DILTAZEM	TEV	<b>FGNQSW</b>

60MG TABLET

00771384	AA-DILTIAZ	AAA	<b>FGNQSW</b>
00862932	TEVA-DILTAZEM	TEV	<b>FGNQSW</b>

**⑤DISOPYRAMIDE**

100MG CAPSULE

02224801	RYTHMODAN	XPI	<b>FNQSW</b>
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**⑤FLECAINIDE ACETATE**

50MG TABLET

02275538	FLECAINIDE	APX	<b>FGNQSW</b>
02459957	AURO-FLECAINIDE	ARO	<b>FGNQSW</b>
02476177	MAR-FLECAINIDE	MAR	<b>FGNQSW</b>
02493705	JAMP-FLECAINIDE	JPC	<b>FGNQSW</b>
02534800	FLECAINIDE	SNS	<b>FGNQSW</b>

100MG TABLET

02275546	FLECAINIDE	APX	<b>FGNQSW</b>
02459965	AURO-FLECAINIDE	ARO	<b>FGNQSW</b>
02476185	MAR-FLECAINIDE	MAR	<b>FGNQSW</b>
02493713	JAMP-FLECAINIDE	JPC	<b>FGNQSW</b>
02534819	FLECAINIDE	SNS	<b>FGNQSW</b>

**⑤IVABRADINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG TABLET

02459973	LANCORA (SA)	SER	<b>FNQSW</b>
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7.5MG TABLET

02459981	LANCORA (SA)	SER	<b>FNQSW</b>
<b>⑤ LABETALOL HCL</b>			
100MG TABLET			
02106272	TRANDATE	PAL	<b>FNQSW</b>
02243538	APO-LABETALOL	APX	<b>FGNQSW</b>
02489406	RIVA-LABETALOL	RIV	<b>FGNQSW</b>
200MG TABLET			
02106280	TRANDATE	PAL	<b>FNQSW</b>
02243539	APO-LABETALOL	APX	<b>FGNQSW</b>
02489414	RIVA-LABETALOL	RIV	<b>FGNQSW</b>
<b>⑤ METOPROLOL TARTRATE</b>			
100MG SUSTAINED RELEASE TABLET			
02285169	AA-METOPROLOL SR	AAA	<b>FGNQSW</b>
200MG SUSTAINED RELEASE TABLET			
02285177	AA-METOPROLOL SR	AAA	<b>FGNQSW</b>
25MG TABLET			
02246010	APO-METOPROLOL	APX	<b>FGNQSW</b>
02248855	PMS-METOPROLOL-L	PMS	<b>FGNQSW</b>
02356813	JAMP-METOPROLOL-L	JPC	<b>FGNQSW</b>
50 MG TABLET			
00618632	APO-METOPROLOL	APX	<b>FGNQSW</b>
00648035	TEVA-METOPROL	TEV	<b>FGNQSW</b>
00749354	APO-METOPROLOL (TYPE L)	APX	<b>FGNQSW</b>
00842648	TEVA-METOPROL (UNCOATED)	TEV	<b>FGNQSW</b>
02230803	PMS-METOPROLOL-L	PMS	<b>FGNQSW</b>
02350394	METOPROLOL	SNS	<b>FGNQSW</b>
02356821	JAMP-METOPROLOL-L	JPC	<b>FGNQSW</b>
02442124	METOPROLOL-L	SIV	<b>FGNQSW</b>
02481316	AG-METOPROLOL-L	ANG	<b>FGNQSW</b>
100MG TABLET			
00618640	APO-METOPROLOL	APX	<b>FGNQSW</b>
00648043	TEVA-METOPROL	TEV	<b>FGNQSW</b>
00751170	APO-METOPROLOL (TYPE L)	APX	<b>FGNQSW</b>
00842656	TEVA-METOPROL (UNCOATED)	TEV	<b>FGNQSW</b>
02230804	PMS-METOPROLOL-L	PMS	<b>FGNQSW</b>
02350408	METOPROLOL	SNS	<b>FGNQSW</b>
02356848	JAMP-METOPROLOL-L	JPC	<b>FGNQSW</b>
02442132	METOPROLOL-L	SIV	<b>FGNQSW</b>
02481324	AG-METOPROLOL-L	ANG	<b>FGNQSW</b>

**⑤MEXILETINE HCL**

100MG CAPSULE

02230359 TEVA-MEXILETINE  
02536846 MINT-MEXILETINE

TEV FGNQSW  
MNT FGNQSW

200MG CAPSULE

02230360 TEVA-MEXILETINE  
02536854 MINT-MEXILETINE

TEV FGNQSW  
MNT FGNQSW

**⑤NADOLOL**

40MG TABLET

00782505 APO-NADOLOL  
02496380 MINT-NADOLOL

APX FGNQSW  
MNT FGNQSW

80MG TABLET

00782467 APO-NADOLOL  
02496399 MINT-NADOLOL

APX FGNQSW  
MNT FGNQSW

160MG TABLET

00782475 APO-NADOLOL

APX FGNQSW

**⑤NIFEDIPINE**

5MG CAPSULE

00725110 NIFEDIPINE

AAA FGNQSW

10MG CAPSULE

00755907 NIFEDIPINE

AAA FGNQSW

30MG EXTENDED RELEASE TABLET

02155907 ADALAT XL  
02349167 MYLAN-NIFEDIPINE ER

BAY FNQSW  
MYL FGNQSW

60MG EXTENDED RELEASE TABLET

02321149 MYLAN-NIFEDIPINE ER

MYL FGNQSW

**⑤PINDOLOL**

5MG TABLET

00417270 VISKEN  
00869007 TEVA-PINDOL

XPI FNQSW  
TEV FGNQSW

10MG TABLET

00443174 VISKEN  
00869015 TEVA-PINDOL

XPI FNQSW  
TEV FGNQSW

15MG TABLET

00755893 APO-PINDOL

APX FGNQSW

00869023 TEVA-PINDOL TEV FGNQSW

**⑤PROPAFENONE HCL**

150MG TABLET

00603708	RYTHMOL	BGP	FNQSW
02243324	APO-PROPAFENONE	APX	FGNQSW
02343053	PROPAFENONE	SNS	FGNQSW
02457172	MYLAN-PROPAFENONE	MYL	FGNQSW

300MG TABLET

00603716	RYTHMOL	BGP	FNQSW
02243325	APO-PROPAFENONE	APX	FGNQSW
02343061	PROPAFENONE	SNS	FGNQSW
02457164	MYLAN-PROPAFENONE	MYL	FGNQSW

**⑤PROPRANOLOL**

10MG TABLET

00496480	TEVA-PROPRANOLOL	TEV	FGNQSW
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20MG TABLET

00740675	TEVA-PROPRANOLOL	TEV	FGNQSW
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40MG TABLET

00496499	TEVA-PROPRANOLOL	TEV	FGNQSW
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80MG TABLET

00496502	TEVA-PROPRANOLOL	TEV	FGNQSW
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**PROPRANOLOL ORAL SOLUTION**

[SEE APPENDIX A](#) FOR SA CRITERIA

3.75MG/ML ORAL SOLUTION

02457857	HEMANGIOL (SA)	PFB	FQW
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**⑤SOTALOL HCL**

80MG TABLET

02210428	APO-SOTALOL	APX	FGNQSW
02368617	JAMP-SOTALOL	JPC	FGNQSW
02238326	PMS-SOTALOL	PMS	FGNQSW

160MG TABLET

02167794	APO-SOTALOL	APX	FGNQSW
02238327	PMS-SOTALOL	PMS	FGNQSW
02368625	JAMP-SOTALOL	JPC	FGNQSW

**TAFAMIDIS**

[SEE APPENDIX A](#) FOR SA CRITERIA

61MG TABLET

02517841 VYNDAMAX (SA)  
00904778 VYNDAMAX (SA)\*  
\*use when drug cost in excess of CPHA maximum

PFI MNQW  
MNQW

**TAFAMIDIS MEGLUMINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG CAPSULE

02495732 VYNDAQEL (SA)  
00904637 VYNDAQEL (SA)\*  
\*use when drug cost in excess of CPHA maximum

PFI MNQW  
MNQW

**⑤TIMOLOL MALEATE**

5MG TABLET

00755842 TIMOLOL

AAA FGNQSW

10MG TABLET

00755850 TIMOLOL

AAA FGNQSW

20MG TABLET

00755869 TIMOLOL

AAA FGNQSW

**⑤VERAPAMIL HCL**

80MG TABLET

00782483 APO-VERAP  
02237921 MYLAN-VERAPAMIL

APX FGNQSW  
MYL FGNQSW

120MG TABLET

00782491 APO-VERAP  
02237922 MYLAN-VERAPAMIL

APX FGNQSW  
MYL FGNQSW

120MG SUSTAINED RELEASE TABLET

01907123 ISOPTIN SR  
02210347 MYLAN-VERAPAMIL SR  
02246893 APO-VERAP SR

BGP FNQSW  
MYL FGNQSW  
APX FGNQSW

180MG SUSTAINED RELEASE TABLET

01934317 ISOPTIN SR  
02450488 MYLAN-VERAPAMIL SR

BGP FNQSW  
MYL FGNQSW

240MG SUSTAINED RELEASE TABLET

00742554 ISOPTIN SR  
02450496 MYLAN-VERAPAMIL SR

BGP FNQSW  
MYL FGNQSW

## **24:06.00 ANTILIPEMIC DRUGS**

### **⑤ ALIROCUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

75MG/ML PREFILLED PEN

02453819 PRALUENT (SA) SAV **FNQSW**

150MG/ML PREFILLED PEN

02453835 PRALUENT (SA) SAV **FNQSW**

### **⑤ ATORVASTATIN CALCIUM**

10MG TABLET

02230711	LIPITOR	UJC	<b>FNQSW</b>
02295261	APO-ATORVASTATIN	APX	<b>FGNQSW</b>
02310899	TEVA-ATORVASTATIN	TEV	<b>FGNQSW</b>
02313707	TARO-ATORVASTATIN	SUN	<b>FGNQSW</b>
02324946	SANDOZ-ATORVASTATIN	SDZ	<b>FGNQSW</b>
02348705	ATORVASTATIN	SNS	<b>FGNQSW</b>
02391058	JAMP-ATORVASTATIN	JPC	<b>FGNQSW</b>
02392933	MYLAN-ATORVASTATIN	MYL	<b>FGNQSW</b>
02407256	AURO-ATORVASTATIN	ARO	<b>FGNQSW</b>
02411350	ATORVASTATIN	SIV	<b>FGNQSW</b>
02417936	REDDY-ATORVASTATIN	RCH	<b>FGNQSW</b>
02454017	MAR-ATORVASTATIN	MAR	<b>FGNQSW</b>
02457741	ACH-ATORVASTATIN	ACH	<b>FGNQSW</b>
02471167	M-ATORVASTATIN	MRA	<b>FGNQSW</b>
02475022	ATORVASTATIN	RIV	<b>FGNQSW</b>
02476517	NRA-ATORVASTATIN	NRA	<b>FGNQSW</b>
02477149	PMS-ATORVASTATIN	PMS	<b>FGNQSW</b>
02478145	AG-ATORVASTATIN	ANG	<b>FGNQSW</b>
02479508	MINT-ATORVASTATIN	MNT	<b>FGNQSW</b>
02504197	JAMP-ATORVASTATIN	JPC	<b>FGNQSW</b>
02507234	PMSC-ATORVASTATIN	PMS	<b>FGNQSW</b>
02521555	PRZ-ATORVASTATIN	PRZ	<b>FGNQSW</b>

20MG TABLET

02230713	LIPITOR	UJC	<b>FNQSW</b>
02295288	APO-ATORVASTATIN	APX	<b>FGNQSW</b>
02310902	TEVA-ATORVASTATIN	TEV	<b>FGNQSW</b>
02313715	TARO-ATORVASTATIN	SUN	<b>FGNQSW</b>
02324954	SANDOZ-ATORVASTATIN	SDZ	<b>FGNQSW</b>
02348713	ATORVASTATIN	SNS	<b>FGNQSW</b>
02391066	JAMP-ATORVASTATIN	JPC	<b>FGNQSW</b>



02392941	MYLAN-ATORVASTATIN	MYL	<b>FGNQSW</b>
02407264	AURO-ATORVASTATIN	ARO	<b>FGNQSW</b>
02411369	ATORVASTATIN	SIV	<b>FGNQSW</b>
02417944	REDDY-ATORVASTATIN	RCH	<b>FGNQSW</b>
02454025	MAR-ATORVASTATIN	MAR	<b>FGNQSW</b>
02457768	ACH-ATORVASTATIN	ACH	<b>FGNQSW</b>
02471175	M-ATORVASTATIN	MRA	<b>FGNQSW</b>
02475030	ATORVASTATIN	RIV	<b>FGNQSW</b>
02476525	NRA-ATORVASTATIN	NRA	<b>FGNQSW</b>
02477157	PMS-ATORVASTATIN	PMS	<b>FGNQSW</b>
02478153	AG-ATORVASTATIN	ANG	<b>FGNQSW</b>
02479516	MINT-ATORVASTATIN	MNT	<b>FGNQSW</b>
02504200	JAMP-ATORVASTATIN	JPC	<b>FGNQSW</b>
02507242	PMSC-ATORVASTATIN	PMS	<b>FGNQSW</b>
02521563	PRZ-ATORVASTATIN	PRZ	<b>FGNQSW</b>

40MG TABLET

02230714	LIPITOR	UJC	<b>FNQSW</b>
02295296	APO-ATORVASTATIN	APX	<b>FGNQSW</b>
02310910	TEVA-ATORVASTATIN	TEV	<b>FGNQSW</b>
02313723	TARO-ATORVASTATIN	SUN	<b>FGNQSW</b>
02324962	SANDOZ-ATORVASTATIN	SDZ	<b>FGNQSW</b>
02348721	ATORVASTATIN	SNS	<b>FGNQSW</b>
02391074	JAMP-ATORVASTATIN	JPC	<b>FGNQSW</b>
02392968	MYLAN-ATORVASTATIN	MYL	<b>FGNQSW</b>
02407272	AURO-ATORVASTATIN	ARO	<b>FGNQSW</b>
02411377	ATORVASTATIN	SIV	<b>FGNQSW</b>
02417952	REDDY-ATORVASTATIN	RCH	<b>FGNQSW</b>
02454033	MAR-ATORVASTATIN	MAR	<b>FGNQSW</b>
02457776	ACH-ATORVASTATIN	ACH	<b>FGNQSW</b>
02471183	M-ATORVASTATIN	MRA	<b>FGNQSW</b>
02475049	ATORVASTATIN	RIV	<b>FGNQSW</b>
02476533	NRA-ATORVASTATIN	NRA	<b>FGNQSW</b>
02477165	PMS-ATORVASTATIN	PMS	<b>FGNQSW</b>
02478161	AG-ATORVASTATIN	ANG	<b>FGNQSW</b>
02479524	MINT-ATORVASTATIN	MNT	<b>FGNQSW</b>
02504219	JAMP-ATORVASTATIN	JPC	<b>FGNQSW</b>
02507250	PMSC-ATORVASTATIN	PMS	<b>FGNQSW</b>
02521571	PRZ-ATORVASTATIN	PRZ	<b>FGNQSW</b>

80MG TABLET

02243097	LIPITOR	UJC	<b>FNQSW</b>
02295318	APO-ATORVASTATIN	APX	<b>FGNQSW</b>
02310929	TEVA-ATORVASTATIN	TEV	<b>FGNQSW</b>
02313758	TARO-ATORVASTATIN	SUN	<b>FGNQSW</b>
02324970	SANDOZ-ATORVASTATIN	SDZ	<b>FGNQSW</b>

02348748	ATORVASTATIN	SNS	FGNQSW
02391082	JAMP-ATORVASTATIN	JPC	FGNQSW
02392976	MYLAN-ATORVASTATIN	MYL	FGNQSW
02407280	AURO-ATORVASTATIN	ARO	FGNQSW
02411385	ATORVASTATIN	SIV	FGNQSW
02417960	REDDY-ATORVASTATIN	RCH	FGNQSW
02454041	MAR-ATORVASTATIN	MAR	FGNQSW
02457784	ACH-ATORVASTATIN	ACH	FGNQSW
02471191	M-ATORVASTATIN	MRA	FGNQSW
02475057	ATORVASTATIN	RIV	FGNQSW
02476541	NRA-ATORVASTATIN	NRA	FGNQSW
02479532	MINT-ATORVASTATIN	MNT	FGNQSW
02478188	AG-ATORVASTATIN	ANG	FGNQSW
02504235	JAMP-ATORVASTATIN	JPC	FGNQSW
02507269	PMSC-ATORVASTATIN	PMS	FGNQSW
02521598	PRZ-ATORVASTATIN	PRZ	FGNQSW

**⑤CHOLESTYRAMINE**

REGULAR - 4G/POUCH X 30 POUCHES - 120G/PK ORAL POWDER (POUCHES)

02210320	OLESTYR	PMS	FGNQSW
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\*price per gram for cholestyramine powder pouches

LIGHT - 4G/POUCH X 30 POUCHES- 120G/PK

02478595	JAMP-CHOLESTYRAMINE	JPC	FGNQSW
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**⑤COLESEVELAM**

625MG TABLET

02373955	LODALIS	VAL	FNQSW
02494051	APO-COLESEVELAM	APX	FGNQSW

3.75G PACKET

02432463	LODALIS	VAL	FNQSW
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**⑤EVOLOCUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

140MG/ML PEN INJECTOR

02446057	REPATHA (SA)	AMG	FNQSW
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**⑤EZETIMIBE**

10MG TABLET

02247521	EZETROL	MSD	FNQSW
02354101	TEVA-EZETIMIBE	TEV	FGNQSW
02416409	PMS-EZETIMIBE	PMS	FGNQSW
02416778	SANDOZ-EZETIMIBE	SDZ	FGNQSW
02419548	RAN-EZETIMIBE	RAN	FGNQSW
02422662	MAR-EZETIMIBE	MAR	FGNQSW

02423235	JAMP-EZETIMIBE	JPC	FGNQSW
02423243	MINT-EZETIMIBE	MNT	FGNQSW
02425610	ACH-EZETIMIBE	ACH	FGNQSW
02427826	APO-EZETIMIBE	APX	FGNQSW
02429659	EZETIMIBE	SIV	FGNQSW
02431300	EZETIMIBE	SNS	FGNQSW
02460750	GLN-EZETIMIBE	GLM	FGNQSW
02467437	M-EZETIMIBE	MRA	FGNQSW
02469286	AURO-EZETIMIBE	ARO	FGNQSW
02475898	AG-EZETIMIBE	AGP	FGNQSW
02481669	NRA-EZETIMIBE	NRA	FGNQSW

**⑤FENOFIBRATE**

67MG CAPSULE

02243180	AA-FENO-MICRO	AAA	FGNQSW
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100MG TABLET

02246859	AA-FENO-SUPER	AAA	FGNQSW
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160MG TABLET

02241602	LIPIDIL SUPRA	BGP	FNQSW
02246860	AA-FENO-SUPER	AAA	FGNQSW

200MG CAPSULE

02239864	AA-FENO-MICRO	AAA	FGNQSW
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**⑤FLUVASTATIN SODIUM**

20MG CAPSULE

02299224	TEVA-FLUVASTATIN	TEV	FGNQSW
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40MG CAPSULE

02299232	TEVA-FLUVASTATIN	TEV	FGNQSW
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**⑤GEMFIBROZIL**

600MG TABLET

02142074	TEVA-GEMFIBROZIL	TEV	FGNQSW
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**⑤LOVASTATIN**

20MG TABLET

02220172	LOVASTATIN	AAA	FGNQSW
02248572	ACT-LOVASTATIN	TEV	FGNQSW

40MG TABLET

02220180	LOVASTATIN	AAA	FGNQSW
02248573	ACT-LOVASTATIN	TEV	FGNQSW

## ⑤ PRAVASTATIN

### 10MG TABLET

02243506	APO-PRAVASTATIN	APX	FGNQSW
02247008	TEVA-PRAVASTATIN	TEV	FGNQSW
02247655	PMS-PRAVASTATIN	PMS	FGNQSW
02284421	RAN-PRAVASTATIN	RAN	FGNQSW
02317451	MINT-PRAVASTATIN	MNT	FGNQSW
02330954	JAMP PRAVASTATIN	JPC	FGNQSW
02356546	PRAVASTATIN	SNS	FGNQSW
02389703	PRAVASTATIN	SVI	FGNQSW
02432048	MAR-PRAVASTATIN	MAR	FGNQSW
02440644	ACH-PRAVASTATIN	ACH	FGNQSW
02458977	AURO-PRAVASTATIN	ARO	FGNQSW
02468700	SANDOZ-PRAVASTATIN	SDZ	FGNQSW
02476142	AG-PRAVASTATIN	ANG	FGNQSW
02476274	M-PRAVASTATIN	MRA	FGNQSW

### 20MG TABLET

02243507	APO-PRAVASTATIN	APX	FGNQSW
02247009	TEVA-PRAVASTATIN	TEV	FGNQSW
02247656	PMS-PRAVASTATIN	PMS	FGNQSW
02284448	RAN-PRAVASTATIN	RAN	FGNQSW
02317478	MINT-PRAVASTATIN	MNT	FGNQSW
02330962	JAMP PRAVASTATIN	JPC	FGNQSW
02356554	PRAVASTATIN	SNS	FGNQSW
02389738	PRAVASTATIN	SIV	FGNQSW
02432056	MAR-PRAVASTATIN	MAR	FGNQSW
02440652	ACH-PRAVASTATIN	ACH	FGNQSW
02458985	AURO-PRAVASTATIN	ARO	FGNQSW
02468719	SANDOZ-PRAVASTATIN	SDZ	FGNQSW
02476150	AG-PRAVASTATIN	ANG	FGNQSW
02476282	M-PRAVASTATIN	MRA	FGNQSW

### 40MG TABLET

02243508	APO-PRAVASTATIN	APX	FGNQSW
02247010	TEVA-PRAVASTATIN	TEV	FGNQSW
02247657	PMS-PRAVASTATIN	PMS	FGNQSW
02284456	RAN-PRAVASTATIN	RAN	FGNQSW
02317486	MINT-PRAVASTATIN	MNT	FGNQSW
02330970	JAMP PRAVASTATIN	JPC	FGNQSW
02356562	PRAVASTATIN	SNS	FGNQSW
02389746	PRAVASTATIN	SIV	FGNQSW
02432064	MAR-PRAVASTATIN	MAR	FGNQSW
02458993	AURO-PRAVASTATIN	ARO	FGNQSW
02468727	SANDOZ-PRAVASTATIN	SDZ	FGNQSW
02476169	AG-PRAVASTATIN	ANG	FGNQSW

02476290 M-PRAVASTATIN MRA FGNQSW

**⑤ROSUVASTATIN**

**5MG TABLET**

02265540	CRESTOR	AZE	FNQSW
02337975	APO-ROSUVASTATIN	APX	FGNQSW
02338726	SANDOZ-ROSUVASTATIN	SDZ	FGNQSW
02354608	TEVA-ROSUVASTATIN	TEV	FGNQSW
02378523	PMS-ROSUVASTATIN	PMS	FGNQSW
02382644	TARO-ROSUVASTATIN	SUN	FGNQSW
02391252	JAMP-ROSUVASTATIN	JPC	FGNQSW
02405628	ROSUVASTATIN	SNS	FGNQSW
02397781	MINT-ROSUVASTATIN	MNT	FGNQSW
02399164	MED-ROSUVASTATIN	GMP	FGNQSW
02411628	ROSUVASTATIN-5	SIV	FGNQSW
02413051	MAR-ROSUVASTATIN	MAR	FGNQSW
02438917	ACH-ROSUVASTATIN	ACH	FGNQSW
02442574	AURO-ROSUVASTATIN	ARO	FGNQSW
02477483	NRA-ROSUVASTATIN	NRA	FGNQSW
02496534	M-ROSUVASTATIN	MRA	FGNQSW
02498332	JAMP-ROSUVASTATIN CALCIUM	JPC	FGNQSW
02505576	PRZ-ROSUVASTATIN	PRZ	FGNQSW

**10MG TABLET**

02247162	CRESTOR	AZE	FNQSW
02337983	APO-ROSUVASTATIN	APX	FGNQSW
02338734	SANDOZ-ROSUVASTATIN	SDZ	FGNQSW
02354616	TEVA-ROSUVASTATIN	TEV	FGNQSW
02378531	PMS-ROSUVASTATIN	PMS	FGNQSW
02382652	TARO-ROSUVASTATIN	SUN	FGNQSW
02391260	JAMP-ROSUVASTATIN	JPC	FGNQSW
02405636	ROSUVASTATIN	SNS	FGNQSW
02397803	MINT-ROSUVASTATIN	MNT	FGNQSW
02399172	MED-ROSUVASTATIN	GMP	FGNQSW
02411636	ROSUVASTATIN-10	SIV	FGNQSW
02413078	MAR-ROSUVASTATIN	MAR	FGNQSW
02438925	ACH-ROSUVASTATIN	ACH	FGNQSW
02442582	AURO-ROSUVASTATIN	ARO	FGNQSW
02477491	NRA-ROSUVASTATIN	NRA	FGNQSW
02496542	M-ROSUVASTATIN	MRA	FGNQSW
02498340	JAMP-ROSUVASTATIN CALCIUM	JPC	FGNQSW
02505584	PRZ-ROSUVASTATIN	PRZ	FGNQSW

**20MG TABLET**

02247163	CRESTOR	AZE	FNQSW
02337991	APO-ROSUVASTATIN	APX	FGNQSW

02338742	SANDOZ-ROSUVASTATIN	SDZ	FGNQSW
02354624	TEVA-ROSUVASTATIN	TEV	FGNQSW
02378558	PMS-ROSUVASTATIN	PMS	FGNQSW
02382660	TARO-ROSUVASTATIN	SUN	FGNQSW
02391279	JAMP-ROSUVASTATIN	JPC	FGNQSW
02405644	ROSUVASTATIN	SNS	FGNQSW
02399180	MED-ROSUVASTATIN	GMP	FGNQSW
02411644	ROSUVASTATIN-20	SIV	FGNQSW
02413086	MAR-ROSUVASTATIN	MAR	FGNQSW
02438933	ACH-ROSUVASTATIN	ACH	FGNQSW
02442590	AURO-ROSUVASTATIN	ARO	FGNQSW
02477505	NRA-ROSUVASTATIN	NRA	FGNQSW
02496550	M-ROSUVASTATIN	MRA	FGNQSW
02498359	JAMP-ROSUVASTATIN	JPC	FGNQSW
02505592	PRZ-ROSUVASTATIN	PRZ	FGNQSW

40MG TABLET

02247164	CRESTOR	AZE	FNQSW
02338009	APO-ROSUVASTATIN	APX	FGNQSW
02338750	SANDOZ-ROSUVASTATIN	SDZ	FGNQSW
02354632	TEVA-ROSUVASTATIN	TEV	FGNQSW
02378566	PMS-ROSUVASTATIN	PMS	FGNQSW
02382679	TARO-ROSUVASTATIN	SUN	FGNQSW
02391287	JAMP-ROSUVASTATIN	JPC	FGNQSW
02405652	ROSUVASTATIN	SNS	FGNQSW
02399199	MED-ROSUVASTATIN	GMP	FGNQSW
02411652	ROSUVASTATIN-40	SIV	FGNQSW
02413108	MAR-ROSUVASTATIN	MAR	FGNQSW
02438941	ACH-ROSUVASTATIN	ACH	FGNQSW
02442604	AURO-ROSUVASTATIN	ARO	FGNQSW
02477513	NRA-ROSUVASTATIN	NRA	FGNQSW
02496569	M-ROSUVASTATIN	MRA	FGNQSW
02498367	JAMP-ROSUVASTATIN	JPC	FGNQSW
02505606	PRZ-ROSUVASTATIN	PRZ	FGNQSW

⑤SIMVASTATIN

5MG TABLET

02247011	APO-SIMVASTATIN	APX	FGNQSW
02250144	TEVA-SIMVASTATIN	TEV	FGNQSW
02284723	SIMVASTATIN	SNS	FGNQSW
02329131	RAN-SIMVASTATIN	RAN	FGNQSW
02372932	MINT-SIMVASTATIN	MNT	FGNQSW
02375036	MAR-SIMVASTATIN	MAR	FGNQSW
02375591	JAMP-SIMVASTATIN	JPC	FGNQSW
02386291	SIMVASTATIN	SIV	FGNQSW
02405148	AURO-SIMVASTATIN	ARO	FGNQSW

02469979	PHARMA-SIMVASTATIN	PMS	<b>FGNQSW</b>
02480050	AG-SIMVASTATIN	ANG	<b>FGNQSW</b>

10MG TABLET

00884332	ZOCOR	MSD	<b>FNQSW</b>
02247012	APO-SIMVASTATIN	APX	<b>FGNQSW</b>
02250152	TEVA-SIMVASTATIN	TEV	<b>FGNQSW</b>
02284731	SIMVASTATIN	SNS	<b>FGNQSW</b>
02329158	RAN-SIMVASTATIN	RAN	<b>FGNQSW</b>
02372940	MINT-SIMVASTATIN	MNT	<b>FGNQSW</b>
02375044	MAR-SIMVISTATIN	MAR	<b>FGNQSW</b>
02375605	JAMP-SIMVASTATIN	JPC	<b>FGNQSW</b>
02386305	SIMVASTATIN	SIV	<b>FGNQSW</b>
02405156	AURO-SIMVASTATIN	ARO	<b>FGNQSW</b>
02469987	PHARMA-SIMVASTATIN	PMS	<b>FGNQSW</b>
02480069	AG-SIMVASTATIN	ANG	<b>FGNQSW</b>

20MG TABLET

00884340	ZOCOR	MSD	<b>FNQSW</b>
02247013	APO-SIMVASTATIN	APX	<b>FGNQSW</b>
02250160	TEVA-SIMVASTATIN	TEV	<b>FGNQSW</b>
02284758	SIMVASTATIN	SNS	<b>FGNQSW</b>
02329166	RAN-SIMVASTATIN	RAN	<b>FGNQSW</b>
02372959	MINT-SIMVASTATIN	MNT	<b>FGNQSW</b>
02375052	MAR-SIMVISTATIN	MAR	<b>FGNQSW</b>
02375613	JAMP-SIMVASTATIN	JPC	<b>FGNQSW</b>
02386313	SIMVASTATIN	SIV	<b>FGNQSW</b>
02405164	AURO-SIMVASTATIN	ARO	<b>FGNQSW</b>
02469995	PHARMA-SIMVASTATIN	PMS	<b>FGNQSW</b>
02480077	AG-SIMVASTATIN	ANG	<b>FGNQSW</b>

40MG TABLET

00884359	ZOCOR	MSD	<b>FNQSW</b>
02247014	APO-SIMVASTATIN	APX	<b>FGNQSW</b>
02250179	TEVA-SIMVASTATIN	TEV	<b>FGNQSW</b>
02284766	SIMVASTATIN	SNS	<b>FGNQSW</b>
02329174	RAN-SIMVASTATIN	RAN	<b>FGNQSW</b>
02372967	MINT-SIMVASTATIN	MNT	<b>FGNQSW</b>
02375060	MAR-SIMVISTATIN	MAR	<b>FGNQSW</b>
02375621	JAMP-SIMVASTATIN	JPC	<b>FGNQSW</b>
02386321	SIMVASTATIN	SIV	<b>FGNQSW</b>
02405172	AURO-SIMVASTATIN	ARO	<b>FGNQSW</b>
02470004	PHARMA-SIMVASTATIN	PMS	<b>FGNQSW</b>
02480085	AG-SIMVASTATIN	ANG	<b>FGNQSW</b>

80MG TABLET

02247015	APO-SIMVASTATIN	APX	FGNQSW
02250187	TEVA-SIMVASTATIN	TEV	FGNQSW
02284774	SIMVASTATIN	SNS	FGNQSW
02329182	RAN-SIMVASTATIN	RAN	FGNQSW
02372975	MINT-SIMVASTATIN	MNT	FGNQSW
02375079	MAR-SIMVISTATIN	MAR	FGNQSW
02375648	JAMP-SIMVASTATIN	JPC	FGNQSW
02386348	SIMVASTATIN	SIV	FGNQSW
02405180	AURO-SIMVASTATIN	ARO	FGNQSW
02470012	PHARMA-SIMVASTATIN	PMS	FGNQSW
02480093	AG-SIMVASTATIN	ANG	FGNQSW

## **24:08.00 HYPOTENSIVE DRUGS**

### **⑤EPLERENONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

#### **25MG TABLET**

02471442	MINT-EPLERENONE (SA)	MNT	FGNQSW
02543389	JAMP-EPLERENONE (SA)	JPC	FGNQSW

#### **50MG TABLET**

02471450	MINT-EPLERENONE (SA)	MNT	FGNQSW
02543397	JAMP-EPLERENONE (SA)	JPC	FGNQSW

### **⑤HYDRALAZINE HCL**

#### **10MG TABLET**

00441619	APO-HYDRALAZINE	APX	FGNQSW
02457865	JAMP-HYDRALAZINE	JPC	FGNQSW
02468778	MINT-HYDRALAZINE	MNT	FGNQSW
02539802	HYDRALAZINE	SNS	FGNQSW

#### **25MG TABLET**

00441627	APO-HYDRALAZINE	APX	FGNQSW
02457873	JAMP-HYDRALAZINE	JPC	FGNQSW
02468786	MINT-HYDRALAZINE	MNT	FGNQSW
02539810	HYDRALAZINE	SNS	FGNQSW

#### **50MG TABLET**

00441635	APO-HYDRALAZINE	APX	FGNQSW
02457881	JAMP-HYDRALAZINE	JPC	FGNQSW
02468794	MINT-HYDRALAZINE	MNT	FGNQSW
02539829	HYDRALAZINE	SNS	FGNQSW



**⑤PERINDOPRIL****2MG TABLET**

02123274	COVERSYL	SEV	<b>FNQSW</b>
02289261	APO-PERINDOPRIL	APX	<b>FGNQSW</b>
02459817	AURO-PERINDOPRIL	ARO	<b>FGNQSW</b>
02464985	TEVA-PERINDOPRIL	TEV	<b>FGNQSW</b>
02470225	SANDOZ-PERINDOPRIL ERBUMINE	SDZ	<b>FGNQSW</b>
02470675	PMS-PERINDOPRIL	PMS	<b>FGNQSW</b>
02474824	MAR-PERINDOPRIL	MAR	<b>FGNQSW</b>
02476762	MINT-PERINDOPRIL	MNT	<b>FGNQSW</b>
02477009	JAMP-PERINDOPRIL	JPC	<b>FGNQSW</b>
02479877	PERINDOPRIL ERBUMINE	SIV	<b>FGNQSW</b>
02481634	PERINDOPRIL ERBUMINE	SNS	<b>FGNQSW</b>
02481677	AG-PERINDOPRIL	ANG	<b>FGNQSW</b>
02482924	M-PERINDOPRIL ERBUMINE	MRA	<b>FGNQSW</b>
02489015	NRA-PERINDOPRIL	NRA	<b>FGNQSW</b>
02527200	JAMP-PERINDOPRIL	JPC	<b>FGNQSW</b>

**4MG TABLET**

02123282	COVERSYL	SEV	<b>FNQSW</b>
02289288	APO-PERINDOPRIL	APX	<b>FGNQSW</b>
02459825	AURO-PERINDOPRIL	ARO	<b>FGNQSW</b>
02464993	TEVA-PERINDOPRIL	TEV	<b>FGNQSW</b>
02470233	SANDOZ-PERINDOPRIL ERBUMINE	SDZ	<b>FGNQSW</b>
02470683	PMS-PERINDOPRIL	PMS	<b>FGNQSW</b>
02474832	MAR-PERINDOPRIL	MAR	<b>FGNQSW</b>
02477017	JAMP-PERINDOPRIL	JPC	<b>FGNQSW</b>
02476770	MINT-PERINDOPRIL	MNT	<b>FGNQSW</b>
02479885	PERINDOPRIL ERBUMINE	SIV	<b>FGNQSW</b>
02481642	PERINDOPRIL ERBUMINE	SNS	<b>FGNQSW</b>
02481685	AG-PERINDOPRIL	ANG	<b>FGNQSW</b>
02482932	M-PERINDOPRIL ERBUMINE	MRA	<b>FGNQSW</b>
02489023	NRA-PERINDOPRIL	NRA	<b>FGNQSW</b>
02527219	JAMP-PERINDOPRIL	JPC	<b>FGNQSW</b>

**8MG TABLET**

02246624	COVERSYL	SEV	<b>FNQSW</b>
02289296	APO-PERINDOPRIL	APX	<b>FGNQSW</b>
02459833	AURO-PERINDOPRIL	ARO	<b>FGNQSW</b>
02465000	TEVA-PERINDOPRIL	TEV	<b>FGNQSW</b>
02470241	SANDOZ-PERINDOPRIL ERBUMINE	SDZ	<b>FGNQSW</b>
02470691	PMS-PERINDOPRIL	PMS	<b>FGNQSW</b>
02474840	MAR-PERINDOPRIL	MAR	<b>FGNQSW</b>
02477025	JAMP-PERINDOPRIL	JPC	<b>FGNQSW</b>
02476789	MINT-PERINDOPRIL	MNT	<b>FGNQSW</b>
02479893	PERINDOPRIL ERBUMINE	SIV	<b>FGNQSW</b>

02481650	PERINDOPRIL ERBUMINE	SNS	FGNQSW
02481693	AG-PERINDOPRIL	ANG	FGNQSW
02482940	M-PERINDOPRIL ERBUMINE	MRA	FGNQSW
02489031	NRA-PERINDOPRIL	NRA	FGNQSW
02527227	JAMP-PERINDOPRIL	JPC	FGNQSW

**⑤ PERINDOPRIL & INDAPAMIDE**

4MG & 1.25MGMG TABLET

02246569	COVERSYL PLUS	SEV	FNQSW
02297574	APO-PERINDOPRIL/INDAPAMIDE	APX	FGNQSW
02464020	TEVA-PERINDOPRIL/INDAPAMIDE	TEV	FGNQSW
02470438	PERINDOPRIL ERBUMIN-INDAPAMIDE	SDZ	FGNQSW
02479834	PERINDOPRIL ERBUMIN-INDAPAMIDE	SIV	FGNQSW
02519720	PERINDOPRIL-INDAPAMIDE	SNS	FGNQSW
02538008	PMS-PERINDOPRIL INDAPAMIDE	PMS	FGNQSW

**⑤ PERINDOPRIL ERBUMIN/INDAPAMIDE**

8MG & 2.5MG TABLET

02321653	COVERSYL PLUS HD	SEV	FNQSW
02453061	APO-PERINDOPRIL/INDAPAMIDE	APX	FGNQSW
02464039	TEVA-PERINDOPRIL/INDAPAMIDE	TEV	FGNQSW
02470446	PERINDOPRIL ERBUMIN-INDAPAMIDE HD	SDZ	FGNQSW
02479842	PERINDOPRIL ERBUMIN-INDAPAMIDE HD	SIV	FGNQSW
02519739	PERINDOPRIL-INDAPAMIDE	SNS	FGNQSW
02537982	PMS-PERINDOPRIL INDAPAMIDE	PMS	FGNQSW

**PINDOLOL & HYDROCHLOROTHIAZIDE**

10MG & 50MG TABLET

00568635	VISKAZIDE	XPI	FNQSW
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**24:08.16 CENTRAL ALPHA AGONISTS**

**⑤ CLONIDINE HCL**

0.025MG TABLET

02304163	TEVA-CLONIDINE	TEV	FGNQSW
02516217	SANDOZ-CLONIDINE	SDZ	FGNQSW
02524198	MAR-CLONIDINE	MAR	FGNQSW
02528207	JAMP-CLONIDINE	JPC	FGNQSW
02534738	MINT-CLONIDINE	MNT	FGNQSW
02540061	CLONIDINE	SIV	FGNQSW

0.1MG TABLET

02046121	TEVA-CLONIDINE	TEV	FGNQSW
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02462192	MINT-CLONIDINE	MNT	FGNQSW
02515784	SANDOZ-CLONIDINE	SDZ	FGNQSW
02538490	CLONIDINE	SIV	FGNQSW

0.2MG TABLET			
02046148	TEVA-CLONIDINE	TEV	FGNQSW
02462206	MINT-CLONIDINE	MNT	FGNQSW
02515792	SANDOZ-CLONIDINE	SDZ	FGNQSW
02538504	CLONIDINE	SIV	FGNQSW

**⑤ METHYLDOPA**

125MG TABLET			
00360252	METHYLDOPA	AAA	FGNQSW

250MG TABLET			
00360260	METHYLDOPA	AAA	FGNQSW

500MG TABLET			
00426830	METHYLDOPA	AAA	FGNQSW

**24:12.00 MISCELLANEOUS VASODILATING AGENTS**

**AMBRISANTAN**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG TABLET

02307065	VOLIBRIS (SA)	GSK	NMQW
02475375	APO-AMBRISANTAN (SA)	APX	NMQW
02521938	JAMP-AMBRISANTAN (SA)	JPC	NMQW
02526875	SANDOZ-AMBRISANTAN (SA)	SDZ	NMQW

10MG TABLET

02307073	VOLIBRIS (SA)	GSK	NMQW
02475383	APO-AMBRISANTAN (SA)	APX	NMQW
02521946	JAMP-AMBRISANTAN (SA)	JPC	NMQW
02526883	SANDOZ-AMBRISANTAN (SA)	SDZ	NMQW

**BETAHISTINE HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA (EXCEPT NURSING HOME PROGRAM)

16MG TABLET

02243878	SERC (SA)	BGP	FNQSW
02280191	TEVA-BETAHISTINE (SA)	TEV	FGNQSW
02330210	PMS-BETAHISTINE (SA)	PMS	FGNQSW
02449153	AURO-BETAHISTINE (SA)	ARO	FGNQSW

02466449	BETAHISTINE (SA)	SNS	<b>FGNQSW</b>
02519690	M-BETAHISTINE (SA)	MRA	<b>FGNQSW</b>
02538148	MINT-BETAHISTINE (SA)	MNT	<b>FGNQSW</b>

**24MG TABLET**

02247998	SERC (SA)	BGP	<b>FNQSW</b>
02280205	TEVA-BETAHISTINE (SA)	TEV	<b>FGNQSW</b>
02330237	PMS-BETAHISTINE (SA)	PMS	<b>FGNQSW</b>
02449161	AURO-BETAHISTINE (SA)	ARO	<b>FGNQSW</b>
02466457	BETAHISTINE (SA)	SNS	<b>FGNQSW</b>
02519704	M-BETAHISTINE (SA)	MRA	<b>FGNQSW</b>
02538156	MINT-BETAHISTINE (SA)	MNT	<b>FGNQSW</b>

**DIPYRIDAMOLE**

**25MG TABLET**

00895644	APO-DIPYRIDAMOLE-FC	APX	<b>FGNQSW</b>
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**50MG TABLET**

00895652	APO-DIPYRIDAMOLE-FC	APX	<b>FGNQSW</b>
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**75MG TABLET**

00895660	APO-DIPYRIDAMOLE-FC	APX	<b>FGNQSW</b>
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**EPOPROSTENOL SODIUM (GLYCINE)**

[SEE APPENDIX A](#) FOR SA CRITERIA

**0.5 MG INJECTION**

02230845	FLOLAN (SA)	GSK	<b>NMQW</b>
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**1.5 MG INJECTION**

02230848	FLOLAN (SA)	GSK	<b>NMQW</b>
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**EPOPROSTENOL SODIUM (ARGININE)**

[SEE APPENDIX A](#) FOR SA CRITERIA

**0.5MG INJECTION**

02397447	CARIPUL (SA)	JAN	<b>NMQW</b>
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**1.5MG INJECTION**

02397455	CARIPUL (SA)	JAN	<b>NMQW</b>
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**⑤ ISOSORBIDE DINITRATE**

**10MG TABLET**

00441686	ISDN	AAA	<b>FGNQSW</b>
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**30MG TABLET**

00441694	ISDN	AAA	<b>FGNQSW</b>
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### ⑤ ISOSORBIDE MONONITRATE

60MG TABLET

02126559	IMDUR	AST	<b>FNQSW</b>
02272830	APO-ISMM	APX	<b>FGNQSW</b>
02301288	PMS-ISMN	PMS	<b>FGNQSW</b>

### MACITENTAN

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET

02415690	OPSUMIT (SA)	JAN	<b>NMQW</b>
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### NIMODIPINE

30MG TABLET

02325926	NIMOTOP	BAY	<b>FNQSW</b>
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### NITROGLYCERIN

#### NOTES:

1. To prevent development of tolerance, patches should be removed after 12-14 hours to provide daily NITRATE-FREE periods of 10-12 hours. The NITRATE-FREE period should be timed to coincide with the period in which angina is least likely to occur (USUALLY AT NIGHT).

### ⑤ NITROGLYCERIN TRANSDERMAL

Eligible for a 90 day supply

0.2MG/HR TRANSDERMAL PATCH

01911910	NITRO-DUR 0.2	RCH	<b>FNQSW</b>
02407442	MYLAN-NITRO PATCH	MYL	<b>FGNQSW</b>

0.2MG/HR TRANSDERMAL PATCH

02230732	TRINIPATCH 0.2	PAL	<b>FQSW</b>
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0.4 MG/HR TRANSDERMAL PATCH

01911902	NITRO-DUR 0.4	RCH	<b>FNQSW</b>
02407450	MYLAN-NITRO PATCH	MYL	<b>FGNQSW</b>

0.4 MG/HR TRANSDERMAL PATCH

02230733	TRINIPATCH 0.4	PAL	<b>FQSW</b>
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0.6 MG/HR TRANSDERMAL PATCH

01911929	NITRO-DUR 0.6	RCH	<b>FNQSW</b>
02407469	MYLAN-NITRO PATCH	MYL	<b>FGNQSW</b>

0.6 MG/HR TRANSDERMAL PATCH

02046156	TRANSDERM - NITRO 0.6	NVR	<b>FQSW</b>
02230734	TRINIPATCH 0.6	PAL	<b>FQSW</b>

0.8MG/HR TRANSDERMAL PATCH			
02011271	NITRO-DUR 0.8	RCH	<b>FNQSW</b>
02407477	MYLAN-NITRO PATCH	MYL	<b>FGNQSW</b>

**NITROGLYCERIN**

0.3MG SUBLINGUAL TABLET			
00037613	NITROSTAT	UJC	<b>NQW</b>

0.6MG SUBLINGUAL TABLET			
00037621	NITROSTAT	UJC	<b>NQW</b>

0.4MG/DOSE METERED DOSE LINGUAL SPRAY			
02231441	NITROLINGUAL PUMPSPRAY	AVN	<b>NQW</b>
02238998	RHO-NITRO PUMPSPRAY	SDZ	<b>NQW</b>
02243588	MYLAN-NITRO SL SPRAY	MYL	<b>NQW</b>
02393433	APO-NITROGLYCERIN	APX	<b>NQW</b>

**RIOCIGUAT**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.5MG TABLET			
02412764	ADEMPAS (SA)	BAY	<b>NMQW</b>
02533545	SANDOZ-RIOCIGUAT (SA)	SDZ	<b>NMQW</b>

1MG TABLET			
02412772	ADEMPAS (SA)	BAY	<b>NMQW</b>
02533561	SANDOZ-RIOCIGUAT (SA)	SDZ	<b>NMQW</b>

1.5MG TABLET			
02412799	ADEMPAS (SA)	BAY	<b>NMQW</b>
02533588	SANDOZ-RIOCIGUAT (SA)	SDZ	<b>NMQW</b>

2MG TABLET			
02412802	ADEMPAS (SA)	BAY	<b>NMQW</b>
02533596	SANDOZ-RIOCIGUAT (SA)	SDZ	<b>NMQW</b>

2.5MG TABLET			
02412810	ADEMPAS (SA)	BAY	<b>NMQW</b>
02533618	SANDOZ-RIOCIGUAT (SA)	SDZ	<b>NMQW</b>

**SELEXIPAG**

[SEE APPENDIX A](#) FOR SA CRITERIA

200MCG TABLET			
02451158	UPTRAVI (SA)	JAN	<b>NMQW</b>

400MCG TABLET			
02451166	UPTRAVI (SA)	JAN	<b>NMQW</b>

600MCG TABLET 02451174	UPTRAVI (SA)	JAN	<b>NMQW</b>
800MCG TABLET 02451182	UPTRAVI (SA)	JAN	<b>NMQW</b>
1000MCG TABLET 02451190	UPTRAVI (SA)	JAN	<b>NMQW</b>
1200MCG TABLET 02451204	UPTRAVI (SA)	JAN	<b>NMQW</b>
1400MCG TABLET 02451212	UPTRAVI (SA)	JAN	<b>NMQW</b>
1600MCG TABLET 02451220	UPTRAVI (SA)	JAN	<b>NMQW</b>

### **SILDENAFIL CITRATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG TABLET			
02279401	REVATIO (SA)	UJC	<b>MNSQW</b>
02319500	TEVA-SILDENAFIL R (SA)	TEV	<b>GMNSQW</b>
02412179	PMS-SILDENAFIL-R (SA)	PMS	<b>GMNSQW</b>
02469669	JAMP-SILDENAFIL R (SA)	JPC	<b>GMNSQW</b>

## **24:20.00 ALPHA ADRENERGIC BLOCKING AGENTS**

### **⑤ DOXAZOSIN**

1MG TABLET			
02240588	APO-DOXAZOSIN	APX	<b>FGNQSW</b>
02242728	TEVA-DOXAZOSIN	TEV	<b>FGNQSW</b>
02489937	JAMP-DOXAZOSIN	JPC	<b>FGNQSW</b>
2MG TABLET			
02240589	APO-DOXAZOSIN	APX	<b>FGNQSW</b>
02242729	TEVA-DOXAZOSIN	TEV	<b>FGNQSW</b>
02489945	JAMP-DOXAZOSIN	JPC	<b>FGNQSW</b>
4MG TABLET			
02240590	APO-DOXAZOSIN	APX	<b>FGNQSW</b>
02242730	TEVA-DOXAZOSIN	TEV	<b>FGNQSW</b>

02489953 JAMP-DOXAZOSIN JPC FGNQSW

**⑤ PRAZOSIN HCL**

1MG TABLET  
01934198 TEVA-PRAZOSIN TEV FGNQSW

2MG TABLET  
01934201 TEVA-PRAZOSIN TEV FGNQSW

5MG TABLET  
01934228 TEVA-PRAZOSIN TEV FGNQSW

**⑤ TERAZOSIN HCL**

1MG TABLET  
02234502 APO-TERAZOSIN APX FGNQSW  
02243518 PMS-TERAZOSIN PMS FGNQSW

2MG TABLET  
02234503 APO-TERAZOSIN APX FGNQSW  
02243519 PMS-TERAZOSIN PMS FGNQSW

5MG TABLET  
02230807 TEVA-TERAZOSIN TEV FGNQSW  
02234504 APO-TERAZOSIN APX FGNQSW  
02243520 PMS-TERAZOSIN PMS FGNQSW

10MG TABLET  
02234505 APO-TERAZOSIN APX FGNQSW  
02243521 PMS-TERAZOSIN PMS FGNQSW

**24:28.08 DIHYDROPYRIDINES (CALCIUM CHANNEL BLOCKERS)**

**⑤ FELODIPINE**

2.5MG SUSTAINED RELEASE TABLET  
02057778 PLENDIL AZE FNQSW  
02452367 APO-FELODIPINE APX FGNQSW

5MG SUSTAINED RELEASE TABLET  
00851779 PLENDIL AZE FNQSW  
02280264 SANDOZ FELODIPINE SDZ FGNQSW  
02452375 APO-FELODIPINE APX FGNQSW

10MG SUSTAINED RELEASE TABLET



00851787	PLENDIL	AZE	<b>FNQSW</b>
02280272	SANDOZ FELODIPINE	SDZ	<b>FGNQSW</b>
02452383	APO-FELODIPINE	APX	<b>FGNQSW</b>

**24:32.04 ANGIOTENSIN-CONVERTING ENZYME INHIBITORS**

**⑤BENAZEPRIL HCL**

5MG TABLET			
02290332	BENAZEPRIL	AAA	<b>FGNQSW</b>
10MG TABLET			
02290340	BENAZEPRIL	AAA	<b>FGNQSW</b>
20MG TABLET			
02273918	BENAZEPRIL	AAA	<b>FGNQSW</b>

**⑤CAPTOPRIL**

12.5MG TABLET			
01942964	TEVA-CAPTOPRIL	TEV	<b>FGNQSW</b>
25MG TABLET			
01942972	TEVA-CAPTOPRIL	TEV	<b>FGNQSW</b>
50MG TABLET			
01942980	TEVA-CAPTOPRIL	TEV	<b>FGNQSW</b>
100MG TABLET			
01942999	TEVA-CAPTORIL	TEV	<b>FGNQSW</b>

**⑤CILAZAPRIL**

1MG TABLET			
02283778	MYLAN-CILAZAPRIL	MYL	<b>FGNQSW</b>
2.5MG TABLET			
02283786	MYLAN-CILAZAPRIL	MYL	<b>FGNQSW</b>
02291142	APO-CILAZAPRIL	APX	<b>FGNQSW</b>
5MG TABLET			
01911481	INHIBACE	XPI	<b>FNQSW</b>
02283794	MYLAN-CILAZAPRIL	MYL	<b>FGNQSW</b>
02291150	APO-CILAZAPRIL	APX	<b>FGNQSW</b>

**⑤ CILAZAPRIL & HYDROCHLOROTHIAZIDE**

**5MG & 12.5MG TABLET**

02181479	INHIBACE PLUS	XPI	<b>FNQSW</b>
02284987	APO-CILAZAPRIL/HCTZ	APX	<b>FGNQSW</b>
02313731	TEVA-CILAZAPRIL/HCTZ	TEV	<b>FGNQSW</b>

**⑤ ENALAPRIL MALEATE**

**2.5MG TABLET**

02020025	APO-ENALAPRIL	APX	<b>FGNQSW</b>
02291878	ACT-ENALAPRIL	TEV	<b>FGNQSW</b>
02299933	SANDOZ-ENALAPRIL	SDZ	<b>FGNQSW</b>
02352230	RAN-ENALAPRIL	RAN	<b>FGNQSW</b>
02400650	ENALAPRIL	SNS	<b>FGNQSW</b>
02442957	ENALAPRIL	SIV	<b>FGNQSW</b>
02459450	MAR-ENALAPRIL	MAR	<b>FGNQSW</b>
02474786	JAMP-ENALAPRIL	JPC	<b>FGNQSW</b>

**5MG TABLET**

00708879	VASOTEC	MSD	<b>FNQSW</b>
02019884	APO-ENALAPRIL	APX	<b>FGNQSW</b>
02291886	ACT-ENALAPRIL	TEV	<b>FGNQSW</b>
02299941	SANDOZ-ENALAPRIL	SDZ	<b>FGNQSW</b>
02352249	RAN-ENALAPRIL	RAN	<b>FGNQSW</b>
02400669	ENALAPRIL	SNS	<b>FGNQSW</b>
02442965	ENALAPRIL	SIV	<b>FGNQSW</b>
02459469	MAR-ENALAPRIL	MAR	<b>FGNQSW</b>
02474794	JAMP-ENALAPRIL	JPC	<b>FGNQSW</b>

**10MG TABLET**

00670901	VASOTEC	MSD	<b>FNQSW</b>
02019892	APO-ENALAPRIL	APX	<b>FGNQSW</b>
02291894	ACT-ENALAPRIL	TEV	<b>FGNQSW</b>
02299968	SANDOZ-ENALAPRIL	SDZ	<b>FGNQSW</b>
02352257	RAN-ENALAPRIL	RAN	<b>FGNQSW</b>
02400677	ENALAPRIL	SNS	<b>FGNQSW</b>
02442973	ENALAPRIL	SIV	<b>FGNQSW</b>
02444771	MAR-ENALAPRIL	MAR	<b>FGNQSW</b>
02474808	JAMP-ENALAPRIL	JPC	<b>FGNQSW</b>

**20MG TABLET**

00670928	VASOTEC	MSD	<b>FNQSW</b>
02019906	APO-ENALAPRIL	APX	<b>FGNQSW</b>
02291908	ACT-ENALAPRIL	TEV	<b>FGNQSW</b>
02299976	SANDOZ-ENALAPRIL	SDZ	<b>FGNQSW</b>
02352265	RAN-ENALAPRIL	RAN	<b>FGNQSW</b>
02400685	ENALAPRIL	SNS	<b>FGNQSW</b>

02442981	ENALAPRIL	SIV	FGNQSW
02444798	MAR-ENALAPRIL	MAR	FGNQSW
02474816	JAMP-ENALAPRIL	JPC	FGNQSW

**⑤ ENALAPRIL & HYDROCHLOROTHIAZIDE**

5MG & 12.5MG TABLET

02352923	ENALAPRIL MALEATE/HCTZ	AAA	FGNQSW
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10MG & 25MG TABLET

00657298	VASERETIC	MSD	FNQSW
02352931	ENALAPRIL MALEATE/HCTZ	AAA	FGNQSW

**⑤ FOSINOPRIL**

10MG TABLET

02247802	TEVA-FOSINOPRIL	TEV	FGNQSW
02266008	APO-FOSINOPRIL	APX	FGNQSW
02331004	JAMP-FOSINOPRIL	JPC	FGNQSW
02294524	RAN-FOSINOPRIL	RAN	FGNQSW
02459388	FOSINOPRIL	SNS	FGNQSW

20MG TABLET

02247803	TEVA-FOSINOPRIL	TEV	FGNQSW
02266016	APO-FOSINOPRIL	APX	FGNQSW
02331012	JAMP-FOSINOPRIL	JPC	FGNQSW
02294532	RAN-FOSINOPRIL	RAN	FGNQSW
02459396	FOSINOPRIL	SNS	FGNQSW

**⑤ LISINOPRIL**

5MG TABLET

02049333	ZESTRIL	AZE	FNQSW
02217481	APO-LISINOPRIL	APX	FGNQSW
02285118	TEVA-LISINOPRIL (TYPE Z)	TEV	FGNQSW
02361531	JAMP-LISINOPRIL	JPC	FGNQSW
02386232	LISINOPRIL	SIV	FGNQSW
02394472	AURO-LISINOPRIL	ARO	FGNQSW
02525186	LISINOPRIL	SNS	FGNQSW

10MG TABLET

02049376	ZESTRIL	AZE	FNQSW
02217503	APO-LISINOPRIL	APX	FGNQSW
02285126	TEVA-LISINOPRIL (TYPE Z)	TEV	FGNQSW
02294249	RAN-LISINOPRIL	RAN	FGNQSW
02361558	JAMP-LISINOPRIL	JPC	FGNQSW
02386240	LISINOPRIL	SIV	FGNQSW
02394480	AURO-LISINOPRIL	ARO	FGNQSW
02525194	LISINOPRIL	SNS	FGNQSW

20MG TABLET

02049384	ZESTRIL	AZE	FNQSW
02217511	APO-LISINOPRIL	APX	FGNQSW
02285134	TEVA-LISINOPRIL (TYPE Z)	TEV	FGNQSW
02294257	RAN-LISINOPRIL	RAN	FGNQSW
02361566	JAMP-LISINOPRIL	JPC	FGNQSW
02386259	LISINOPRIL	SIV	FGNQSW
02394499	AURO-LISINOPRIL	ARO	FGNQSW
02525208	LISINOPRIL	SNS	FGNQSW

⑤ LISINOPRIL & HYDROCHLOROTHIAZIDE

10MG & 12.5MG TABLET

02103729	ZESTORESTIC	AZE	FNQSW
02301768	TEVA-LISINOPRIL/HCTZ (TYPE Z)	TEV	FGNQSW
02302136	TEVA-LISINOPRIL/HCTZ (TYPE P)	TEV	FGNQSW
02302365	SANDOZ LISINOPRIL/HCT	SDZ	FGNQSW
02362945	LISINOPRIL	SNS	FGNQSW

20MG & 12.5MG TABLET

02045737	ZESTORESTIC	AZE	FNQSW
02301776	TEVA-LISINOPRIL/HCTZ (TYPE Z)	TEV	FGNQSW
02302144	TEVA-LISINOPRIL/HCTZ (TYPE P)	TEV	FGNQSW
02302373	SANDOZ LISINOPRIL/HCT	SDZ	FGNQSW
02362953	LISINOPRIL	SNS	FGNQSW

20MG & 25MG TABLET

02045729	ZESTORESTIC	AZE	FNQSW
02301784	TEVA-LISINOPRIL/HCTZ (TYPE Z)	TEV	FGNQSW
02302152	TEVA-LISINOPRIL/HCTZ (TYPE P)	TEV	FGNQSW
02302381	SANDOZ-LISINOPRIL/HCT	SDZ	FGNQSW
02362961	LISINOPRIL	SNS	FGNQSW

⑤ QUINAPRIL HCL

5MG TABLET

02248499	APO-QUINAPRIL	APX	FGNQSW
02340550	PMS-QUINAPRIL	PMS	FGNQSW

10MG TABLET

02248500	APO-QUINAPRIL	APX	FGNQSW
02340569	PMS-QUINAPRIL	PMS	FGNQSW
02517450	JAMP-QUINAPRIL	JPC	FGNQSW

20MG TABLET

02248501	APO-QUINAPRIL	APX	FGNQSW
02340577	PMS-QUINAPRIL	PMS	FGNQSW

02517469	JAMP-QUINAPRIL	JPC	<b>FGNQSW</b>
40MG TABLET			
02248502	APO-QUINAPRIL	APX	<b>FGNQSW</b>
02340585	PMS-QUINAPRIL	PMS	<b>FGNQSW</b>
02517477	JAMP-QUINAPRIL	JPC	<b>FGNQSW</b>

**⑤QUINAPRIL HCL & HYDROCHLOROTHIAZIDE**

10MG & 12.5MG TABLET			
02408767	APO-QUINAPRIL HCTZ	APX	<b>FGNQSW</b>
02473291	AURO-QUINAPRIL HCTZ	ARO	<b>FGNQSW</b>

20MG & 12.5MG TABLET			
02408775	APO-QUINAPRIL HCTZ	APX	<b>FGNQSW</b>
02473305	AURO-QUINAPRIL HCTZ	ARO	<b>FGNQSW</b>

20MG & 25MG TABLET			
02408783	APO-QUINAPRIL HCTZ	APX	<b>FGNQSW</b>
02473321	AURO-QUINAPRIL HCTZ	ARO	<b>FGNQSW</b>

**⑤RAMIPRIL**

1.25MG CAPSULE			
02221829	ALTACE	VAL	<b>FNQSW</b>
02251515	APO-RAMIPRIL	APX	<b>FGNQSW</b>
02308363	RAMIPRIL	SIV	<b>FGNQSW</b>
02310503	RAN-RAMIPRIL	RAN	<b>FGNQSW</b>
02331101	JAMP RAMIPRIL	JPC	<b>FGNQSW</b>
02387387	AURO-RAMIPRIL	ARO	<b>FGNQSW</b>
02420457	MAR-RAMIPRIL	MAR	<b>FGNQSW</b>
02469057	PHARMA-RAMIPRIL	PMS	<b>FGNQSW</b>

2.5MG CAPSULE			
02221837	ALTACE	VAL	<b>FNQSW</b>
02247945	TEVA-RAMIPRIL	TEV	<b>FGNQSW</b>
02251531	APO-RAMIPRIL	APX	<b>FGNQSW</b>
02287927	RAMIPRIL	SIV	<b>FGNQSW</b>
02310511	RAN-RAMIPRIL	RAN	<b>FGNQSW</b>
02331128	JAMP-RAMIPRIL	JPC	<b>FGNQSW</b>
02374846	RAMIPRIL	SNS	<b>FGNQSW</b>
02387395	AURO-RAMIPRIL	ARO	<b>FGNQSW</b>
02420465	MAR-RAMIPRIL	MAR	<b>FGNQSW</b>
02421305	MINT-RAMIPRIL	MNT	<b>FGNQSW</b>
02469065	PHARMA-RAMIPRIL	PMS	<b>FGNQSW</b>
02477572	AG-RAMIPRIL	ANG	<b>FGNQSW</b>
02486172	NRA-RAMIPRIL	NRA	<b>FGNQSW</b>

5MG CAPSULE

02221845	ALTACE	VAL	FNQSW
02247946	TEVA-RAMIPRIL	TEV	FGNQSW
02251574	APO-RAMIPRIL	APX	FGNQSW
02287935	RAMIPRIL	SIV	FGNQSW
02310538	RAN-RAMIPRIL	RAN	FGNQSW
02331136	JAMP-RAMIPRIL	JPC	FGNQSW
02374854	RAMIPRIL	SNS	FGNQSW
02387409	AURO-RAMIPRIL	ARO	FGNQSW
02420473	MAR-RAMIPRIL	MAR	FGNQSW
02421313	MINT-RAMIPRIL	MNT	FGNQSW
02469073	PHARMA-RAMIPRIL	PMS	FGNQSW
02477580	AG-RAMIPRIL	ANG	FGNQSW
02486180	NRA-RAMIPRIL	NRA	FGNQSW

10MG CAPSULE

02221853	ALTACE	VAL	FNQSW
02247947	TEVA-RAMIPRIL	TEV	FGNQSW
02251582	APO-RAMIPRIL	APX	FGNQSW
02287943	RAMIPRIL	SIV	FGNQSW
02310546	RAN-RAMIPRIL	RAN	FGNQSW
02331144	JAMP-RAMIPRIL	JPC	FGNQSW
02374862	RAMIPRIL	SNS	FGNQSW
02387417	AURO-RAMIPRIL	ARO	FGNQSW
02420481	MAR-RAMIPRIL	MAR	FGNQSW
02421321	MINT-RAMIPRIL	MNT	FGNQSW
02469081	PHARMA-RAMIPRIL	PMS	FGNQSW
02477599	AG-RAMIPRIL	ANG	FGNQSW
02486199	NRA-RAMIPRIL	NRA	FGNQSW

**⑤ RAMIPRIL & HYDROCHLOROTHIAZIDE**

2.5MG & 12.5MG TABLET

02283131	ALTACE HCT	VAL	FNQSW
02449439	RAN-RAMIPRIL HCTZ	RAN	FGNQSW

5MG & 12.5MG TABLET

02283158	ALTACE HCT	VAL	FNQSW
02449447	RAN-RAMIPRIL HCTZ	RAN	FGNQSW

10MG & 12.5MG TABLET

02283166	ALTACE HCT	VAL	FNQSW
02342154	PMS-RAMIPRIL HCTZ	PMS	FGNQSW
02449455	RAN-RAMIPRIL HCTZ	RAN	FGNQSW

5MG & 25MG TABLET

02283174	ALTACE HCT	VAL	FNQSW
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02449463	RAN-RAMIPRIL HCTZ	RAN	<b>FGNQSW</b>
10MG & 25MG TABLET			
02283182	ALTACE HCT	VAL	<b>FNQSW</b>
02342170	PMS-RAMIPRIL-HCTZ	PMS	<b>FGNQSW</b>
02449471	RAN-RAMIPRIL HCTZ	RAN	<b>FGNQSW</b>

**⑤ TRANDOLAPRIL**

0.5MG CAPSULE			
02231457	MAVIK	BGP	<b>FNQSW</b>
02325721	SANDOZ-TRANDOLAPRIL	SDZ	<b>FGNQSW</b>
02357755	PMS-TRANDOLAPRIL	PMS	<b>FGNQSW</b>
02471868	AURO-TRANDOLAPRIL	ARO	<b>FGNQSW</b>

1MG CAPSULE			
02231459	MAVIK	BGP	<b>FNQSW</b>
02325748	SANDOZ-TRANDOLAPRIL	SDZ	<b>FGNQSW</b>
02357763	PMS-TRANDOLAPRIL	PMS	<b>FGNQSW</b>
02471876	AURO-TRANDOLAPRIL	ARO	<b>FGNQSW</b>
02525046	TRANDOLAPRIL	SNS	<b>FGNQSW</b>
02526565	TRANDOLAPRIL	SIV	<b>FGNQSW</b>

2MG CAPSULE			
02231460	MAVIK	BGP	<b>FNQSW</b>
02325756	SANDOZ-TRANDOLAPRIL	SDZ	<b>FGNQSW</b>
02357771	PMS-TRANDOLAPRIL	PMS	<b>FGNQSW</b>
02471884	AURO-TRANDOLAPRIL	ARO	<b>FGNQSW</b>
02525054	TRANDOLAPRIL	SNS	<b>FGNQSW</b>
02526573	TRANDOLAPRIL	SIV	<b>FGNQSW</b>

4MG CAPSULE			
02239267	MAVIK	BGP	<b>FNQSW</b>
02325764	SANDOZ-TRANDOLAPRIL	SDZ	<b>FGNQSW</b>
02357798	PMS-TRANDOLAPRIL	PMS	<b>FGNQSW</b>
02471892	AURO-TRANDOLAPRIL	ARO	<b>FGNQSW</b>
02525070	TRANDOLAPRIL	SNS	<b>FGNQSW</b>
02526581	TRANDOLAPRIL	SIV	<b>FGNQSW</b>

**24:32.08 ANGIOTENSIN II RECEPTOR ANTAGONISTS**

**⑤ CANDESARTAN CILEXETIL**

4MG TABLET			
02239090	ATACAND	XPI	<b>FNQSW</b>
02326957	SANDOZ-CANDESARTAN	SDZ	<b>FGNQSW</b>

02365340	APO-CANDESARTAN	APX	<b>FGNQSW</b>
02379260	CANDESARTAN CILEXETIL	ACH	<b>FGNQSW</b>
02380684	RAN-CANDESARTAN	RAN	<b>FGNQSW</b>
02388901	CANDESARTAN	SNS	<b>FGNQSW</b>
02391171	PMS-CANDESARTAN	PMS	<b>FGNQSW</b>
02445786	AURO-CANDESARTAN	ARO	<b>FGNQSW</b>
02476908	MINT-CANDESARTAN	MNT	<b>FGNQSW</b>
02528258	CANDESARTAN	SIV	<b>FGNQSW</b>
02541289	APO-CANDESARTAN	APX	<b>FGNQSW</b>

**8MG TABLET**

02239091	ATACAND	XPI	<b>FNQSW</b>
02326965	SANDOZ-CANDESARTAN	SDZ	<b>FGNQSW</b>
02365359	APO-CANDESARTAN	APX	<b>FGNQSW</b>
02366312	TEVA-CANDESARTAN	TEV	<b>FGNQSW</b>
02379279	CANDESARTAN CILEXETIL	ACH	<b>FGNQSW</b>
02380692	RAN-CANDESARTAN	RAN	<b>FGNQSW</b>
02386518	JAMP-CANDESARTAN	JPC	<b>FGNQSW</b>
02388707	CANDESARTAN	SIV	<b>FGNQSW</b>
02388928	CANDESARTAN	SNS	<b>FGNQSW</b>
02391198	PMS-CANDESARTAN	PMS	<b>FGNQSW</b>
02445794	AURO-CANDESARTAN	ARO	<b>FGNQSW</b>
02476916	MINT-CANDESARTAN	MNT	<b>FGNQSW</b>
02527014	NRA-CANDESARTAN	NRA	<b>FGNQSW</b>
02541297	APO-CANDESARTAN	APX	<b>FGNQSW</b>

**16MG TABLET**

02239092	ATACAND	XPI	<b>FNQSW</b>
02326973	SANDOZ-CANDESARTAN	SDZ	<b>FGNQSW</b>
02365367	APO-CANDESARTAN	APX	<b>FGNQSW</b>
02366320	TEVA-CANDESARTAN	TEV	<b>FGNQSW</b>
02379287	CANDESARTAN CILEXETIL	ACH	<b>FGNQSW</b>
02380706	RAN-CANDESARTAN	RAN	<b>FGNQSW</b>
02386526	JAMP-CANDESARTAN	JPC	<b>FGNQSW</b>
02388715	CANDESARTAN	SIV	<b>FGNQSW</b>
02388936	CANDESARTAN	SNS	<b>FGNQSW</b>
02391201	PMS-CANDESARTAN	PMS	<b>FGNQSW</b>
02445808	AURO-CANDESARTAN	ARO	<b>FGNQSW</b>
02476924	MINT-CANDESARTAN	MNT	<b>FGNQSW</b>
02527022	NRA-CANDESARTAN	NRA	<b>FGNQSW</b>
02541300	APO-CANDESARTAN	APX	<b>FGNQSW</b>

**32MG TABLET**

02311658	ATACAND	XPI	<b>FNQSW</b>
02366339	TEVA-CANDESARTAN	TEV	<b>FGNQSW</b>
02379295	CANDESARTAN CILEXETIL	ACH	<b>FGNQSW</b>



02380714	RAN-CANDESARTAN	RAN	<b>FGNQSW</b>
02386534	JAMP-CANDESARTAN	JPC	<b>FGNQSW</b>
02391228	PMS-CANDESARTAN	PMS	<b>FGNQSW</b>
02399105	APO-CANDESARTAN	APX	<b>FGNQSW</b>
02417340	SANDOZ-CANDESARTAN	SDZ	<b>FGNQSW</b>
02435845	CANDESARTAN	SNS	<b>FGNQSW</b>
02445816	AURO-CANDESARTAN	ARO	<b>FGNQSW</b>
02476932	MINT-CANDESARTAN	MNT	<b>FGNQSW</b>
02528266	CANDESARTAN	SIV	<b>FGNQSW</b>
02527030	NRA-CANDESARTAN	NRA	<b>FGNQSW</b>
02541319	APO-CANDESARTAN	APX	<b>FGNQSW</b>

**⑤ CANDESARTAN CILEXETIL & HYDROCHLOROTHIAZIDE**

**16MG & 12.5MG TABLET**

02244021	ATACAND PLUS	XPI	<b>FNQSW</b>
02327902	SANDOZ-CANDESARTAN PLUS	SDZ	<b>FGNQSW</b>
02391295	PMS-CANDESARTAN HCTZ	PMS	<b>FGNQSW</b>
02394804	CANDESARTAN/HCTZ	SNS	<b>FGNQSW</b>
02394812	CANDESARTAN/HCTZ	SIV	<b>FGNQSW</b>
02395541	TEVA-CANDESARTAN HCTZ	TEV	<b>FGNQSW</b>
02421038	AURO-CANDESARTAN HCT	ARO	<b>FGNQSW</b>
02473240	JAMP-CANDESARTAN HCT	JPC	<b>FGNQSW</b>
02531240	NRA-CANDESARTAN HCTZ	NRA	<b>FGNQSW</b>

**32MG & 12.5MG TABLET**

02332922	ATACAND PLUS	XPI	<b>FNQSW</b>
02395568	TEVA-CANDESARTAN HCTZ	TEV	<b>FGNQSW</b>
02420732	SANDOZ-CANDESARTAN HCTZ	SDZ	<b>FGNQSW</b>
02421046	AURO-CANDESARTAN HCT	ARO	<b>FGNQSW</b>
02473259	JAMP-CANDESARTAN HCT	JPC	<b>FGNQSW</b>
02536064	CANDESARTAN HCTZ	SNS	<b>FGNQSW</b>
02531259	NRA-CANDESARTAN HCTZ	NRA	<b>FGNQSW</b>

**32MG & 25MG TABLET**

02332957	ATACAND PLUS	XPI	<b>FNQSW</b>
02420740	SANDOZ-CANDESARTAN HCTZ	SDZ	<b>FGNQSW</b>
02421054	AURO-CANDESARTAN HCT	ARO	<b>FGNQSW</b>
02473267	JAMP-CANDESARTAN HCT	JPC	<b>FGNQSW</b>
02531267	NRA-CANDESARTAN HCTZ	NRA	<b>FGNQSW</b>

**⑤ EPROSARTAN MESYLATE**

**400MG TABLET**

02240432	TEVETEN	BGP	<b>FNQSW</b>
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**600MG TABLET**

02243942	TEVETEN	BGP	<b>FNQSW</b>
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02337428	SANDOZ-IRBESARTAN HCT	SDZ	<b>FGNQSW</b>
02372886	IRBESARTAN HCTZ	SNS	<b>FGNQSW</b>
02385317	IRBESARTAN HCT	SIV	<b>FGNQSW</b>
02447878	AURO-IRBESARTAN HCT	ARO	<b>FGNQSW</b>

**300MG & 12.5MG TABLET**

02241819	AVALIDE	AVN	<b>FNQSW</b>
02328526	PMS-IRBESARTAN HCTZ	PMS	<b>FGNQSW</b>
02330520	TEVA-IRBESARTAN HCTZ	TEV	<b>FGNQSW</b>
02337436	SANDOZ-IRBESARTAN HCT	SDZ	<b>FGNQSW</b>
02372894	IRBESARTAN HCTZ	SNS	<b>FGNQSW</b>
02385325	IRBESARTAN HCT	SIV	<b>FGNQSW</b>
02447886	AURO-IRBESARTAN HCT	ARO	<b>FGNQSW</b>

**300MG & 25MG TABLET**

02328534	PMS-IRBESARTAN HCTZ	PMS	<b>FGNQSW</b>
02330539	TEVA-IRBESARTAN HCTZ	TEV	<b>FGNQSW</b>
02337444	SANDOZ-IRBESARTAN HCT	SDZ	<b>FGNQSW</b>
02372908	IRBESARTAN HCTZ	SNS	<b>FGNQSW</b>
02385333	IRBESARTAN HCT	SIV	<b>FGNQSW</b>
02447894	AURO-IRBESARTAN HCT	ARO	<b>FGNQSW</b>

**⑤ LOSARTAN POTASSIUM**

**25MG TABLET**

02182815	COZAAR	MSD	<b>FNQSW</b>
02313332	SANDOZ-LOSARTAN	SDZ	<b>FGNQSW</b>
02309750	PMS-LOSARTAN	PMS	<b>FGNQSW</b>
02380838	TEVA-LOSARTAN	TEV	<b>FGNQSW</b>
02388790	LOSARTAN	SIV	<b>FGNQSW</b>
02388863	LOSARTAN	SNS	<b>FGNQSW</b>
02398834	JAMP-LOSARTAN	JPC	<b>FGNQSW</b>
02403323	AURO-LOSARTAN	ARO	<b>FGNQSW</b>
02405733	MINT-LOSARTAN	MNT	<b>FGNQSW</b>

**50MG TABLET**

02182874	COZAAR	MSD	<b>FNQSW</b>
02309769	PMS-LOSARTAN	PMS	<b>FGNQSW</b>
02313340	SANDOZ-LOSARTAN	SDZ	<b>FGNQSW</b>
02357968	TEVA-LOSARTAN	TEV	<b>FGNQSW</b>
02388804	LOSARTAN	SIV	<b>FGNQSW</b>
02388871	LOSARTAN	SNS	<b>FGNQSW</b>
02398842	JAMP-LOSARTAN	JPC	<b>FGNQSW</b>
02403331	AURO-LOSARTAN	ARO	<b>FGNQSW</b>
02405741	MINT-LOSARTAN	MNT	<b>FGNQSW</b>

**100MG TABLET**

02182882	COZAAR	MSD	<b>FNQSW</b>
02309777	PMS-LOSARTAN	PMS	<b>FGNQSW</b>
02313359	SANDOZ-LOSARTAN	SDZ	<b>FGNQSW</b>
02357976	TEVA-LOSARTAN	TEV	<b>FGNQSW</b>
02388812	LOSARTAN	SIV	<b>FGNQSW</b>
02388898	LOSARTAN	SNS	<b>FGNQSW</b>
02398850	JAMP-LOSARTAN	JPC	<b>FGNQSW</b>
02403358	AURO-LOSARTAN	ARO	<b>FGNQSW</b>
02405768	MINT-LOSARTAN	MNT	<b>FGNQSW</b>

**⑤ LOSARTAN POTASSIUM & HYDROCHLOROTHIAZIDE**

**50MG & 12.5MG TABLET**

02230047	HYZAAR	MSD	<b>FNQSW</b>
02313375	SANDOZ-LOSARTAN HCT	SDZ	<b>FGNQSW</b>
02358263	TEVA-LOSARTAN HCTZ	TEV	<b>FGNQSW</b>
02388960	LOSARTAN HCTZ	SIV	<b>FGNQSW</b>
02389657	MINT-LOSARTAN HCTZ	MNT	<b>FGNQSW</b>
02392224	PMS-LOSARTAN HCTZ	PMS	<b>FGNQSW</b>
02423642	AURO-LOSARTAN HCT	ARO	<b>FGNQSW</b>
02427648	LOSARTAN-HCTZ	SNS	<b>FGNQSW</b>

**100MG & 12.5MG TABLET**

02297841	HYZAAR	MSD	<b>FNQSW</b>
02362449	SANDOZ-LOSARTAN HCT	SDZ	<b>FGNQSW</b>
02377144	TEVA-LOSARTAN HCTZ	TEV	<b>FGNQSW</b>
02388979	LOSARTAN HCTZ	SIV	<b>FGNQSW</b>
02389665	MINT-LOSARTAN HCTZ	MNT	<b>FGNQSW</b>
02392232	PMS-LOSARTAN HCTZ	PMS	<b>FGNQSW</b>
02423650	AURO-LOSARTAN HCT	ARO	<b>FGNQSW</b>
02427656	LOSARTAN-HCTZ	SNS	<b>FGNQSW</b>

**100MG & 25MG TABLET**

02241007	HYZAAR DS	MSD	<b>FNQSW</b>
02313383	SANDOZ-LOSARTAN HCT	SDZ	<b>FGNQSW</b>
02377152	TEVA-LOSARTAN HCTZ	TEV	<b>FGNQSW</b>
02388987	LOSARTAN HCTZ	SIV	<b>FGNQSW</b>
02389673	MINT-LOSARTAN HCTZ	MNT	<b>FGNQSW</b>
02392240	PMS-LOSARTAN HCTZ	PMS	<b>FGNQSW</b>
02423669	AURO-LOSARTAN HCT	ARO	<b>FGNQSW</b>
02427664	LOSARTAN-HCTZ	SNS	<b>FGNQSW</b>

**⑤ OLMESARTAN**

**20MG TABLET**

02318660	OLMETEC	MSD	<b>FNQSW</b>
02442191	TEVA-OLMESARTAN	TEV	<b>FGNQSW</b>
02443414	SANDOZ-OLMESARTAN	SDZ	<b>FGNQSW</b>

02443864	AURO-OLMESARTAN	ARO	FGNQSW
02453452	APO-OLMESARTAN	APX	FGNQSW
02456311	ACH-OLMESARTAN	ACH	FGNQSW
02461307	PMS-OLMESARTAN	PMS	FGNQSW
02461641	JAMP-OLMESARTAN	JPC	FGNQSW
02469812	GLN-OLMESARTAN	GLM	FGNQSW
02481057	OLMESARTAN	SNS	FGNQSW
02499258	NRA-OLMESARTAN	NRA	FGNQSW

40MG TABLET

02318679	OLMETEC	MSD	FNQSW
02442205	TEVA-OLMESARTAN	TEV	FGNQSW
02443422	SANDOZ-OLMESARTAN	SDZ	FGNQSW
02443872	AURO-OLMESARTAN	ARO	FGNQSW
02453460	APO-OLMESARTAN	APX	FGNQSW
02456338	ACH-OLMESARTAN	ACH	FGNQSW
02461315	PMS-OLMESARTAN	PMS	FGNQSW
02461668	JAMP-OLMESARTAN	JPC	FGNQSW
02469820	GLN-OLMESARTAN	GLM	FGNQSW
02481065	OLMESARTAN	SNS	FGNQSW
02499266	NRA-OLMESARTAN	NRA	FGNQSW

**⑤ OLMESARTAN & HYDROCHLOROTHIAZIDE**

20MG & 12.5MG TABLET

02319616	OLMETEC PLUS	MSD	FNQSW
02443112	ACT-OLMESARTAN/HCTZ	TEV	FGNQSW
02453606	APO-OLMESARTAN/HCTZ	APX	FGNQSW
02468948	ACH-OLMESARTAN/HCTZ	ACH	FGNQSW
02476487	AURO-OLMESARTAN HCTZ	ARO	FGNQSW
02508273	NRA-OLMESARTAN/HCTZ	NRA	FGNQSW
02509601	OLMESARTAN-HCTZ	SNS	FGNQSW
02526468	PRZ-OLMESARTAN-HCTZ	PRZ	FGNQSW

40MG & 12.5MG TABLET

02319624	OLMETEC PLUS	MSD	FNQSW
02443120	ACT-OLMESARTAN/HCTZ	TEV	FGNQSW
02453614	APO-OLMESARTAN/HCTZ	APX	FGNQSW
02468956	ACH-OLMESARTAN/HCTZ	ACH	FGNQSW
02476495	AURO-OLMESARTAN HCTZ	ARO	FGNQSW
02508281	NRA-OLMESARTAN/HCTZ	NRA	FGNQSW
02509636	OLMESARTAN-HCTZ	SNS	FGNQSW
02526476	PRZ-OLMESARTAN-HCTZ	PRZ	FGNQSW

40MG & 25MG TABLET

02319632	OLMETEC PLUS	MSD	FNQSW
02443139	ACT-OLMESARTAN/HCTZ	TEV	FGNQSW

02453622	APO-OLMESARTAN/HCTZ	APX	<b>FGNQSW</b>
02468964	ACH-OLMESARTAN/HCTZ	ACH	<b>FGNQSW</b>
02476509	AURO-OLMESARTAN HCTZ	ARO	<b>FGNQSW</b>
02508303	NRA-OLMESARTAN/HCTZ	NRA	<b>FGNQSW</b>
02509628	OLMESARTAN-HCTZ	SNS	<b>FGNQSW</b>
02526484	PRZ-OLMESARTAN-HCTZ	PRZ	<b>FGNQSW</b>

**⑤ SACUBITRIL & VALSARTAN**  
[SEE APPENDIX A](#) FOR SA CRITERIA  
 24MG & 26MG TABLET

02446928	ENTRESTO (SA)	NVR	<b>FNQSW</b>
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49MG & 51MG TABLET

02446936	ENTRESTO (SA)	NVR	<b>FNQSW</b>
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97MG & 103MG TABLET

02446944	ENTRESTO (SA)	NVR	<b>FNQSW</b>
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**⑤ TELMISARTAN**

40MG TABLET

02240769	MICARDIS	BOE	<b>FNQSW</b>
02320177	TEVA-TELMISARTAN	TEV	<b>FGNQSW</b>
02375958	SANDOZ-TELMISARTAN	SDZ	<b>FGNQSW</b>
02386755	JAMP-TELMISARTAN	JPC	<b>FGNQSW</b>
02388944	TELMISARTAN	SNS	<b>FGNQSW</b>
02390345	TELMISARTAN	SIV	<b>FGNQSW</b>
02407485	TELMISARTAN	ACH	<b>FGNQSW</b>
02453568	AURO-TELMISARTAN	ARO	<b>FGNQSW</b>
02486369	MINT-TELMISARTAN	MNT	<b>FGNQSW</b>
02499622	PMS-TELMISARTAN	PMS	<b>FGNQSW</b>
02503794	NRA-TELMISARTAN	NRA	<b>FGNQSW</b>

80MG TABLET

02240770	MICARDIS	BOE	<b>FNQSW</b>
02320185	TEVA-TELMISARTAN	TEV	<b>FGNQSW</b>
02375966	SANDOZ-TELMISARTAN	SDZ	<b>FGNQSW</b>
02386763	JAMP-TELMISARTAN	JPC	<b>FGNQSW</b>
02388952	TELMISARTAN	SNS	<b>FGNQSW</b>
02390353	TELMISARTAN	SIV	<b>FGNQSW</b>
02407493	TELMISARTAN	ACH	<b>FGNQSW</b>
02453576	AURO-TELMISARTAN	ARO	<b>FGNQSW</b>
02486377	MINT-TELMISARTAN	MNT	<b>FGNQSW</b>
02499630	PMS-TELMISARTAN	PMS	<b>FGNQSW</b>
02503808	NRA-TELMISARTAN	NRA	<b>FGNQSW</b>

**⑤ TELMISARTAN & AMLODIPINE**

40/5MG TABLET			
02371022	TWYNSTA	BOE	FNQSW
40/10MG TABLET			
02371030	TWYNSTA	BOE	FNQSW
80/5MG TABLET			
02371049	TWYNSTA	BOE	FNQSW
80/10MG TABLET			
02371057	TWYNSTA	BOE	FNQSW

**⑤ TELMISARTAN & HYDROCHLOROTHIAZIDE**

80MG & 12.5MG TABLET			
02244344	MICARDIS PLUS	BOE	FNQSW
02330288	TEVA-TELMISARTAN HCTZ	TEV	FGNQSW
02389940	JAMP-TELMISARTAN HCT	JPC	FGNQSW
02390302	TELMISARTAN-HCTZ	SIV	FGNQSW
02393557	SANDOZ-TELMISARTAN HCT	SDZ	FGNQSW
02395355	TELMISARTAN HCTZ	SNS	FGNQSW
02419114	ACH-TELMISARTAN HCTZ	ACH	FGNQSW
02456389	AURO-TELMISARTAN HCTZ	ARO	FGNQSW
02504146	NRA-TELMISARTAN HCTZ	NRA	FGNQSW

80MG & 25 MG TABLET			
02318709	MICARDIS PLUS	BOE	FNQSW
02379252	TEVA-TELMISARTAN HCTZ	TEV	FGNQSW
02389959	JAMP-TELMISARTAN HCT	JPC	FGNQSW
02390310	TELMISARTAN-HCTZ	SIV	FGNQSW
02393565	SANDOZ-TELMISARTAN HCT	SDZ	FGNQSW
02395363	TELMISARTAN HCTZ	SNS	FGNQSW
02419122	ACH-TELMISARTAN HCTZ	ACH	FGNQSW
02456397	AURO-TELMISARTAN HCTZ	ARO	FGNQSW
02504138	NRA-TELMISARTAN HCTZ	NRA	FGNQSW

**⑤ VALSARTAN**

40MG TABLET			
02270528	DIOVAN	NVR	FNQSW
02356740	SANDOZ-VALSARTAN	SDZ	FGNQSW
02356643	TEVA-VALSARTAN	TEV	FGNQSW
02363062	TARO-VALSARTAN	SUN	FGNQSW
02366940	VALSARTAN	SNS	FGNQSW
02384523	VALSARTAN	SIV	FGNQSW
02414201	AURO-VALSARTAN	ARO	FGNQSW
02524511	M-VALSARTAN	MRA	FGNQSW

80MG TABLET

02244781	DIOVAN	NVR	<b>FNQSW</b>
02356759	SANDOZ-VALSARTAN	SDZ	<b>FGNQSW</b>
02356651	TEVA-VALSARTAN	TEV	<b>FGNQSW</b>
02363100	TARO-VALSARTAN	SUN	<b>FGNQSW</b>
02366959	VALSARTAN	SNS	<b>FGNQSW</b>
02384531	VALSARTAN	SIV	<b>FGNQSW</b>
02414228	AURO-VALSARTAN	ARO	<b>FGNQSW</b>
02524538	M-VALSARTAN	MRA	<b>FGNQSW</b>

160MG TABLET

02244782	DIOVAN	NVR	<b>FNQSW</b>
02356767	SANDOZ-VALSARTAN	SDZ	<b>FGNQSW</b>
02356678	TEVA-VALSARTAN	TEV	<b>FGNQSW</b>
02363119	TARO-VALSARTAN	SUN	<b>FGNQSW</b>
02366967	VALSARTAN	SNS	<b>FGNQSW</b>
02384558	VALSARTAN	SIV	<b>FGNQSW</b>
02414236	AURO-VALSARTAN	ARO	<b>FGNQSW</b>
02524546	M-VALSARTAN	MRA	<b>FGNQSW</b>

320MG TABLET

02289504	DIOVAN	NVR	<b>FNQSW</b>
02356775	SANDOZ-VALSARTAN	SDZ	<b>FGNQSW</b>
02356686	TEVA-VALSARTAN	TEV	<b>FGNQSW</b>
02366975	VALSARTAN	SNS	<b>FGNQSW</b>
02384566	VALSARTAN	SIV	<b>FGNQSW</b>
02414244	AURO-VALSARTAN	ARO	<b>FGNQSW</b>

**⑤ VALSARTAN & HYDROCHLORTHIAZIDE**

80MG & 12.5MG TABLET

02241900	DIOVAN-HCT	NVR	<b>FNQSW</b>
02356694	SANDOZ-VALSARTAN HCT	SDZ	<b>FGNQSW</b>
02356996	TEVA-VALSARTAN HCTZ	TEV	<b>FGNQSW</b>
02367009	VALSARTAN HCTZ	SNS	<b>FGNQSW</b>
02384736	VALSARTAN HCT	SIV	<b>FGNQSW</b>
02408112	AURO-VALSARTAN HCT	ARO	<b>FGNQSW</b>

160MG & 12.5MG TABLET

02241901	DIOVAN-HCT	NVR	<b>FNQSW</b>
02356708	SANDOZ-VALSARTAN HCT	SDZ	<b>FGNQSW</b>
02357003	TEVA-VALSARTAN HCTZ	TEV	<b>FGNQSW</b>
02367017	VALSARTAN HCTZ	SNS	<b>FGNQSW</b>
02384744	VALSARTAN HCT	SIV	<b>FGNQSW</b>
02408120	AURO-VALSARTAN HCT	ARO	<b>FGNQSW</b>



160MG & 25MG TABLET

02246955	DIOVAN-HCT	NVR	FNQSW
02356716	SANDOZ-VALSARTAN HCT	SDZ	FGNQSW
02357011	TEVA-VALSARTAN HCTZ	TEV	FGNQSW
02367025	VALSARTAN HCTZ	SNS	FGNQSW
02384752	VALSARTAN HCT	SIV	FGNQSW
02408139	AURO-VALSARTAN HCT	ARO	FGNQSW

320MG & 12.5MG TABLET

02308908	DIOVAN-HCT	NVR	FNQSW
02356724	SANDOZ-VALSARTAN HCT	SDZ	FGNQSW
02357038	TEVA-VALSARTAN HCTZ	TEV	FGNQSW
02367033	VALSARTAN HCTZ	SNS	FGNQSW
02384760	VALSARTAN HCT	SIV	FGNQSW
02408147	AURO-VALSARTAN HCT	ARO	FGNQSW

320MG & 25MG TABLET

02308916	DIOVAN-HCT	NVR	FNQSW
02356732	SANDOZ-VALSARTAN HCT	SDZ	FGNQSW
02357046	TEVA-VALSARTAN HCTZ	TEV	FGNQSW
02367041	VALSARTAN HCTZ	SNS	FGNQSW
02408155	AURO-VALSARTAN HCT	ARO	FGNQSW

**28:08.04 NONSTEROIDAL ANTI INFLAMMATORY AGENTS**

**ACETYLSALICYLIC ACID**

325MG TABLET

00999963	ASA (DIN for billing purposes only)	NW
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81MG ENTERIC COATED TABLET

00999971	ASA (DIN for billing purposes only)	NW
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**CELECOXIB**

100MG CAPSULE

02239941	CELEBREX	UJC	FNQSW
02355442	PMS-CELECOXIB	PMS	FGNQSW
02412373	RAN-CELECOXIB	RAN	FGNQSW
02412497	MINT-CELECOXIB	MNT	FGNQSW
02418932	APO-CELECOXIB	APX	FGNQSW
02420058	MAR-CELECOXIB	MAR	FGNQSW
02420155	ACT-CELECOXIB	TEV	FGNQSW
02424533	JAMP-CELECOXIB	JPC	FGNQSW
02429675	CELECOXIB	SIV	FGNQSW

02436299	CELECOXIB	SNS	<b>FGNQSW</b>
02437570	AG-CELECOXIB	ANG	<b>FGNQSW</b>
02445670	AURO-CELECOXIB	ARO	<b>FGNQSW</b>
02479737	NRA-CELECOXIB	NRA	<b>FGNQSW</b>
02495465	M-CELECOXIB	MRA	<b>FGNQSW</b>
02517116	PMSC-CELECOXIB	PMS	<b>FGNQSW</b>

**200MG CAPSULE**

02239942	CELEBREX	UJC	<b>FNQSW</b>
02355450	PMS-CELECOXIB	PMS	<b>FGNQSW</b>
02412381	RAN-CELECOXIB	RAN	<b>FGNQSW</b>
02412500	MINT-CELECOXIB	MNT	<b>FGNQSW</b>
02418940	APO-CELECOXIB	APO	<b>FGNQSW</b>
02420066	MAR-CELECOXIB	MAR	<b>FGNQSW</b>
02420163	ACT-CELECOXIB	TEV	<b>FGNQSW</b>
02424541	JAMP-CELECOXIB	JPC	<b>FGNQSW</b>
02429683	CELECOXIB	SIV	<b>FGNQSW</b>
02436302	CELECOXIB	SNS	<b>FGNQSW</b>
02437589	AG-CELECOXIB	ANG	<b>FGNQSW</b>
02445689	AURO-CELECOXIB	ARO	<b>FGNQSW</b>
02479745	NRA-CELECOXIB	NRA	<b>FGNQSW</b>
02495473	M-CELECOXIB	MRA	<b>FGNQSW</b>
02517124	PMSC-CELECOXIB	PMS	<b>FGNQSW</b>

**DICLOFENAC SODIUM**

**25MG ENTERIC COATED TABLET**

00808539	TEVA-DICLOFENAC EC	TEV	<b>FGNQSW</b>
00839175	APO-DICLO	APX	<b>FGNQSW</b>
02302616	PMS-DICLOFENAC	PMS	<b>FGNQSW</b>

**50MG ENTERIC COATED TABLET**

00808547	TEVA DIFENAC	TEV	<b>FGNQSW</b>
00839183	APO-DICLO	APX	<b>FGNQSW</b>
02302624	PMS-DICLOFENAC	PMS	<b>FGNQSW</b>

**75MG SUSTAINED RELEASE TABLET**

02158582	TEVA DIFENAC SR	TEV	<b>FGNQSW</b>
02162814	APO-DICLO SR	APX	<b>FGNQSW</b>
02261901	SANDOZ-DICLOFENAC	SDZ	<b>FGNQSW</b>

**100MG SUSTAINED RELEASE TABLET**

02091194	APO-DICLO SR	APX	<b>FGNQSW</b>
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**50MG SUPPOSITORY**

00632724	VOLTAREN	NVR	<b>FNQSW</b>
02261928	SANDOZ-DICLOFENAC	SDZ	<b>FGNQSW</b>

**DICLOFENAC & MISOPROSTOL**

50MG/200MG TABLET

01917056	ARTHROTEC	PFI	<b>FNQSW</b>
02341689	GD-DICLOFENAC/MISOPROSTOL	GMD	<b>FGNQSW</b>
02413469	PMS-DICLOFENAC/MISOPROSTOL	PMS	<b>FGNQSW</b>

75MG/200MG TABLET

02229837	ARTHROTEC	PFI	<b>FNQSW</b>
02341697	GD-DICLOFENAC/MISOPROSTOL	GMD	<b>FGNQSW</b>
02413477	PMS-DICLOFENAC/MISOPROSTOL	PMS	<b>FGNQSW</b>

**FLURBIPROFEN**

50MG TABLET

01912046	FLURBIPROFEN	AAA	<b>FGNQSW</b>
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100MG TABLET

01912038	FLURBIPROFEN	AAA	<b>FGNQSW</b>
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**IBUPROFEN**

300MG TABLET

00999986	IBUPROFEN (DIN for billing purposes only)		<b>NW</b>
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400MG TABLET

00999987	IBUPROFEN (DIN for billing purposes only)		<b>NW</b>
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600MG TABLET

00585114	APO-IBUPROFEN	APX	<b>FNQSW</b>
00629359	TEVA-PROFEN	TEV	<b>FNQSW</b>

**INDOMETHACIN**

25MG CAPSULE

00337420	TEVA-METHACIN	TEV	<b>FGNQSW</b>
02461811	MINT-INDOMETHACIN	MNT	<b>FGNQSW</b>

50MG CAPSULE

00337439	TEVA-METHACIN	TEV	<b>FGNQSW</b>
02461536	MINT-INDOMETHACIN	MNT	<b>FGNQSW</b>
02499223	AURO-INDOMETHACIN	ARO	<b>FGNQSW</b>

**KETOPROFEN**

50MG CAPSULE

00790427	KETOPROFEN	AAA	<b>FGNQSW</b>
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50MG ENTERIC COATED TABLET

00790435	KETOPROFEN-E	AAA	<b>FGNQSW</b>
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100MG ENTERIC COATED TABLET 00842664	KETOPROFEN-E	AAA	<b>FGNQSW</b>
<b>MEFENAMIC ACID</b>			
250MG CAPSULE 00155225	PONSTAN	AAA	<b>FQW</b>
<b>MELOXICAM</b>			
7.5MG TABLET 02248267	PMS-MELOXICAM	PMS	<b>FGNQSW</b>
02248973	APO-MELOXICAM	APX	<b>FGNQSW</b>
02258315	TEVA-MELOXICAM	TEV	<b>FGNQSW</b>
02353148	MELOXICAM	SNS	<b>FGNQSW</b>
02390884	AURO-MELOXICAM	ARO	<b>FGNQSW</b>
15MG TABLET 02248268	PMS-MELOXICAM	PMS	<b>FGNQSW</b>
02248974	APO-MELOXICAM	APX	<b>FGNQSW</b>
02258323	TEVA-MELOXICAM	TEV	<b>FGNQSW</b>
02353156	MELOXICAM	SNS	<b>FGNQSW</b>
02390892	AURO-MELOXICAM	ARO	<b>FGNQSW</b>
<b>NABUMETONE</b>			
500MG TABLET 02238639	NABUMETONE	AAA	<b>FGNQSW</b>
<b>NAPROXEN</b>			
125MG TABLET 00522678	APO-NAPROXEN	APX	<b>FGNQSW</b>
250MG TABLET 00522651	APO-NAPROXEN	APX	<b>FGNQSW</b>
00565350	TEVA-NAPROX	TEV	<b>FGNQSW</b>
375MG TABLET 00600806	APO-NAPROXEN	APX	<b>FGNQSW</b>
00627097	TEVA NAPROX	TEV	<b>FGNQSW</b>
500MG TABLET 00589861	TEVA-NAPROX	TEV	<b>FGNQSW</b>
00592277	APO-NAPROXEN	APX	<b>FGNQSW</b>
250MG ENTERIC COATED TABLET 02243312	TEVA-NAPROXEN EC	TEV	<b>FGNQSW</b>

375MG ENTERIC COATED TABLET			
02162415	NAPROSYN-E	MTP	<b>FNQSW</b>
02243313	TEVA-NAPROXEN EC	TEV	<b>FGNQSW</b>
02246700	APO-NAPROXEN EC	APX	<b>FGNQSW</b>

500MG ENTERIC COATED TABLET			
02162423	NAPROSYN-E	MTP	<b>FNQSW</b>
02243314	TEVA-NAPROXEN EC	TEV	<b>FGNQSW</b>
02246701	APO-NAPROXEN EC	APX	<b>FGNQSW</b>

750MG SUSTAINED RELEASE TABLET			
02162466	NAPROSYN SR	MTP	<b>FNQSW</b>

**PIROXICAM**

10MG CAPSULE			
00695718	TEVA-PIROXICAM	TEV	<b>FGNQSW</b>

20MG CAPSULE			
00695696	TEVA-PIROXICAM	TEV	<b>FGNQSW</b>

**SULINDAC**

150MG TABLET			
00745588	TEVA-SULINDAC	TEV	<b>FGNQSW</b>

200MG TABLET			
00745596	TEVA-SULINDAC	TEV	<b>FGNQSW</b>

**TIAPROFENIC ACID**

200MG TABLET			
02179679	TEVA-TIAPROFENIC ACID	TEV	<b>FGNQSW</b>

300MG TABLET			
02179687	TEVA-TIAPROFENIC ACID	TEV	<b>FGNQSW</b>

**28:08.08 OPIATE AGONISTS (NARCOTIC ANALGESICS)**

**ACETAMINOPHEN & CODEINE**

300MG & 60MG TABLET			
00621463	TEVA-LENOLTEC NO.4	TEV	<b>FNQSW</b>

**ACETAMINOPHEN COMPOUND WITH CODEINE**

15MG CODEINE TABLET			
00653241	TEVA-LENOLTEC NO.2	TEV	<b>FNQSW</b>

30MG CODEINE TABLET  
00653276 TEVA-LENOLTEC NO.3 TEV **FNQSW**

**CODEINE**

15MG TABLET  
00593435 TEVA-CODEINE TEV **FNQSW**

30MG TABLET  
00593451 TEVA-CODEINE TEV **FNQSW**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG CONTROLLED RELEASE TABLET  
02230302 CODEINE CONTIN (SA) PFR **FNQSW**

100MG CONTROLLED RELEASE TABLET  
02163748 CODEINE CONTIN (SA) PFR **FNQSW**

150MG CONTROLLED RELEASE TABLET  
02163780 CODEINE CONTIN (SA) PFR **FNQSW**

200MG CONTROLLED RELEASE TABLET  
02163799 CODEINE CONTIN (SA) PFR **FNQSW**

**FENTANYL**

[SEE APPENDIX A](#) FOR SA CRITERIA

12MCG/HR TRANSDERMAL PATCH  
02311925 TEVA-FENTANYL (SA) TEV **FNQSW**

02327112 SANDOZ-FENTANYL (SA) SDZ **FNQSW**

25MCG/HR TRANSDERMAL PATCH  
02282941 TEVA-FENTANYL (SA) TEV **FNQSW**

02327120 SANDOZ-FENTANYL (SA) SDZ **FNQSW**

37MCG/HR TRANSDERMAL PATCH  
02327139 SANDOZ-FENTANYL (SA) SDZ **FNQSW**

50MCG/HR TRANSDERMAL PATCH  
02282968 TEVA-FENTANYL (SA) TEV **FNQSW**

02327147 SANDOZ-FENTANYL (SA) SDZ **FNQSW**

75MCG/HR TRANSDERMAL PATCH  
02282976 TEVA-FENTANYL (SA) TEV **FNQSW**

02327155 SANDOZ-FENTANYL (SA) SDZ **FNQSW**

100MCG/HR TRANSDERMAL PATCH

02282984	TEVA-FENTANYL (SA)	TEV	<b>FNQSW</b>
02327163	SANDOZ-FENTANYL (SA)	SDZ	<b>FNQSW</b>

**HYDROMORPHONE HCL**

**1MG TABLET**

00705438	DILAUDID	PFR	<b>FNQSW</b>
00885444	PMS-HYDROMORPHONE	PMS	<b>FNQSW</b>
02364115	APO-HYDROMORPHONE	APX	<b>FNQSW</b>

**2MG TABLET**

00125083	DILAUDID	PFR	<b>FNQSW</b>
00885436	PMS-HYDROMORPHONE	PMS	<b>FNQSW</b>
02364123	APO-HYDROMORPHONE	APX	<b>FNQSW</b>

**4MG TABLET**

00125121	DILAUDID	PFR	<b>FNQSW</b>
00885401	PMS-HYDROMORPHONE	PMS	<b>FNQSW</b>
02364131	APO-HYDROMORPHONE	APX	<b>FNQSW</b>

**8MG TABLET**

00786543	DILAUDID	PFR	<b>FNQSW</b>
00885428	PMS-HYDROMORPHONE	PMS	<b>FNQSW</b>
02364158	APO-HYDROMORPHONE	APX	<b>FNQSW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA

**3MG CONTROLLED-RELEASE CAPSULE**

02125323	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>
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**4.5MG CONTROLLED-RELEASE CAPSULE**

02359502	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>
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**6MG CONTROLLED-RELEASE CAPSULE**

02125331	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>
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**9MG CONTROLLED-RELEASE CAPSULE**

02359510	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>
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**12MG CONTROLLED-RELEASE CAPSULE**

02125366	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>
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**18MG CONTROLLED-RELEASE CAPSULE**

02243562	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>
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**24MG CONTROLLED-RELEASE CAPSULE**

02125382	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>
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30MG CONTROLLED-RELEASE CAPSULE			
02125390	HYDROMORPH CONTIN (SA)	PFR	<b>FNQSW</b>

1MG/ML ORAL LIQUID			
01916386	PMS-HYDROMORPHONE	PMS	<b>FNQSW</b>

2MG/ML INJECTION SOLUTION (1ML)			
02145901	HYDROMORPHONE	SDZ	<b>NQ</b>

[SEE APPENDIX A](#) FOR SA CRITERIA. **NOTE:** SA NOT REQUIRED FOR NURSING HOME PROGRAM.

10MG/ML INJECTION SOLUTION (1ML, 5ML, AND 50ML)			
02145928	HYDROMORPHONE (SA)	SDZ	<b>FNQSW</b>

20MG/ML INJECTION			
02145936	HYDROMORPHONE (SA)	SDZ	<b>FNQSW</b>

50MG/ML INJECTION SOLUTION (50ML)			
02146126	HYDROMORPHONE HP (SA)	SDZ	<b>FNQSW</b>

**METHADONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

1MG TABLET			
02247698	METADOL (SA)	PAL	<b>FNQSW</b>
02533642	APO-METHADONE (SA)	APX	<b>FNQSW</b>

5MG TABLET			
02247699	METADOL (SA)	PAL	<b>FNQSW</b>
02533650	APO-METHADONE (SA)	APX	<b>FNQSW</b>

10MG TABLET			
02247700	METADOL (SA)	PAL	<b>FNQSW</b>
02533669	APO-METHADONE (SA)	APX	<b>FNQSW</b>

25MG TABLET			
02247701	METADOL (SA)	PAL	<b>FNQSW</b>
02533677	APO-METHADONE (SA)	APX	<b>FNQSW</b>

Tablets Only - For the management of severe chronic or malignant pain as an alternative to other opiates

**⑤METHADONE SOLUTION**

10MG/ML			
02244290	METADOL-D	PAL	<b>FLNQSW</b>
02495872	ODAN-METHADONE	ODN	<b>FLNQSW</b>
02495880	ODAN-METHADONE	ODN	<b>FLNQSW</b>



**MORPHINE**

1MG/ML ORAL SYRUP

00614491 DOLORAL 1 ATL **FNQSW**

5MG TABLET

00594652 STATEX PAL **FNQSW**

02549794 PMS-MORPHINE SULFATE PMS **FNQSW**

02014203 MSIR PFR **FNQSW**

10MG TABLET

00594644 STATEX PAL **FNQSW**

02549808 PMS-MORPHINE SULFATE PMS **FNQSW**

02014211 MSIR PFR **FNQSW**

20MG TABLET

02014238 MSIR PFR **FNQSW**

25MG TABLET

00594636 STATEX PAL **FNQSW**

30MG TABLET

02014254 MSIR PFR **FNQSW**

50MG TABLET

00675962 STATEX PAL **FNQSW**

10MG EXTENDED RELEASE CAPSULE

02019930 M-ESLON ETH **FNQSW**

15MG EXTENDED RELEASE CAPSULE

02177749 M-ESLON ETH **FNQSW**

30MG EXTENDED RELEASE CAPSULE

02019949 M-ESLON ETH **FNQSW**

60MG EXTENDED RELEASE CAPSULE

02019957 M-ESLON ETH **FNQSW**

100MG EXTENDED RELEASE CAPSULE

02019965 M-ESLON ETH **FNQSW**

200MG EXTENDED RELEASE CAPSULE

02177757 M-ESLON ETH **FNQSW**

15MG SUSTAINED RELEASE TABLET			
02015439	MS CONTIN	PFR	<b>FNQSW</b>
02244790	SANDOZ-MORPHINE SR	SDZ	<b>FNQSW</b>
02302764	TEVA-MORPHINE SR	TEV	<b>FNQSW</b>

30MG SUSTAINED RELEASE TABLET			
02014297	MS CONTIN	PFR	<b>FNQSW</b>
02244791	SANDOZ-MORPHINE SR	SDZ	<b>FNQSW</b>
02302772	TEVA-MORPHINE SR	TEV	<b>FNQSW</b>

60MG SUSTAINED RELEASE TABLET			
02014300	MS CONTIN	PFR	<b>FNQSW</b>
02244792	SANDOZ-MORPHINE SR	SDZ	<b>FNQSW</b>
02302780	TEVA-MORPHINE SR	TEV	<b>FNQSW</b>

100MG SUSTAINED RELEASE TABLET			
02014319	MS CONTIN	PFR	<b>FNQSW</b>
02302799	TEVA-MORPHINE SR	TEV	<b>FNQSW</b>
02478889	SANDOZ-MORPHINE SR	SDZ	<b>FNQSW</b>

200MG SUSTAINED RELEASE TABLET			
02014327	MS CONTIN	PFR	<b>FNQSW</b>
02302802	TEVA-MORPHINE SR	TEV	<b>FNQSW</b>
02478897	SANDOZ-MORPHINE SR	SDZ	<b>FNQSW</b>

10MG/ML INJECTION SOLUTION (1ML)			
00392588	MORPHINE SULFATE	SDZ	<b>NQ</b>

15MG/ML INJECTION SOLUTION (1ML)			
00392561	MORPHINE SULFATE	SDZ	<b>NQ</b>

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG/ML INJECTION SOLUTION(5ML AND 10ML)			
00617288	MORPHINE SULFATE (SA)	SDZ	<b>NQ</b>

**OXYCODONE**

5MG TABLET			
00789739	SUPEUDOL	SDZ	<b>FNQSW</b>
02231934	OXY-IR	PFR	<b>FNQSW</b>
02319977	PMS-OXYCODONE	PMS	<b>FNQSW</b>

10MG TABLET			
00443948	SUPEUDOL	SDZ	<b>FNQSW</b>
02240131	OXY-IR	PFR	<b>FNQSW</b>
02319985	PMS-OXYCODONE	PMS	<b>FNQSW</b>

20MG TABLET			
02240132	OXY-IR	PFR	<b>FNQSW</b>
02262983	SUPEUDOL	SDZ	<b>FNQSW</b>
02319993	PMS-OXYCODONE	PMS	<b>FNQSW</b>

**OXYCODONE HCL & ACETAMINOPHEN**

5MG & 325MG TABLET			
00608165	TEVA-OXYCOCET	TEV	<b>FNQSW</b>
02307898	SANDOZ-OXYCODONE ACET	SDZ	<b>FNQSW</b>
02324628	APO-OXYCODONE/ACET	APX	<b>FNQSW</b>

**OXYCODONE HCL & ACETYLSALICYLIC ACID**

5MG & 325MG TAB			
00608157	TEVA-OXYCODAN	TEV	<b>FQSW</b>

**28:08.12 OPIATE PARTIAL AGONISTS**

**⑤BUPRENORPHINE**

100MG/0.5ML SYRINGE			
02483084	SUBLOCADE	ICL	<b>FLNQSW</b>

300MG/1.5ML SYRINGE			
02483092	SUBLOCADE	ICL	<b>FLNQSW</b>

**⑤BUPRENORPHINE & NALOXONE**

2MG/0.5MG TABLET			
02295695	SUBOXONE	ICL	<b>FLNQSW</b>
02424851	PMS-BUPRENORPHINE/NALOXONE	PMS	<b>FLNQSW</b>
02453908	TEVA-BUPRENORPHINE/NALOXONE	TEV	<b>FLNQSW</b>

8MG/2MG TABLET			
02295709	SUBOXONE	ICL	<b>FLNQSW</b>
02424878	PMS-BUPRENORPHINE/NALOXONE	PMS	<b>FLNQSW</b>
02453916	TEVA-BUPRENORPHINE/NALOXONE	TEV	<b>FLNQSW</b>

2MG/0.5MG FILM			
02502313	SUBOXONE	ICL	<b>FLNQSW</b>

4MG/1MG FILM			
02502321	SUBOXONE	ICL	<b>FLNQSW</b>

8MG/2MG FILM			
02502348	SUBOXONE	ICL	<b>FLNQSW</b>

12MG/3MG FILM  
02502356

SUBOXONE

ICL FLNQSW

**28:08.92 MISCELLANEOUS ANALGESICS AND ANTIPIYRETICS**

**ACETAMINOPHEN**

32MG/ML ELIXIR

00999929 ACETAMINOPHEN

**NW**

Note: The Drug Identification Number listed is for billing purposes only.

80MG/ML DROPS

00999719 ACETAMINOPHEN

**W**

Note: The Drug Identification Number listed is for billing purposes only.

325MG TABLET

00999939 ACETAMINOPHEN

**NW**

Note: The Drug Identification Number listed is for billing purposes only.

500MG TABLET

00999949 ACETAMINOPHEN

**NW**

Note: The Drug Identification Number listed is for billing purposes only.

120MG RECTAL SUPPOSITORY

02230434 ACET-120

PEN **W**

325MG RECTAL SUPPOSITORY

02230436 ACET-325

PEN **NW**

650MG RECTAL SUPPOSITORY

02230437 ACET-650

PEN **NW**

**28:10:00 OPIATE ANTAGONISTS**

**NALOXONE HCL**

0.4MG/ML INJECTION SOLUTION

02148706 NALOXONE

SDZ **NQ**

**⑤ NALTREXONE HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA (FOR SUBSTANCE USE HARM REDUCTION DRUG PROGRAM, NO SA IS REQUIRED)

50MG TABLET

02213826	REVIA (SA)	TEV	<b>FLNQSW</b>
02444275	APO-NALTREXONE (SA)	APX	<b>FGLNQSW</b>
02451883	NALTREXONE (SA)	JPC	<b>FGLNQSW</b>

**28:12.04 ANTICONVULSANTS (BARBITURATES)**

**⑤ PHENOBARBITAL**

Eligible for a 90 day supply

15MG TABLET

00178799	PHENOBARB	PEN	<b>FNQSW</b>
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30MG TABLET

00178802	PHENOBARB	PEN	<b>FNQSW</b>
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60MG TABLET

00178810	PHENOBARB	PEN	<b>FNQSW</b>
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100MG TABLET

00178829	PHENOBARB	PEN	<b>FNQSW</b>
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5MG/ML ELIXIR

00645575	PHENOBARB	PEN	<b>FNQSW</b>
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**PRIMIDONE**

125MG TABLET

00399310	PRIMIDONE	AAA	<b>FGQW</b>
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250MG TABLET

00396761	PRIMIDONE	AAA	<b>FGNQSW</b>
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**28:12.08 ANTICONVULSANTS (BENZODIAZEPINES)**

**CLONAZEPAM**

Eligible for a 90 day supply

0.5MG TABLET

00382825	RIVOTRIL	XPI	<b>FNQSW</b>
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02048701	PMS-CLONAZEPAM	PMS	<b>FNQSW</b>
02177889	APO-CLONAZEPAM	APX	<b>FNQSW</b>
02207818	PMS-CLONAZEPAM-R	PMS	<b>FNQSW</b>

1MG TABLET			
02048728	PMS-CLONAZEPAM	PMS	<b>FNQSW</b>

2MG TABLET			
00382841	RIVOTRIL	XPI	<b>FNQSW</b>
02048736	PMS-CLONAZEPAM	PMS	<b>FNQSW</b>
02177897	APO-CLONAZEPAM	APX	<b>FNQSW</b>

**LORAZEPAM**

4MG/ML INJECTION SOLUTION			
02243278	LORAZEPAM	SDZ	<b>NQ</b>

**28:12.12 ANTICONVULSANTS (HYDANTOINS)**

**PHENYTOIN**

50MG TABLET			
00023698	DILANTIN	UJC	<b>FNQSW</b>

30MG CAPSULE			
00022772	DILANTIN	UJC	<b>FNQSW</b>

100MG CAPSULE			
00022780	DILANTIN	UJC	<b>FNQSW</b>
02460912	PHENYTOIN SODIUM	AAA	<b>FGNQSW</b>

25MG/ML ORAL SUSPENSION			
00023450	DILANTIN	UJC	<b>FNQSW</b>
02250896	TARO-PHENYTOIN	TAR	<b>FGNQSW</b>

50MG/ML INJECTION SOLUTION			
00780626	PHENYTOIN SODIUM	SDZ	<b>NQ</b>

**28:12.20 ANTICONVULSANTS (SUCCINIMIDES)**

**ETHOSUXIMIDE**

50MG/ML SYRUP			
00023485	ZARONTIN	ERF	<b>FNQSW</b>

250MG CAPSULE			
00022799	ZARONTIN	ERF	<b>FNQSW</b>
02545772	MAR-ETHOSUXIMIDE	MAR	<b>FGNQSW</b>
02547171	ODAN-ETHOSUXIMIDE	ODN	<b>FGNQSW</b>

**28:12.92 ANTICONVULSANTS (MISCELLANEOUS)**

**BRIVARACETAM**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET			
02452936	BRIVLERA (SA)	UCB	<b>FNQSW</b>
25MG TABLET			
02452944	BRIVLERA (SA)	UCB	<b>FNQSW</b>
50MG TABLET			
02452952	BRIVLERA (SA)	UCB	<b>FNQSW</b>
75MG TABLET			
02452960	BRIVLERA (SA)	UCB	<b>FNQSW</b>
100MG TABLET			
02452979	BRIVLERA (SA)	UCB	<b>FNQSW</b>

**⑤ CARBAMAZEPINE**

100MG CHEWABLE TABLET			
02244403	TARO-CARBAMAZEPINE	TAR	<b>FGQW</b>
200MG CHEWABLE TABLET			
02244404	TARO-CARBAMAZEPINE	TAR	<b>FGQW</b>
200MG TABLET			
00010405	TEGRETOL	NVR	<b>FNQSW</b>
00782718	TEVA-CARBAMAZEPINE	TEV	<b>FGNQSW</b>
02541238	MINT-CARBAMAZEPINE	MNT	<b>FGNQSW</b>
200MG CONTROLLED RELEASE TABLET			
00773611	TEGRETOL CR	NVR	<b>FNQSW</b>
02261839	SANDOZ-CARBAMAZEPINE CR	SDZ	<b>FGNQSW</b>
400MG CONTROLLED RELEASE TABLET			
00755583	TEGRETOL CR	NVR	<b>FNQSW</b>

02261847 SANDOZ-CARBAMAZEPINE CR SDZ FGNQSW

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG/5ML SUSPENSION

02194333 TEGRETOL (SA) NVR FNQSW  
02367394 TARO-CARBAMAZEPINE (SA) TAR FGNQSW

**CLOBAZAM**

Eligible for a 90 day supply

10MG TABLET

02238334 TEVA-CLOBAZAM TEV FGNQSW  
02244638 APO-CLOBAZAM APX FGNQSW

**Ⓢ DIVALPROEX SODIUM**

125MG ENTERIC COATED TABLET

00596418 EPIVAL BGP FNQSW  
02239698 APO-DIVALPROEX APX FGNQSW  
02458926 MYLAN-DIVALPROEX MYL FGNQSW

250MG ENTERIC COATED TABLET

00596426 EPIVAL BGP FNQSW  
02239699 APO-DIVALPROEX APX FGNQSW  
02458934 MYLAN-DIVALPROEX MYL FGNQSW

500MG ENTERIC COATED TABLET

00596434 EPIVAL BGP FNQSW  
02239700 APO-DIVALPROEX APX FGNQSW  
02459019 MYLAN-DIVALPROEX MYL FGNQSW

**ESLICARBAZEPINE ACETATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

200MG TABLET

02426862 APTIOM (SA) SNV FNQSW

400MG TABLET

02426870 APTIOM (SA) SNV FNQSW

600MG TABLET

02426889 APTIOM (SA) SNV FNQSW

800MG TABLET

02426897 APTIOM (SA) SNV FNQSW

**GABAPENTIN**

100MG CAPSULE



02084260	NEURONTIN	UJC	<b>FNQSW</b>
02243446	PMS-GABAPENTIN	PMS	<b>FGNQSW</b>
02244304	APO-GABAPENTIN	APX	<b>FGNQSW</b>
02244513	TEVA-GABAPENTIN	TEV	<b>FGNQSW</b>
02246314	GABAPENTIN	SIV	<b>FGNQSW</b>
02321203	AURO-GABAPENTIN	ARO	<b>FGNQSW</b>
02353245	GABAPENTIN	SNS	<b>FGNQSW</b>
02361469	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>
02391473	MAR-GABAPENTIN	MAR	<b>FGNQSW</b>
02408880	MINT-GABAPENTIN	MNT	<b>FGNQSW</b>
02416840	GABAPENTIN	ACH	<b>FGNQSW</b>
02535246	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>

300MG CAPSULE

02084279	NEURONTIN	UJC	<b>FNQSW</b>
02243447	PMS-GABAPENTIN	PMS	<b>FGNQSW</b>
02244305	APO-GABAPENTIN	APX	<b>FGNQSW</b>
02244514	TEVA-GABAPENTIN	TEV	<b>FGNQSW</b>
02246315	GABAPENTIN	SIV	<b>FGNQSW</b>
02321211	AURO-GABAPENTIN	ARO	<b>FGNQSW</b>
02319063	RAN-GABAPENTIN	RAN	<b>FGNQSW</b>
02353253	GABAPENTIN	SNS	<b>FGNQSW</b>
02361485	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>
02391481	MAR-GABAPENTIN	MAR	<b>FGNQSW</b>
02408899	MINT-GABAPENTIN	MNT	<b>FGNQSW</b>
02416859	GABAPENTIN	ACH	<b>FGNQSW</b>
02535254	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>

400MG CAPSULE

02084287	NEURONTIN	UJC	<b>FNQSW</b>
02243448	PMS-GABAPENTIN	PMS	<b>FGNQSW</b>
02244306	APO-GABAPENTIN	APX	<b>FGNQSW</b>
02244515	TEVA-GABAPENTIN	TEV	<b>FGNQSW</b>
02246316	GABAPENTIN	SIV	<b>FGNQSW</b>
02321238	AURO-GABAPENTIN	ARO	<b>FGNQSW</b>
02353261	GABAPENTIN	SNS	<b>FGNQSW</b>
02361493	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>
02391503	MAR-GABAPENTIN	MAR	<b>FGNQSW</b>
02408902	MINT-GABAPENTIN	MNT	<b>FGNQSW</b>
02416867	GABAPENTIN	ACH	<b>FGNQSW</b>
02535262	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>

600MG TABLET

02239717	NEURONTIN	UJC	<b>FNQSW</b>
02248457	TEVA-GABAPENTIN	TEV	<b>FGNQSW</b>
02293358	APO-GABAPENTIN	APX	<b>FGNQSW</b>

02388200	GABAPENTIN	SIV	<b>FGNQSW</b>
02392526	GABAPENTIN	ACH	<b>FGNQSW</b>
02402289	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>
02410990	GLN-GABAPENTIN	GLM	<b>FGNQSW</b>
02428334	AURO-GABAPENTIN	ARO	<b>FGNQSW</b>
02431289	GABAPENTIN	SNS	<b>FGNQSW</b>
02432072	GABAPENTIN	JPC	<b>FGNQSW</b>

**800MG TABLET**

02239718	NEURONTIN	UJC	<b>FNQSW</b>
02247346	TEVA-GABAPENTIN	TEV	<b>FGNQSW</b>
02293366	APO-GABAPENTIN	APX	<b>FGNQSW</b>
02388219	GABAPENTIN	SIV	<b>FGNQSW</b>
02392534	GABAPENTIN	ACH	<b>FGNQSW</b>
02402297	JAMP-GABAPENTIN	JPC	<b>FGNQSW</b>
02411008	GLN-GABAPENTIN	GLM	<b>FGNQSW</b>
02428342	AURO-GABAPENTIN	ARO	<b>FGNQSW</b>
02431297	GABAPENTIN	SNS	<b>FGNQSW</b>
02432080	GABAPENTIN	JPC	<b>FGNQSW</b>

**LACOSAMIDE**

**50MG TABLET**

02357615	VIMPAT	UCB	<b>FNQSW</b>
02472902	TEVA-LACOSAMIDE	TEV	<b>FGNQSW</b>
02474670	SANDOZ-LACOSAMIDE	SDZ	<b>FGNQSW</b>
02475332	AURO-LACOSAMIDE	ARO	<b>FGNQSW</b>
02478196	PHARMA-LACOSAMIDE	PMS	<b>FGNQSW</b>
02487802	MAR-LACOSAMIDE	MAR	<b>FGNQSW</b>
02488388	JAMP-LACOSAMIDE	JPC	<b>FGNQSW</b>
02489287	ACH-LACOSAMIDE	ACH	<b>FGNQSW</b>
02490544	MINT-LACOSAMIDE	MNT	<b>FGNQSW</b>
02499568	NRA-LACOSAMIDE	NRA	<b>FGNQSW</b>
02512874	LACOSAMIDE	SNS	<b>FGNQSW</b>

**100MG TABLET**

02357623	VIMPAT	UCB	<b>FNQSW</b>
02472910	TEVA-LACOSAMIDE	TEV	<b>FGNQSW</b>
02474689	SANDOZ-LACOSAMIDE	SDZ	<b>FGNQSW</b>
02475340	AURO-LACOSAMIDE	ARO	<b>FGNQSW</b>
02478218	PHARMA-LACOSAMIDE	PMS	<b>FGNQSW</b>
02487810	MAR-LACOSAMIDE	MAR	<b>FGNQSW</b>
02488396	JAMP-LACOSAMIDE	JPC	<b>FGNQSW</b>
02489295	ACH-LACOSAMIDE	ACH	<b>FGNQSW</b>
02490552	MINT-LACOSAMIDE	MNT	<b>FGNQSW</b>
02499576	NRA-LACOSAMIDE	NRA	<b>FGNQSW</b>
02512882	LACOSAMIDE	SNS	<b>FGNQSW</b>

150MG TABLET

02357631	VIMPAT	UCB	FNQSW
02472929	TEVA-LACOSAMIDE	TEV	FGNQSW
02474697	SANDOZ-LACOSAMIDE	SDZ	FGNQSW
02475359	AURO-LACOSAMIDE	ARO	FGNQSW
02478226	PHARMA-LACOSAMIDE	PMS	FGNQSW
02487829	MAR-LACOSAMIDE	MAR	FGNQSW
02488418	JAMP-LACOSAMIDE	JPC	FGNQSW
02489309	ACH-LACOSAMIDE	ACH	FGNQSW
02490560	MINT-LACOSAMIDE	MNT	FGNQSW
02499584	NRA-LACOSAMIDE	NRA	FGNQSW
02512890	LACOSAMIDE	SNS	FGNQSW

200MG TABLET

02357658	VIMPAT	UCB	FNQSW
02472937	TEVA-LACOSAMIDE	TEV	FGNQSW
02474700	SANDOZ-LACOSAMIDE	SDZ	FGNQSW
02475367	AURO-LACOSAMIDE	ARO	FGNQSW
02478234	PHARMA-LACOSAMIDE	PMS	FGNQSW
02487837	MAR-LACOSAMIDE	MAR	FGNQSW
02488426	JAMP-LACOSAMIDE	JPC	FGNQSW
02489317	ACH-LACOSAMIDE	ACH	FGNQSW
02490579	MINT-LACOSAMIDE	MNT	FGNQSW
02499592	NRA-LACOSAMIDE	NRA	FGNQSW
02512904	LACOSAMIDE	SNS	FGNQSW

**LAMOTRIGINE**

25MG TABLET

02142082	LAMICTAL	GSK	FNQSW
02245208	APO-LAMOTRIGINE	APX	FGNQSW
02246897	PMS-LAMOTRIGINE	PMS	FGNQSW
02265494	MYLAN-LAMOTRIGINE	MYL	FGNQSW
02343010	LAMOTRIGINE	SNS	FGNQSW
02381354	AURO-LAMOTRIGINE	ARO	FGNQSW
02428202	LAMOTRIGINE	SIV	FGNQSW
02542730	JAMP-LAMOTRIGINE	JPC	FGNQSW

100MG TABLET

02142104	LAMICTAL	GSK	FNQSW
02245209	APO-LAMOTRIGINE	APX	FGNQSW
02246898	PMS-LAMOTRIGINE	PMS	FGNQSW
02248233	TEVA-LAMOTRIGINE	TEV	FGNQSW
02265508	MYLAN-LAMOTRIGINE	MYL	FGNQSW
02343029	LAMOTRIGINE	SNS	FGNQSW
02381362	AURO-LAMOTRIGINE	ARO	FGNQSW

02428210	LAMOTRIGINE	SIV	<b>FGNQSW</b>
02542749	JAMP-LAMOTRIGINE	JPC	<b>FGNQSW</b>

150MG TABLET

02142112	LAMICTAL	GSK	<b>FNQSW</b>
02245210	APO-LAMOTRIGINE	APX	<b>FGNQSW</b>
02246899	PMS-LAMOTRIGINE	PMS	<b>FGNQSW</b>
02248234	TEVA-LAMOTRIGINE	TEV	<b>FGNQSW</b>
02265516	MYLAN-LAMOTRIGINE	MYL	<b>FGNQSW</b>
02343037	LAMOTRIGINE	SNS	<b>FGNQSW</b>
02381370	AURO-LAMOTRIGINE	ARO	<b>FGNQSW</b>
02428229	LAMOTRIGINE	SIV	<b>FGNQSW</b>
02542757	JAMP-LAMOTRIGINE	JPC	<b>FGNQSW</b>

**LEVETIRACETAM**

250MG TABLET

02247027	KEPPRA	UCB	<b>FNQSW</b>
02274183	TEVA-LEVETIRACETAM	TEV	<b>FGNQSW</b>
02285924	APO-LEVETIRACETAM	APX	<b>FGNQSW</b>
02296101	PMS-LEVETIRACETAM	PMS	<b>FGNQSW</b>
02353342	LEVETIRACETAM	SNS	<b>FGNQSW</b>
02375249	AURO-LEVETIRACETAM	ARO	<b>FGNQSW</b>
02399776	LEVETIRACETAM	ACH	<b>FGNQSW</b>
02403005	JAMP-LEVETIRACETAM	JPC	<b>FGNQSW</b>
02440202	NAT-LEVETIRACETAM	NAT	<b>FGNQSW</b>
02442388	MINT-LEVETIRACETAM	MNT	<b>FGNQSW</b>
02442531	LEVETIRACETAM	SIV	<b>FGNQSW</b>
02461986	SANDOZ-LEVETIRACETAM	SDZ	<b>FGNQSW</b>
02482274	RIVA-LEVETIRACETAM	RIV	<b>FGNQSW</b>
02499193	NRA-LEVETIRACETAM	NRA	<b>FGNQSW</b>
02504553	JAMP-LEVETIRACETAM	JPC	<b>FGNQSW</b>
02524562	M-LEVETIRACETAM	MRA	<b>FGNQSW</b>

500MG TABLET

02247028	KEPPRA	UCB	<b>FNQSW</b>
02274191	TEVA-LEVETIRACETAM	TEV	<b>FGNQSW</b>
02285932	APO-LEVETIRACETAM	APX	<b>FGNQSW</b>
02296128	PMS-LEVETIRACETAM	PMS	<b>FGNQSW</b>
02353350	LEVETIRACETAM	SNS	<b>FGNQSW</b>
02375257	AURO-LEVETIRACETAM	ARO	<b>FGNQSW</b>
02399784	LEVETIRACETAM	ACH	<b>FGNQSW</b>
02403021	JAMP-LEVETIRACETAM	JPC	<b>FGNQSW</b>
02440210	NAT-LEVETIRACETAM	NAT	<b>FGNQSW</b>
02442396	MINT-LEVETIRACETAM	MNT	<b>FGNQSW</b>
02442558	LEVETIRACETAM	SIV	<b>FGNQSW</b>
02461994	SANDOZ-LEVETIRACETAM	SDZ	<b>FGNQSW</b>

02482282	RIVA-LEVERTIRACETAM	RIV	<b>FGNQSW</b>
02499207	NRA-LEVETIRACETAM	NRA	<b>FGNQSW</b>
02504561	JAMP-LEVETIRACETAM	JPC	<b>FGNQSW</b>
02524570	M-LEVETIRACETAM	MRA	<b>FGNQSW</b>

**750MG TABLET**

02247029	KEPPRA	UCB	<b>FNQSW</b>
02274205	TEVA-LEVETIRACETAM	TEV	<b>FGNQSW</b>
02285940	APO-LEVETIRACETAM	APX	<b>FGNQSW</b>
02296136	PMS-LEVETIRACETAM	PMS	<b>FGNQSW</b>
02353369	LEVETIRACETAM	SNS	<b>FGNQSW</b>
02375265	AURO-LEVETIRACETAM	ARO	<b>FGNQSW</b>
02399792	LEVETIRACETAM	ACH	<b>FGNQSW</b>
02403048	JAMP-LEVETIRACETAM	JPC	<b>FGNQSW</b>
02440229	NAT-LEVETIRACETAM	NAT	<b>FGNQSW</b>
02442418	MINT-LEVETIRACETAM	MNT	<b>FGNQSW</b>
02442566	LEVETIRACETAM	SIV	<b>FGNQSW</b>
02462001	SANDOZ-LEVETIRACETAM	SDZ	<b>FGNQSW</b>
02482290	RIVA-LEVETIRACETAM	RIV	<b>FGNQSW</b>
02499215	NRA-LEVETIRACETAM	NRA	<b>FGNQSW</b>
02504588	JAMP-LEVETIRACETAM	JPC	<b>FGNQSW</b>
02524589	M-LEVETIRACETAM	MRA	<b>FGNQSW</b>

**1000MG TABLET**

02462028	SANDOZ-LEVETIRACETAM	SDZ	<b>FGNQSW</b>
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[SEE APPENDIX A](#) FOR SA CRITERIA

**100MG/ML ORAL SOLUTION**

02490447	PDP-LEVETIRACETAM (SA)	PEN	<b>FGNQSW</b>
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**OXCARBAZEPINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**150MG TABLET**

02284294	APO-OXCARBAZEPINE (SA)	APX	<b>FGNQSW</b>
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**300MG TABLET**

02242068	TRILEPTAL (SA)	NVR	<b>FNQSW</b>
02284308	APO-OXCARBAZEPINE (SA)	APX	<b>FGNQSW</b>

**600MG TABLET**

02242069	TRILEPTAL (SA)	NVR	<b>FNQSW</b>
02284316	APO-OXCARBAZEPINE (SA)	APX	<b>FGNQSW</b>

**PERAMPANEL**

[SEE APPENDIX A](#) FOR SA CRITERIA

**2MG TABLET**

02404516	FYCOMPA (SA)	EIS	<b>FNQSW</b>
02522632	TARO-PERAMPANEL (SA)	TAR	<b>FGNQSW</b>
<b>4MG TABLET</b>			
02404524	FYCOMPA (SA)	EIS	<b>FNQSW</b>
02522640	TARO-PERAMPANEL (SA)	TAR	<b>FGNQSW</b>
<b>6MG TABLET</b>			
02404532	FYCOMPA (SA)	EIS	<b>FNQSW</b>
02522659	TARO-PERAMPANEL (SA)	TAR	<b>FGNQSW</b>
<b>8MG TABLET</b>			
02404540	FYCOMPA (SA)	EIS	<b>FNQSW</b>
02522667	TARO-PERAMPANEL (SA)	TAR	<b>FGNQSW</b>
<b>10MG TABLET</b>			
02404559	FYCOMPA (SA)	EIS	<b>FNQSW</b>
02522675	TARO-PERAMPANEL (SA)	TAR	<b>FGNQSW</b>
<b>12MG TABLET</b>			
02404567	FYCOMPA (SA)	EIS	<b>FNQSW</b>
02522683	TARO-PERAMPANEL (SA)	TAR	<b>FGNQSW</b>
<b>PREGABALIN</b>			
<b>25MG CAPSULE</b>			
02268418	LYRICA	UJC	<b>FNQSW</b>
02359596	PMS-PREGABALIN	PMS	<b>FGNQSW</b>
02361159	TEVA-PREGABALIN	TEV	<b>FGNQSW</b>
02390817	SANDOZ-PREGABALIN	SDZ	<b>FGNQSW</b>
02392801	RAN-PREGABALIN	RAN	<b>FGNQSW</b>
02394235	APO-PREGABALIN	APX	<b>FGNQSW</b>
02403692	PREGABALIN	SIV	<b>FGNQSW</b>
02405539	PREGABALIN	SNS	<b>FGNQSW</b>
02417529	MAR-PREGABALIN	MAR	<b>FGNQSW</b>
02423804	MINT-PREGABALIN	MNT	<b>FGNQSW</b>
02433869	AURO-PREGABALIN	ARO	<b>FGNQSW</b>
02435977	JAMP-PREGABALIN	JPC	<b>FGNQSW</b>
02449838	ACH-PREGABALIN	ACH	<b>FGNQSW</b>
02467291	M-PREGABALIN	MRA	<b>FGNQSW</b>
02479117	NRA-PREGABALIN	NRA	<b>FGNQSW</b>
02480727	AG-PREGABALIN	ANG	<b>FGNQSW</b>
02494841	NAT-PREGABALIN	NAT	<b>FGNQSW</b>
<b>50MG CAPSULE</b>			
02268426	LYRICA	UJC	<b>FNQSW</b>
02359618	PMS-PREGABALIN	PMS	<b>FGNQSW</b>

02361175	TEVA-PREGABALIN	TEV	<b>FGNQSW</b>
02390825	SANDOZ-PREGABALIN	SDZ	<b>FGNQSW</b>
02392828	RAN-PREGABALIN	RAN	<b>FGNQSW</b>
02394243	APO-PREGABALIN	APX	<b>FGNQSW</b>
02403706	PREGABALIN	SIV	<b>FGNQSW</b>
02405547	PREGABALIN	SNS	<b>FGNQSW</b>
02417537	MAR-PREGABALIN	MAR	<b>FGNQSW</b>
02423812	MINT-PREGABALIN	MNT	<b>FGNQSW</b>
02433877	AURO-PREGABALIN	ARO	<b>FGNQSW</b>
02435985	JAMP-PREGABALIN	JPC	<b>FGNQSW</b>
02449846	ACH-PREGABALIN	ACH	<b>FGNQSW</b>
02467305	M-PREGABALIN	MRA	<b>FGNQSW</b>
02479125	NRA-PREGABALIN	NRA	<b>FGNQSW</b>
02480735	AG-PREGABALIN	ANG	<b>FGNQSW</b>
02494868	NAT-PREGABALIN	NAT	<b>FGNQSW</b>

75MG CAPSULE

02268434	LYRICA	UJC	<b>FNQSW</b>
02359626	PMS-PREGABALIN	PMS	<b>FGNQSW</b>
02361183	TEVA-PREGABALIN	TEV	<b>FGNQSW</b>
02390833	SANDOZ-PREGABALIN	SDZ	<b>FGNQSW</b>
02392836	RAN-PREGABALIN	RAN	<b>FGNQSW</b>
02394251	APO-PREGABALIN	APX	<b>FGNQSW</b>
02403714	PREGABALIN	SIV	<b>FGNQSW</b>
02405555	PREGABALIN	SNS	<b>FGNQSW</b>
02417545	MAR-PREGABALIN	MAR	<b>FGNQSW</b>
02424185	MINT-PREGABALIN	MNT	<b>FGNQSW</b>
02433885	AURO-PREGABALIN	ARO	<b>FGNQSW</b>
02435993	JAMP-PREGABALIN	JPC	<b>FGNQSW</b>
02449854	ACH-PREGABALIN	ACH	<b>FGNQSW</b>
02467313	M-PREGABALIN	MRA	<b>FGNQSW</b>
02479133	NRA-PREGABALIN	NRA	<b>FGNQSW</b>
02480743	AG-PREGABALIN	ANG	<b>FGNQSW</b>
02494876	NAT-PREGABALIN	NAT	<b>FGNQSW</b>

150MG CAPSULE

02268450	LYRICA	UJC	<b>FNQSW</b>
02359634	PMS-PREGABALIN	PMS	<b>FGNQSW</b>
02361205	TEVA-PREGABALIN	TEV	<b>FGNQSW</b>
02390841	SANDOZ-PREGABALIN	SDZ	<b>FGNQSW</b>
02392844	RAN-PREGABALIN	RAN	<b>FGNQSW</b>
02394278	APO-PREGABALIN	APX	<b>FGNQSW</b>
02403722	PREGABALIN	SIV	<b>FGNQSW</b>
02405563	PREGABALIN	SNS	<b>FGNQSW</b>
02417561	MAR-PREGABALIN	MAR	<b>FGNQSW</b>
02424207	MINT-PREGABALIN	MNT	<b>FGNQSW</b>

02433907	AURO-PREGABALIN	ARO	<b>FGNQSW</b>
02436000	JAMP-PREGABALIN	JPC	<b>FGNQSW</b>
02449870	ACH-PREGABALIN	ACH	<b>FGNQSW</b>
02467321	M-PREGABALIN	MRA	<b>FGNQSW</b>
02479168	NRA-PREGABALIN	NRA	<b>FGNQSW</b>
02480751	AG-PREGABALIN	ANG	<b>FGNQSW</b>
02494884	NAT-PREGABALIN	NAT	<b>FGNQSW</b>

**225MG CAPSULE**

02268477	LYRICA	UJC	<b>FNQSW</b>
02361221	TEVA-PREGABALIN	TEV	<b>FGNQSW</b>
02392852	RAN-PREGABALIN	RAN	<b>FGNQSW</b>
02394286	APO-PREGABALIN	APX	<b>FGNQSW</b>
02398079	PMS-PREGABALIN	PMS	<b>FGNQSW</b>
02449897	ACH-PREGABALIN	ACH	<b>FGNQSW</b>
02494892	NAT-PREGABALIN	NAT	<b>FGNQSW</b>

**300MG CAPSULE**

02268485	LYRICA	UJC	<b>FNQSW</b>
02359642	PMS-PREGABALIN	PMS	<b>FGNQSW</b>
02361248	TEVA-PREGABALIN	TEV	<b>FGNQSW</b>
02390868	SANDOZ-PREGABALIN	SDZ	<b>FGNQSW</b>
02392860	RAN-PREGABALIN	RAN	<b>FGNQSW</b>
02394294	APO-PREGABALIN	APX	<b>FGNQSW</b>
02403730	PREGABALIN	SIV	<b>FGNQSW</b>
02405598	PREGABALIN	SNS	<b>FGNQSW</b>
02436019	JAMP-PREGABALIN	JPC	<b>FGNQSW</b>
02449900	ACH-PREGABALIN	ACH	<b>FGNQSW</b>
02480778	AG-PREGABALIN	ANG	<b>FGNQSW</b>
02494906	NAT-PREGABALIN	NAT	<b>FGNQSW</b>
02541963	M-PREGABALIN	MRA	<b>FGNQSW</b>

**RUFINAMIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**100MG TABLET**

02369613	BANZEL (SA)	EIS	<b>FNQSW</b>
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**200MG TABLET**

02369621	BANZEL (SA)	EIS	<b>FNQSW</b>
02545985	AURO-RUFINAMIDE (SA)	ARO	<b>FGNQSW</b>

**400MG TABLET**

02369648	BANZEL (SA)	EIS	<b>FNQSW</b>
02545993	AURO-RUFINAMIDE (SA)	ARO	<b>FGNQSW</b>



**STIRIPENTOL**

[SEE APPENDIX A](#) FOR SA CRITERIA  
250MG CAPSULE

02398958          DIACOMIT (SA)          BOX   **NMQW**

250MG POWDER FOR SUSPENSION

02398974          DIACOMIT (SA)          BOX   **NMQW**

500MG CAPSULE

02398966          DIACOMIT (SA)          BOX   **NMQW**

**TOPIRAMATE**

25MG TABLET

02230893	TOPAMAX	JAN	<b>FNQSW</b>
02248860	TEVA-TOPIRAMATE	TEV	<b>FGNQSW</b>
02262991	PMS-TOPIRAMATE	PMS	<b>FGNQSW</b>
02263351	MYLAN-TOPIRAMATE	MYL	<b>FGNQSW</b>
02279614	APO-TOPIRAMATE	APX	<b>FGNQSW</b>
02287765	GLN-TOPIRAMATE	GLM	<b>FGNQSW</b>
02315645	MINT-TOPIRAMATE	MNT	<b>FGNQSW</b>
02345250	JAMP-TOPIRAMATE	JPC	<b>FGNQSW</b>
02345803	AURO-TOPIRAMATE	ARO	<b>FGNQSW</b>
02356856	TOPIRAMATE	SNS	<b>FGNQSW</b>
02389460	TOPIRAMATE	SIV	<b>FGNQSW</b>
02395738	TOPIRAMATE	ACH	<b>FGNQSW</b>
02431807	SANDOZ-TOPIRAMATE	SDZ	<b>FGNQSW</b>
02432099	MAR-TOPIRAMATE	MAR	<b>FGNQSW</b>
02435608	JAMP-TOPIRAMATE	JPC	<b>FGNQSW</b>
02475936	AG-TOPIRAMATE	ANG	<b>FGNQSW</b>

100MG TABLET

02230894	TOPAMAX	JAN	<b>FNQSW</b>
02248861	TEVA-TOPIRAMATE	TEV	<b>FGNQSW</b>
02263009	PMS-TOPIRAMATE	PMS	<b>FGNQSW</b>
02263378	MYLAN-TOPIRAMATE	MYL	<b>FGNQSW</b>
02279630	APO-TOPIRAMATE	APX	<b>FGNQSW</b>
02287773	GLN-TOPIRAMATE	GLM	<b>FGNQSW</b>
02315653	MINT-TOPIRAMATE	MNT	<b>FGNQSW</b>
02345269	JAMP-TOPIRAMATE	JPC	<b>FGNQSW</b>
02345838	AURO-TOPIRAMATE	ARO	<b>FGNQSW</b>
02356864	TOPIRAMATE	SNS	<b>FGNQSW</b>
02389487	TOPIRAMATE	SIV	<b>FGNQSW</b>
02395746	TOPIRAMATE	ACH	<b>FGNQSW</b>
02431815	SANDOZ-TOPIRAMATE	SDZ	<b>FNGQSW</b>

02432102	MAR-TOPIRAMATE	MAR	<b>FGNQSW</b>
02435616	JAMP-TOPIRAMATE	JPC	<b>FGNQSW</b>
02475944	AG-TOPIRAMATE	ANG	<b>FGNQSW</b>

**200MG TABLET**

02230896	TOPAMAX	JAN	<b>FNQSW</b>
02248862	TEVA-TOPIRAMATE	TEV	<b>FGNQSW</b>
02263017	PMS-TOPIRAMATE	PMS	<b>FGNQSW</b>
02263386	MYLAN-TOPIRAMATE	MYL	<b>FGNQSW</b>
02279649	APO-TOPIRAMATE	APX	<b>FGNQSW</b>
02287781	GLN-TOPIRAMATE	GLM	<b>FGNQSW</b>
02315661	MINT-TOPIRAMATE	MNT	<b>FGNQSW</b>
02345277	JAMP-TOPIRAMATE	JPC	<b>FGNQSW</b>
02345846	AURO-TOPIRAMATE	ARO	<b>FGNQSW</b>
02356872	TOPIRAMATE	SNS	<b>FGNQSW</b>
02395754	TOPIRAMATE	ACH	<b>FGNQSW</b>
02431823	SANDOZ-TOPIRAMATE	SDZ	<b>FGNQSW</b>
02432110	MAR-TOPIRAMATE	MAR	<b>FGNQSW</b>
02435624	JAMP-TOPIRAMATE	JPC	<b>FGNQSW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA

**15MG SPRINKLE CAPSULE**

02239907	TOPAMAX (SA)	JAN	<b>FQW</b>
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**25MG SPRINKLE CAPSULE**

02239908	TOPAMAX (SA)	JAN	<b>FQW</b>
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**⑤ VALPROATE SODIUM**

**50MG/ML SYRUP**

00443832	DEPAKENE	BGP	<b>FNQSW</b>
02236807	PMS-VALPROIC	PMS	<b>FGNQSW</b>
02238370	APO-VALPROIC ACID	APX	<b>FGNQSW</b>
02532441	JAMP-VALPROIC ACID	JPC	<b>FGNQSW</b>

**⑤ VALPROIC ACID**

**250MG CAPSULE**

02230768	PMS-VALPROIC	PMS	<b>FGNQSW</b>
02238048	APO-VALPROIC	APX	<b>FGNQSW</b>

**500MG ENTERIC COATED CAPSULE**

02229628	PMS-VALPROIC	PMS	<b>FGNQSW</b>
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**VIGABATRIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

**500MG TABLET**

02065819	SABRIL (SA)	LUD	<b>FNQSW</b>
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## **28:16.04 PSYCHOTHERAPEUTIC AGENTS (ANTIDEPRESSANTS)**

### **⑤AMITRIPTYLINE**

#### 10MG TABLET

00335053	ELAVIL	AAA	<b>FNQSW</b>
00654523	PMS-AMITRIPTYLINE	PMS	<b>FGNQSW</b>
02326043	TEVA-AMITRIPTYLINE	TEV	<b>FGNQSW</b>
02403137	APO-AMITRIPTYLINE	APX	<b>FGNQSW</b>
02429861	MAR-AMITRIPTYLINE	MAR	<b>FGNQSW</b>
02457679	JAMP-AMITRIPTYLINE	JPC	<b>FGNQSW</b>

#### 25MG TABLET

00335061	ELAVIL	AAA	<b>FNQSW</b>
00654515	PMS-AMITRIPTYLINE	PMS	<b>FGNQSW</b>
02326051	TEVA-AMITRIPTYLINE	TEV	<b>FGNQSW</b>
02403145	APO-AMITRIPTYLINE	APX	<b>FGNQSW</b>
02429888	MAR-AMITRIPTYLINE	MAR	<b>FGNQSW</b>
02457695	JAMP-AMITRIPTYLINE	JPC	<b>FGNQSW</b>

#### 50MG TABLET

00335088	ELAVIL	AAA	<b>FNQSW</b>
00654507	PMS-AMITRIPTYLINE	PMS	<b>FGNQSW</b>
02326078	TEVA-AMITRIPTYLINE	TEV	<b>FGNQSW</b>
02403153	APO-AMITRIPTYLINE	APX	<b>FGNQSW</b>
02429896	MAR-AMITRIPTYLINE	MAR	<b>FGNQSW</b>
02457687	JAMP-AMITRIPTYLINE	JPC	<b>FGNQSW</b>

#### 75MG TABLET

00754129	ELAVIL	AAA	<b>FNQSW</b>
02403161	APO-AMITRIPTYLINE	APX	<b>FGNQSW</b>
02429918	MAR-AMITRIPTYLINE	MAR	<b>FGNQSW</b>
02457660	JAMP-AMITRIPTYLINE	JPC	<b>FGNQSW</b>

### **⑤BUPROPION HCL**

#### 100MG TABLET

02275074	ODAN-BUPROPION SR	ODN	<b>FGNQSW</b>
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#### 150MG TABLET

02275082	ODAN-BUPROPION SR	ODN	<b>FGNQSW</b>
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#### 150MG EXTENDED RELEASE TABLET

02275090	WELLBUTRIN XL	VAL	<b>FNQSW</b>
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02439654	TEVA-BUPROPION XL	TEV	<b>FGNQSW</b>
02475804	TARO-BUPROPION XL	SUN	<b>FGNQSW</b>
300MG EXTENDED RELEASE TABLET			
02275104	WELLBUTRIN XL	VAL	<b>FNQSW</b>
02439662	TEVA-BUPROPION XL	TEV	<b>FGNQSW</b>
02475812	TARO-BUPROPION XL	SUN	<b>FGNQSW</b>

**⑤CITALOPRAM**

10MG TABLET

02270609	PMS-CITALOPRAM	PMS	<b>FGNQSW</b>
02312336	TEVA-CITALOPRAM	TEV	<b>FGNQSW</b>
02371871	MAR-CITALOPRAM	MAR	<b>FGNQSW</b>
02387948	CITALOPRAM	SIV	<b>FGNQSW</b>
02409003	NATCO-CITALOPRAM	NAT	<b>FGNQSW</b>
02429691	MINT-CITALOPRAM	MNT	<b>FGNQSW</b>
02430517	CITALOPRAM	JPC	<b>FGNQSW</b>
02443880	NATCO-CITALOPRAM	NAT	<b>FGNQSW</b>
02445719	CITALOPRAM	SNS	<b>FGNQSW</b>
02532123	M-CITALOPRAM	MRA	<b>FGNQSW</b>

20MG TABLET

02239607	CELEXA	LUD	<b>FNQSW</b>
02246056	APO-CITALOPRAM	APX	<b>FGNQSW</b>
02248010	PMS-CITALOPRAM	PMS	<b>FGNQSW</b>
02275562	AURO-CITALOPRAM	ARO	<b>FGNQSW</b>
02293218	TEVA-CITALOPRAM	TEV	<b>FGNQSW</b>
02353660	CITALOPRAM	SNS	<b>FGNQSW</b>
02371898	MAR-CITALOPRAM	MAR	<b>FGNQSW</b>
02387956	CITALOPRAM	SIV	<b>FGNQSW</b>
02409011	NAT-CITALOPRAM	NAT	<b>FGNQSW</b>
02429705	MINT-CITALOPRAM	MNT	<b>FGNQSW</b>
02430541	CITALOPRAM	JPC	<b>FGNQSW</b>
02443899	NATCO-CITALOPRAM	NAT	<b>FGNQSW</b>
02467836	M-CITALOPRAM	MRA	<b>FGNQSW</b>

40MG TABLET

02239608	CELEXA	LUD	<b>FNQSW</b>
02246057	APO-CITALOPRAM	APX	<b>FGNQSW</b>
02248011	PMS-CITALOPRAM	PMS	<b>FGNQSW</b>
02275570	AURO-CITALOPRAM	ARO	<b>FGNQSW</b>
02293226	TEVA-CITALOPRAM	TEV	<b>FGNQSW</b>
02353679	CITALOPRAM	SNS	<b>FGNQSW</b>
02371901	MAR-CITALOPRAM	MAR	<b>FGNQSW</b>
02387964	CITALOPRAM	SIV	<b>FGNQSW</b>
02409038	NAT-CITALOPRAM	NAT	<b>FGNQSW</b>

02429713	MINT-CITALOPRAM	MNT	<b>FGNQSW</b>
02430568	CITALOPRAM	JPC	<b>FGNQSW</b>
02467844	M-CITALOPRAM	MRA	<b>FGNQSW</b>

**⑤ CLOMIPRAMINE HCL**

10MG TABLET			
00330566	ANAFRANIL	AAA	<b>FNQSW</b>

25MG TABLET & CAPSULE			
00324019	ANAFRANIL	APX	<b>FNQSW</b>
02497506	TARO-CLOMIPRAMINE	TAR	<b>FGNQSW</b>

50MG TABLET & CAPSULE			
00402591	ANAFRANIL	APX	<b>FNQSW</b>
02497514	TARO-CLOMIPRAMINE	TAR	<b>FGNQSW</b>

**⑤ DESIPRAMINE**

10MG TABLET			
02216248	DESIPRAMINE	AAA	<b>FGNQSW</b>

25MG TABLET			
02216256	DESIPRAMINE	AAA	<b>FGNQSW</b>

50MG TABLET			
02216264	DESIPRAMINE	AAA	<b>FGNQSW</b>

75MG TABLET			
02216272	DESIPRAMINE	AAA	<b>FGNQSW</b>

100MG TABLET			
02216280	DESIPRAMINE	AAA	<b>FGNQSW</b>

**⑤ DOXEPIN HCL**

10MG CAPSULE			
00024325	SINEQUAN	AAA	<b>FNQSW</b>

25MG CAPSULE			
00024333	SINEQUAN	AAA	<b>FNQSW</b>

50MG CAPSULE			
00024341	SINEQUAN	AAA	<b>FNQSW</b>

75MG CAPSULE			
00400750	SINEQUAN	AAA	<b>FNQSW</b>

100MG CAPSULE

00326925

SINEQUAN

AAA FNQSW

**⑤DULOXETINE HYDROCHLORIDE**

30MG DELAYED RELEASE CAPSULE

02301482	CYMBALTA	LIL	FNQSW
02429446	PMS-DULOXETINE	PMS	FGNQSW
02436647	AURO-DULOXETINE	ARO	FGNQSW
02438984	MINT-DULOXETINE	MNT	FGNQSW
02439948	SANDOX-DULOXETINE	SDZ	FGNQSW
02440423	APO-DULOXETINE	APX	FGNQSW
02446081	MAR-DULOXETINE	MAR	FGNQSW
02451913	JAMP-DULOXETINE	JPC	FGNQSW
02453630	DULOXETINE	SIV	FGNQSW
02456753	TEVA-DULOXETINE	TEV	FGNQSW
02473208	M-DULOXETINE	MRA	FGNQSW
02475308	AG-DULOXETINE	ANG	FGNQSW
02482126	NRA-DULOXETINE	NRA	FGNQSW
02490889	DULOXETINE	SNS	FGNQSW

60MG DELAYED RELEASE CAPSULE

02301490	CYMBALTA	LIL	FNQSW
02429454	PMS-DULOXETINE	PMS	FGNQSW
02436655	AURO-DULOXETINE	ARO	FGNQSW
02438992	MINT-DULOXETINE	MNT	FGNQSW
02439956	SANDOZ-DULOXETINE	SDZ	FGNQSW
02440431	APO-DULOXETINE	APX	FGNQSW
02446103	MAR-DULOXETINE	MAR	FGNQSW
02451921	JAMP-DULOXETINE	JPC	FGNQSW
02453649	DULOXETINE	SIV	FGNQSW
02456761	TEVA-DULOXETINE	TEV	FGNQSW
02473216	M-DULOXETINE	MRA	FGNQSW
02475316	AG-DULOXETINE	ANG	FGNQSW
02482134	NRA-DULOXETINE	NRA	FGNQSW
02490897	DULOXETINE	SNS	FGNQSW

**⑤ESCITALOPRAM**

10MG TABLET

02263238	CIPRALEX	LUD	FNQSW
02295016	APO-ESCITALOPRAM	APX	FGNQSW
02318180	TEVA-ESCITALOPRAM	TEV	FGNQSW
02364077	SANDOZ-ESCITALOPRAM	SDZ	FGNQSW
02385481	RAN-ESCITALOPRAM	RAN	FGNQSW
02397358	AURO-ESCITALOPRAM	ARO	FGNQSW
02407418	MINT-ESCITALOPRAM	MNT	FGNQSW
02429039	ESCITALOPRAM	SIV	FGNQSW
02429780	JAMP-ESCITALOPRAM	JPC	FGNQSW

02430118	ESCITALOPRAM	SNS	FGNQSW
02434652	ACH-ESCITALOPRAM	ACH	FGNQSW
02440296	NAT-ESCITALOPRAM	NAT	FGNQSW
02469243	PMS-ESCITALOPRAM	PMS	FGNQSW
02471418	M-ESCITALOPRAM	MRA	FGNQSW
02476851	NRA-ESCITALOPRAM	NRA	FGNQSW
02508893	JAMP-ESCITALOPRAM	JPC	FGNQSW

15MG TABLET

02512653	KYE-ESCITALOPRAM	KYE	FGNQSW
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20MG TABLET

02263254	CIPRALEX	LUD	FNQSW
02295024	APO-ESCITALOPRAM	APX	FGNQSW
02318202	TEVA-ESCITALOPRAM	TEV	FGNQSW
02364085	SANDOZ-ESCITALOPRAM	SDZ	FGNQSW
02385503	RAN-ESCITALOPRAM	RAN	FGNQSW
02397374	AURO-ESCITALOPRAM	ARO	FGNQSW
02407434	MINT-ESCITALOPRAM	MNT	FGNQSW
02429047	ESCITALOPRAM	SIV	FGNQSW
02429799	JAMP-ESCITALOPRAM	JPC	FGNQSW
02430126	ESCITALOPRAM	SNS	FGNQSW
02434660	ACH-ESCITALOPRAM	ACH	FGNQSW
02440318	NAT-ESCITALOPRAM	NAT	FGNQSW
02469251	PMS-ESCITALOPRAM	PMS	FGNQSW
02471426	M-ESCITALOPRAM	MRA	FGNQSW
02476878	NRA-ESCITALOPRAM	NRA	FGNQSW
02508907	JAMP-ESCITALOPRAM	JPC	FGNQSW

⑤ FLUOXETINE HCL

10MG CAPSULE

02018985	PROZAC	LIL	FNQSW
02177579	PMS-FLUOXETINE	PMS	FGNQSW
02216353	APO-FLUOXETINE	APX	FGNQSW
02216582	TEVA-FLUOXETINE	TEV	FGNQSW
02286068	FLUOXETINE	SNS	FGNQSW
02374447	FLUOXETINE	SIV	FGNQSW
02380560	MINT-FLUOXETINE	MNT	FGNQSW
02385627	AURO-FLUOXETINE	ARO	FGNQSW
02393441	FLUOXETINE	ACH	FGNQSW
02401894	JAMP-FLUOXETINE	JPC	FGNQSW
02485052	AG-FLUOXETINE	ANG	FGNQSW
02503875	NRA-FLUOXETINE	NRA	FGNQSW
02529432	M-FLUOXETINE	MRA	FGNQSW

20MG CAPSULE

00636622	PROZAC	LIL	<b>FNQSW</b>
02177587	PMS-FLUOXETINE	PMS	<b>FGNQSW</b>
02216361	APO-FLUOXETINE	APX	<b>FGNQSW</b>
02216590	TEVA-FLUOXETINE	TEV	<b>FGNQSW</b>
02286076	FLUOXETINE	SNS	<b>FGNQSW</b>
02374455	FLUOXETINE	SIV	<b>FGNQSW</b>
02380579	MINT-FLUOXETINE	MNT	<b>FGNQSW</b>
02383241	FLUOXETINE	ACH	<b>FGNQSW</b>
02385635	AURO-FLUOXETINE	ARO	<b>FGNQSW</b>
02386402	JAMP-FLUOXETINE	JPC	<b>FGNQSW</b>
02485060	AG-FLUOXETINE	ANG	<b>FGNQSW</b>
02503883	NRA-FLUOXETINE	NRA	<b>FGNQSW</b>
02529440	M-FLUOXETINE	MRA	<b>FGNQSW</b>

40MG CAPSULE			
02464640	PMS-FLUOXETINE	PMS	<b>FGNQSW</b>

60MG CAPSULE			
02464659	PMS-FLUOXETINE	PMS	<b>FGNQSW</b>

**⑤ FLUOXETINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG/5ML ORAL SOLUTION

02231328	APO-FLUOXETINE (SA)	APX	<b>FGNQSW</b>
02459361	ODAN-FLUOXETINE (SA)	ODN	<b>FGNQSW</b>

**⑤ FLUVOXAMINE MALEATE**

50MG TABLET

01919342	LUVOX	BGP	<b>FNQSW</b>
02231329	APO-FLUVOXAMINE	APX	<b>FGNQSW</b>
02255529	ACT-FLUVOXAMINE	TEV	<b>FGNQSW</b>

100MG TABLET

01919369	LUVOX	BGP	<b>FNQSW</b>
02231330	APO-FLUVOXAMINE	APX	<b>FGNQSW</b>
02255537	ACT-FLUVOXAMINE	TEV	<b>FGNQSW</b>

**⑤ IMIPRAMINE**

10MG TABLET

00360201	IMIPRAMINE	AAA	<b>FGNQSW</b>
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25MG TABLET

00312797	IMIPRAMINE	AAA	<b>FGNQSW</b>
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50MG TABLET

00326852	IMIPRAMINE	AAA	<b>FGNQSW</b>
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75MG TABLET			
00644579	IMIPRAMINE	AAA	<b>FGNQSW</b>

**TRYPTOPHAN**

500MG CAPSULE			
00718149	TRYPTAN	VAL	<b>FNQSW</b>
02248540	APO-TRYPTOPHAN	APX	<b>FGNQSW</b>

500MG TABLET			
02029456	TRYPTAN	VAL	<b>FNQSW</b>
02240333	TEVA-TRYPTOPHAN	TEV	<b>FGNQSW</b>
02248538	APO-TRYPTOPHAN	APX	<b>FGNQSW</b>

1G TABLET			
00654531	TRYPTAN	VAL	<b>FNQSW</b>
02237250	TEVA-TRYPTOPHAN	TEV	<b>FGNQSW</b>
02248539	APO-TRYPTOPHAN	APX	<b>FGNQSW</b>

**⑤MIRTAZAPINE**

15 MG TABLET			
02250594	SANDOZ-MIRTAZAPINE	SDZ	<b>FGNQSW</b>
02256096	MYLAN-MIRTAZAPINE	MYL	<b>FGNQSW</b>
02273942	PMS-MIRTAZAPINE	PMS	<b>FGNQSW</b>
02286610	APO-MIRTAZAPINE	APX	<b>FGNQSW</b>
02411695	AURO-MIRTAZAPINE	ARO	<b>FGNQSW</b>
02496666	MIRTAZAPINE	SIV	<b>FGNQSW</b>
02532689	MIRTAZAPINE	SNS	<b>FGNQSW</b>

30MG TABLET			
02243910	REMERON	MSD	<b>FNQSW</b>
02248762	PMS-MIRTAZAPINE	PMS	<b>FGNQSW</b>
02250608	SANDOZ-MIRTAZAPINE	SDZ	<b>FGNQSW</b>
02256118	MYLAN-MIRTAZAPINE	MYL	<b>FGNQSW</b>
02259354	TEVA-MIRTAZAPINE	TEV	<b>FGNQSW</b>
02286629	APO-MIRTAZAPINE	APX	<b>FGNQSW</b>
02370689	MIRTAZAPINE	SNS	<b>FGNQSW</b>
02411709	AURO-MIRTAZAPINE	ARO	<b>FGNQSW</b>
02496674	MIRTAZAPINE	SIV	<b>FGNQSW</b>

45MG TABLET			
02286637	APO-MIRTAZAPINE	APX	<b>FGNQSW</b>
02411717	MIRTAZAPINE	ARO	<b>FGNQSW</b>
02496682	MIRTAZAPINE	SIV	<b>FGNQSW</b>

15MG ORALLY DISINTEGRATING TABLET

02248542	REMERON RD	MSD	<b>FNQSW</b>
02299801	AURO-MIRTAZAPINE	ARO	<b>FGNQSW</b>
30MG ORALLY DISINTEGRATING TABLET			
02248543	REMERON RD	MSD	<b>FNQSW</b>
02299828	AURO-MIRTAZAPINE	ARO	<b>FGNQSW</b>
45MG ORALLY DISINTEGRATING TABLET			
02248544	REMERON RD	MSD	<b>FNQSW</b>
02299836	AURO-MIRTAZAPINE	ARO	<b>FGNQSW</b>
<b>MOCLOBEMIDE</b>			
100MG TABLET			
02232148	MOCLOBEMIDE	AAA	<b>FGNQSW</b>
150MG TABLET			
00899356	MANERIX	HLR	<b>FNQSW</b>
02232150	MOCLOBEMIDE	AAA	<b>FGNQSW</b>
300MG TABLET			
02166747	MANERIX	HLR	<b>FNQSW</b>
02240456	MOCLOBEMIDE	AAA	<b>FGNQSW</b>
<b>⑤NORTRIPTYLINE</b>			
10MG CAPSULE			
00015229	AVENTYL	AAA	<b>FNQSW</b>
25MG CAPSULE			
00015237	AVENTYL	AAA	<b>FNQSW</b>
<b>⑤PAROXETINE HCL</b>			
20MG TABLET			
01940481	PAXIL	GSK	<b>FNQSW</b>
02240908	APO-PAROXETINE	APX	<b>FGNQSW</b>
02247751	PMS-PAROXETINE	PMS	<b>FGNQSW</b>
02368870	JAMP-PAROXETINE	JPC	<b>FGNQSW</b>
02248557	TEVA-PAROXETINE	TEV	<b>FGNQSW</b>
02282852	PAROXETINE	SNS	<b>FGNQSW</b>
02383284	AURO-PAROXETINE	ARO	<b>FGNQSW</b>
02388235	PAROXETINE	SIV	<b>FGNQSW</b>
02411954	MAR-PAROXETINE	MAR	<b>FGNQSW</b>
02421380	MINT-PAROXETINE	MNT	<b>FGNQSW</b>
02467410	M-PAROXETINE	MRA	<b>FGNQSW</b>
02475545	AG-PAROXETINE	ANG	<b>FGNQSW</b>
02479761	NRA-PAROXETINE	NRA	<b>FGNQSW</b>
02507781	JAMP-PAROXETINE	JPC	<b>FGNQSW</b>

**30MG TABLET**

01940473	PAXIL	GSK	FNQSW
02240909	APO-PAROXETINE	APX	FGNQSW
02247752	PMS-PAROXETINE	PMS	FGNQSW
02248558	TEVA-PAROXETINE	TEV	FGNQSW
02368889	JAMP-PAROXETINE	JPC	FGNQSW
02282860	PAROXETINE	SNS	FGNQSW
02383292	AURO-PAROXETINE	ARO	FGNQSW
02388243	PAROXETINE	SIV	FGNQSW
02411962	MAR-PAROXETINE	MAR	FGNQSW
02421399	MINT-PAROXETINE	MNT	FGNQSW
02467429	M-PAROXETINE	MRA	FGNQSW
02475553	AG-PAROXETINE	ANG	FGNQSW
02479788	NRA-PAROXETINE	NRA	FGNQSW
02507803	JAMP-PAROXETINE	JPC	FGNQSW

**PHENELZINE SULFATE**

**15MG TABLET**

00476552	NARDIL	ERF	FNQSW
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**⑤SERTRALINE HCL**

**25MG CAPSULE**

02132702	ZOLOFT	UJC	FNQSW
02238280	APO-SERTRALINE	APX	FGNQSW
02240485	TEVA-SERTRALINE	TEV	FGNQSW
02244838	PMS-SERTRALINE	PMS	FGNQSW
02245159	SANDOZ-SERTRALINE	SDZ	FGNQSW
02353520	SERTRALINE	SNS	FGNQSW
02386070	SERTRALINE	SIV	FGNQSW
02390906	AURO-SERTRALINE	ARO	FGNQSW
02399415	MAR-SERTRALINE	MAR	FGNQSW
02402378	MINT-SERTRALINE	MNT	FGNQSW
02469626	SERTRALINE	JPC	FGNQSW
02477882	AG-SERTRALINE	ANG	FGNQSW
02488434	NRA-SERTRALINE	NRA	FGNQSW
02530937	M-SERTRALINE	MRA	FGNQSW

**50MG CAPSULE**

01962817	ZOLOFT	UJC	FNQSW
02238281	APO-SERTRALINE	APX	FGNQSW
02240484	TEVA-SERTRALINE	TEV	FGNQSW
02244839	PMS-SERTRALINE	PMS	FGNQSW
02245160	SANDOZ-SERTRALINE	SDZ	FGNQSW
02353539	SERTRALINE	SNS	FGNQSW
02386089	SERTRALINE	SIV	FGNQSW

02390914	AURO-SERTRALINE	ARO	FGNQSW
02399423	MAR-SERTRALINE	MAR	FGNQSW
02402394	MINT-SERTRALINE	MNT	FGNQSW
02469634	SERTRALINE	JPC	FGNQSW
02477890	AG-SERTRALINE	ANG	FGNQSW
02488442	NRA-SERTRALINE	NRA	FGNQSW
02530945	M-SERTRALINE	MRA	FGNQSW

**100MG CAPSULE**

01962779	ZOLOFT	UJC	FNQSW
02238282	APO-SERTRALINE	APX	FGNQSW
02240481	TEVA-SERTRALINE	TEV	FGNQSW
02244840	PMS-SERTRALINE	PMS	FGNQSW
02353547	SERTRALINE	SNS	FGNQSW
02386097	SERTRALINE	SIV	FGNQSW
02390922	AURO-SERTRALINE	ARO	FGNQSW
02399431	MAR-SERTRALINE	MAR	FGNQSW
02402408	MINT-SERTRALINE	MNT	FGNQSW
02469642	SERTRALINE	JPC	FGNQSW
02477904	AG-SERTRALINE	ANG	FGNQSW
02488450	NRA-SERTRALINE	NRA	FGNQSW
02530953	M-SERTRALINE	MRA	FGNQSW

**TRANLYCYPROMINE SULFATE**

**10MG TABLET**

01919598	PARNATE	GSK	FNQSW
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**⑤ TRAZODONE HCL**

**50MG TABLET**

01937227	PMS-TRAZODONE	PMS	FGNQSW
02144263	TEVA-TRAZODONE	TEV	FGNQSW
02147637	APO-TRAZODONE	APX	FGNQSW
02348772	TRAZADONE	SNS	FGNQSW
02442809	JAMP-TRAZADONE	JPC	FGNQSW

**100MG TABLET**

01937235	PMS-TRAZODONE	PMS	FGNQSW
02144271	TEVA-TRAZODONE	TEV	FGNQSW
02147645	APO-TRAZODONE	APX	FGNQSW
02348780	TRAZODONE	SNS	FGNQSW
02442817	JAMP-TRAZADONE	JPC	FGNQSW

**150MG TABLET**

02144298	TEVA-TRAZADONE	TEV	FGNQSW
02147653	APO-TRAZADONE D	APX	FGNQSW
02348799	TRAZADONE	SNS	FGNQSW

02442825 JAMP-TRAZADONE JPC FGNQSW

**⑤ TRIMIPRAMINE**

12.5MG TABLET  
00740799 TRIMIPRAMINE AAA FGNQSW

25MG TABLET  
00740802 TRIMIPRAMINE AAA FGNQSW

50MG TABLET  
00740810 TRIMIPRAMINE AAA FGNQSW

100MG TABLET  
00740829 TRIMIPRAMINE AAA FGNQSW

75MG CAPSULE  
02070987 TRIMIPRAMINE AAA FGNQSW

**⑤ VENLAFAXINE HCL**

37.5MG EXTENDED RELEASE CAPSULE  
02237279 EFFEXOR XR UJC FNQSW  
02275023 TEVA-VENLAFAXINE XR TEV FGNQSW  
02278545 PMS-VENLAFAXINE XR PMS FGNQSW  
02304317 ACT-VENLAFAXINE XR TEV FGNQSW  
02310317 SANDOZ-VENLAFAXINE XR SDZ FGNQSW  
02331683 APO-VENLAFAXINE XR APX FGNQSW  
02354713 VENLAFAXINE XR SNS FGNQSW  
02380072 TARO-VENLAFAXINE XR SUN FGNQSW  
02385929 VENLAFAXINE XR SIV FGNQSW  
02452839 AURO-VENLAFAXINE XR ARO FGNQSW  
02471280 M-VENLAFAXINE XR MRA FGNQSW  
02516535 VENLAFAXINE XR JPC FGNQSW  
02521466 PMSC-VENLAFAXINE XR PMS FGNQSW

75MG EXTENDED RELEASE CAPSULE  
02237280 EFFEXOR XR UJC FNQSW  
02275031 TEVA-VENLAFAXINE XR TEV FGNQSW  
02278553 PMS-VENLAFAXINE XR PMS FGNQSW  
02304325 ACT-VENLAFAXINE XR TEV FGNQSW  
02310325 SANDOZ VENLAFAXINE XR SDZ FGNQSW  
02331691 APO-VENLAFAXINE XR APX FGNQSW  
02354721 VENLAFAXINE XR SNS FGNQSW  
02380080 TARO-VENLAFAXINE XR SUN FGNQSW  
02385937 VENLAFAXINE XR SIV FGNQSW  
02452847 AURO-VENLAFAXINE XR ARO FGNQSW  
02471299 M-VENLAFAXINE XR MRA FGNQSW

02516543	VENLAFAXINE XR	JPC	FGNQSW
02521482	PMSC-VENLAFAXINE XR	PMS	FGNQSW
150MG EXTENDED RELEASE CAPSULE			
02237282	EFFEXOR XR	UJC	FNQSW
02275058	TEVA-VENLAFAXINE XR	TEV	FGNQSW
02278561	PMS-VENLAFAXINE XR	PMS	FGNQSW
02304333	ACT-VENLAFAXINE XR	TEV	FGNQSW
02310333	SANDOZ-VENLAFAXINE XR	SNS	FGNQSW
02331705	APO-VENLAFAXINE XR	APX	FGNQSW
02354748	VENLAFAXINE XR	SNS	FGNQSW
02380099	TARO-VENLAFAXINE XR	SUN	FGNQSW
02385945	VENLAFAXINE XR	SIV	FGNQSW
02452855	AURO-VENLAFAXINE XR	ARO	FGNQSW
02471302	M-VENLAFAXINE XR	MRA	FGNQSW
02516551	VENLAFAXINE XR	JPC	FGNQSW
02521474	PMSC-VENLAFAXINE XR	PMS	FGNQSW

**VORTIOXETINE**

5MG TABLET

02432919	TRINTELLIX	LUD	FNQSW
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10MG TABLET

02432927	TRINTELLIX	LUD	FNQSW
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20MG TABLET

02432943	TRINTELLIX	LUD	FNQSW
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**28:16.08 PSYCHOTHERAPEUTIC AGENTS (ANTIPSYCHOTICS)**

**⑤ ARIPIPRAZOLE**

2MG TABLET

02322374	ABILIFY	OTS	FNQSW
02460025	AURO-ARIPIPRAZOLE	ARO	FGNQSW
02466635	PMS-ARIPIPRAZOLE	PMS	FGNQSW
02471086	APO-ARIPIPRAZOLE	APX	FGNQSW
02473658	SANDOZ-ARIPIPRAZOLE	SDZ	FGNQSW
02483556	MINT-ARIPIPRAZOLE	MNT	FGNQSW
02506688	ARIPIPRAZOLE	SNS	FGNQSW
02534320	ARIPIPRAZOLE	SIV	FGNQSW

5MG TABLET

02322382	ABILIFY	OTS	FNQSW
02460033	AURO-ARIPIPRAZOLE	ARO	FGNQSW

02466643	PMS-ARIPIPRAZOLE	PMS	<b>FGNQSW</b>
02471094	APO-ARIPIPRAZOLE	APX	<b>FGNQSW</b>
02473666	SANDOZ-ARIPIPRAZOLE	SDZ	<b>FGNQSW</b>
02483564	MINT-ARIPIPRAZOLE	MNT	<b>FGNQSW</b>
02506718	ARIPIPRAZOLE	SNS	<b>FGNQSW</b>
02534339	ARIPIPRAZOLE	SIV	<b>FGNQSW</b>

10MG TABLET

02322390	ABILIFY	OTS	<b>FNQSW</b>
02460041	AURO-ARIPIPRAZOLE	ARO	<b>FGNQSW</b>
02466651	PMS-ARIPIPRAZOLE	PMS	<b>FGNQSW</b>
02471108	APO-ARIPIPRAZOLE	APX	<b>FGNQSW</b>
02473674	SANDOZ-ARIPIPRAZOLE	SDZ	<b>FGNQSW</b>
02483572	MINT-ARIPIPRAZOLE	MNT	<b>FGNQSW</b>
02506726	ARIPIPRAZOLE	SNS	<b>FGNQSW</b>
02534347	ARIPIPRAZOLE	SIV	<b>FGNQSW</b>

15MG TABLET

02322404	ABILIFY	OTS	<b>FNQSW</b>
02460068	AURO-ARIPIPRAZOLE	ARO	<b>FGNQSW</b>
02466678	PMS-ARIPIPRAZOLE	PMS	<b>FGNQSW</b>
02471116	APO-ARIPIPRAZOLE	APX	<b>FGNQSW</b>
02473682	SANDOZ-ARIPIPRAZOLE	SDZ	<b>FGNQSW</b>
02483580	MINT-ARIPIPRAZOLE	MNT	<b>FGNQSW</b>
02506734	ARIPIPRAZOLE	SNS	<b>FGNQSW</b>
02534355	ARIPIPRAZOLE	SIV	<b>FGNQSW</b>

20MG TABLET

02322412	ABILIFY	OTS	<b>FNQSW</b>
02460076	AURO-ARIPIPRAZOLE	ARO	<b>FGNQSW</b>
02466686	PMS-ARIPIPRAZOLE	PMS	<b>FGNQSW</b>
02471124	APO-ARIPIPRAZOLE	APX	<b>FGNQSW</b>
02473690	SANDOZ-ARIPIPRAZOLE	SDZ	<b>FGNQSW</b>
02483599	MINT-ARIPIPRAZOLE	MNT	<b>FGNQSW</b>
02506750	ARIPIPRAZOLE	SNS	<b>FGNQSW</b>
02534363	ARIPIPRAZOLE	SIV	<b>FGNQSW</b>

30MG TABLET

02322455	ABILIFY	OTS	<b>FNQSW</b>
02460084	AURO-ARIPIPRAZOLE	ARO	<b>FGNQSW</b>
02466694	PMS-ARIPIPRAZOLE	PMS	<b>FGNQSW</b>
02471132	APO-ARIPIPRAZOLE	APX	<b>FGNQSW</b>
02473704	SANDOZ-ARIPIPRAZOLE	SDZ	<b>FGNQSW</b>
02483602	MINT-ARIPIPRAZOLE	MNT	<b>FGNQSW</b>
02506785	ARIPIPRAZOLE	SNS	<b>FGNQSW</b>
02534371	ARIPIPRAZOLE	SIV	<b>FGNQSW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA (COMMUNITY MENTAL HEALTH DRUG PROGRAM DOES NOT REQUIRE A SA)

300MG INJECTION

02420864 ABILIFY MAINTENA (SA) OTS **BFNQSW**

400MG INJECTION

02420872 ABILIFY MAINTENA (SA) OTS **BFNQSW**

**ASENAPINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG SUBLINGUAL TABLET

02374803 SAPHRIS (SA) LUD **FNQSW**

10MG SUBLINGUAL TABLET

02374811 SAPHRIS (SA) LUD **FNQSW**

**BREXPIRAZOLE**

0.25MG TABLET

02461749 REXULTI OTS **FNQSW**

0.5MG TABLET

02461757 REXULTI OTS **FNQSW**

1MG TABLET

02461765 REXULTI OTS **FNQSW**

2MG TABLET

02461773 REXULTI OTS **FNQSW**

3MG TABLET

02461781 REXULTI OTS **FNQSW**

4MG TABLET

02461803 REXULTI OTS **FNQSW**

**⑤CHLORPROMAZINE**

25MG TABLET

00232823 TEVA-CHLORPROMAZINE TEV **FGNQSW**

50MG TABLET

00232807 TEVA-CHLORPROMAZINE TEV **FGNQSW**

100MG TABLET

00232831 TEVA-CHLORPROMAZINE TEV **FGNQSW**



### ⑤ CLOZAPINE

[SEE APPENDIX A](#) FOR SA CRITERIA

#### 25MG TABLET

00894737	CLOZARIL (SA)	NVR	<b>FNQSW</b>
02247243	GEN-CLOZAPINE (SA)	MYL	<b>FGNQSW</b>
02248034	AA-CLOZAPINE (SA)	AAA	<b>FGNQSW</b>

#### 50MG TABLET

02305003	GEN-CLOZAPINE (SA)	MYL	<b>FGNQSW</b>
02458748	AA-CLOZAPINE (SA)	AAA	<b>FGNQSW</b>

#### 100MG TABLET

00894745	CLOZARIL (SA)	NVR	<b>FNQSW</b>
02247244	GEN-CLOZAPINE (SA)	MYL	<b>FGNQSW</b>
02248035	AA-CLOZAPINE (SA)	AAA	<b>FGNQSW</b>

#### 200MG TABLET

02305011	GEN-CLOZAPINE (SA)	MYL	<b>FGNQSW</b>
02458756	AA-CLOZAPINE (SA)	AAA	<b>FGNQSW</b>

Note: Clozapine is only to be dispensed to patients upon receipt of weekly or bi-weekly hematological test results by the pharmacy.

### FLUPENTHIXOL DECANOATE

#### 20MG/ML DEPOT INJECTION SOLUTION (10ML)

02156032	FLUANXOL DEPOT	LUD	<b>B</b>
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#### 100MG/ML DEPOT INJECTION SOLUTION (2ML)

02156040	FLUANXOL DEPOT	LUD	<b>B</b>
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### ⑤ FLUPENTHIXOL DIHYDROCHLORIDE

#### 0.5MG TABLET

02156008	FLUANXOL	LUD	<b>FNQSW</b>
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#### 3MG TABLET

02156016	FLUANXOL	LUD	<b>FNQSW</b>
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### ⑤ FLUPHENAZINE HCL

#### 1MG TABLET

00405345	FLUPHENAZINE	AAA	<b>FGNQSW</b>
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#### 2MG TABLET

00410632	FLUPHENAZINE	AAA	<b>FGNQSW</b>
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#### 5MG TABLET

00405361 FLUPHENAZINE AAA FGNQSW

**⑤ HALOPERIDOL**

0.5MG TABLET  
00363685 TEVA-HALOPERIDOL TEV FGNQSW

1MG TABLET  
00363677 TEVA-HALOPERIDOL TEV FGNQSW

2MG TABLET  
00363669 TEVA-HALOPERIDOL TEV FGNQSW

5MG TABLET  
00363650 TEVA-HALOPERIDOL TEV FGNQSW

10MG TABLET  
00713449 TEVA-HALOPERIDOL TEV FGNQSW

5MG/ML INJECTION SOLUTION (1ML)  
00808652 HALOPERIDOL SDZ NQ  
02366010 HALOPERIDOL OMG NQ

**HALOPERIDOL DECANOATE**

100MG/ML DEPOT INJECTION SOLUTION (5ML)  
02130300 HALOPERIDOL LA SDZ B

**⑤ LOXAPINE SUCCINATE**

2.5MG TABLET  
02242868 XYLAC PEN FNQSW

10MG TABLET  
02230838 XYLAC PEN FNQSW

25MG TABLET  
02230839 XYLAC PEN FNQSW

**⑤ LURASIDONE**

20MG TABLET  
02422050 LATUDA SNV FNQSW  
02504499 TARO-LURASIDONE TAR FGNQSW  
02505878 PMS-LURASIDONE PMS FGNQSW  
02513986 AURO-LURASIDONE ARO FGNQSW  
02516438 JAMP-LURASIDONE JPC FGNQSW  
02521075 SANDOZ-LURASIDONE SDZ FGNQSW

40MG TABLET

02387751	LATUDA	SNV	<b>FNQSW</b>
02504502	TARO-LURASIDONE	TAR	<b>FGNQSW</b>
02505886	PMS-LURASIDONE	PMS	<b>FGNQSW</b>
02513994	AURO-LURASIDONE	ARO	<b>FGNQSW</b>
02516446	JAMP-LURASIDONE	JPC	<b>FGNQSW</b>
02521091	SANDOZ-LURASIDONE	SDZ	<b>FGNQSW</b>

**60MG TABLET**

02413361	LATUDA	SNV	<b>FNQSW</b>
02504510	TARO-LURASIDONE	TAR	<b>FGNQSW</b>
02505894	PMS-LURASIDONE	PMS	<b>FGNQSW</b>
02514001	AURO-LURASIDONE	ARO	<b>FGNQSW</b>
02516454	JAMP-LURASIDONE	JPC	<b>FGNQSW</b>
02521105	SANDOZ-LURASIDONE	SDZ	<b>FGNQSW</b>

**80MG TABLET**

02387778	LATUDA	SNV	<b>FNQSW</b>
02504529	TARO-LURASIDONE	TAR	<b>FGNQSW</b>
02505908	PMS-LURASIDONE	PMS	<b>FGNQSW</b>
02514028	AURO-LURASIDONE	ARO	<b>FGNQSW</b>
02516462	JAMP-LURASIDONE	JPC	<b>FGNQSW</b>
02521113	SANDOZ-LURASIDONE	SDZ	<b>FGNQSW</b>

**120MG TABLET**

02387786	LATUDA	SNV	<b>FNQSW</b>
02504537	TARO-LURASIDONE	TAR	<b>FGNQSW</b>
02505916	PMS-LURASIDONE	PMS	<b>FGNQSW</b>
02514036	AURO-LURASIDONE	ARO	<b>FGNQSW</b>
02516470	JAMP-LURASIDONE	JPC	<b>FGNQSW</b>
02521121	SANDOZ-LURASIDONE	SDZ	<b>FGNQSW</b>

**⑤ METHOTRIMEPRAZINE**

**2MG TABLET**

02238403	METHOPRAZINE	AAA	<b>FGNQSW</b>
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**5MG TABLET**

02238404	METHOPRAZINE	AAA	<b>FGNQSW</b>
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**25MG TABLET**

02238405	METHOPRAZINE	AAA	<b>FGNQSW</b>
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**50MG TABLET**

02238406	METHOPRAZINE	AAA	<b>FGNQSW</b>
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**25MG/ML AMPUL**

01927698	NOZINAN	XPI	<b>N</b>
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## ⑤ OLANZAPINE

### 2.5MG TABLET

02229250	ZYPREXA	LIL	FNQSW
02276712	TEVA-OLANZAPINE	TEV	FGNQSW
02281791	APO-OLANZAPINE	APX	FGNQSW
02303116	PMS-OLANZAPINE	PMS	FGNQSW
02310341	SANDOZ-OLANZAPINE	SDZ	FGNQSW
02372819	OLANZAPINE	SNS	FGNQSW
02385864	OLANZAPINE	SIV	FGNQSW
02410141	MINT-OLANZAPINE	MNT	FGNQSW
02417243	JAMP-OLANZAPINE	JPC	FGNQSW
02487608	AG-OLANZAPINE	ANG	FGNQSW

### 5MG TABLET

02229269	ZYPREXA	LIL	FNQSW
02276720	TEVA-OLANZAPINE	TEV	FGNQSW
02281805	APO-OLANZAPINE	APX	FGNQSW
02303159	PMS-OLANZAPINE	PMS	FGNQSW
02310368	SANDOZ-OLANZAPINE	SDZ	FGNQSW
02372827	OLANZAPINE	SNS	FGNQSW
02385872	OLANZAPINE	SIV	FGNQSW
02410168	MINT-OLANZAPINE	MNT	FGNQSW
02417251	JAMP-OLANZAPINE	JPC	FGNQSW
02487616	AG-OLANZAPINE	ANG	FGNQSW

### 7.5MG TABLET

02229277	ZYPREXA	LIL	FNQSW
02276739	TEVA-OLANZAPINE	TEV	FGNQSW
02281813	APO-OLANZAPINE	APX	FGNQSW
02303167	PMS-OLANZAPINE	PMS	FGNQSW
02310376	SANDOZ-OLANZAPINE	SDZ	FGNQSW
02372835	OLANZAPINE	SNS	FGNQSW
02385880	OLANZAPINE	SIV	FGNQSW
02410176	MINT-OLANZAPINE	MNT	FGNQSW
02417278	JAMP-OLANZAPINE	JPC	FGNQSW

### 10MG TABLET

02229285	ZYPREXA	LIL	FNQSW
02276747	TEVA-OLANZAPINE	TEV	FGNQSW
02281821	APO-OLANZAPINE	APX	FGNQSW
02303175	PMS-OLANZAPINE	PMS	FGNQSW
02310384	SANDOZ-OLANZAPINE	SDZ	FGNQSW
02372843	OLANZAPINE	SNS	FGNQSW
02385899	OLANZAPINE	SIV	FGNQSW
02410184	MINT-OLANZAPINE	MNT	FGNQSW
02417286	JAMP-OLANZAPINE	JPC	FGNQSW

02487632	AG-OLANZAPINE	ANG	<b>FGNQSW</b>
15MG TABLET			
02238850	ZYPREXA	LIL	<b>FNQSW</b>
02276755	TEVA-OLANZAPINE	TEV	<b>FGNQSW</b>
02281848	APO-OLANZAPINE	APX	<b>FGNQSW</b>
02303183	PMS-OLANZAPINE	PMS	<b>FGNQSW</b>
02310392	SANDOZ-OLANZAPINE	SDZ	<b>FGNQSW</b>
02372851	OLANZAPINE	SNS	<b>FGNQSW</b>
02385902	OLANZAPINE	SIV	<b>FGNQSW</b>
02410192	MINT-OLANZAPINE	MNT	<b>FGNQSW</b>
02417294	JAMP-OLANZAPINE	JPC	<b>FGNQSW</b>
20MG TABLET			
02238851	ZYPREXA	LIL	<b>FNQSW</b>
02333015	APO-OLANZAPINE	APX	<b>FGNQSW</b>
02359707	TEVA-OLANZAPINE	TEV	<b>FGNQSW</b>
02385910	OLANZAPINE	SIV	<b>FGNQSW</b>
02417308	JAMP-OLANZAPINE	JPC	<b>FGNQSW</b>
5MG ORALLY DISINTEGRATING TABLET			
02243086	ZYPREXA ZYDIS	LIL	<b>FNQSW</b>
02303191	PMS-OLANZAPINE ODT	PMS	<b>FGNQSW</b>
02327775	SANDOZ-OLANZAPINE ODT	SDZ	<b>FGNQSW</b>
02343665	OLANZAPINE ODT	SIV	<b>FGNQSW</b>
02352974	OLANZAPINE ODT	SNS	<b>FGNQSW</b>
02360616	APO-OLANZAPINE ODT	APX	<b>FGNQSW</b>
02406624	JAMP-OLANZAPINE ODT	JPC	<b>FGNQSW</b>
02436965	MINT-OLANZAPINE ODT	MNT	<b>FGNQSW</b>
02448726	AURO-OLANZAPINE ODT	ARO	<b>FGNQSW</b>
10MG ORALLY DISINTEGRATING TABLET			
02243087	ZYPREXA ZYDIS	LIL	<b>FNQSW</b>
02303205	PMS-OLANZAPINE ODT	PMS	<b>FGNQSW</b>
02327783	SANDOZ-OLANZAPINE ODT	SDZ	<b>FGNQSW</b>
02343673	OLANZAPINE ODT	SIV	<b>FGNQSW</b>
02352982	OLANZAPINE ODT	SNS	<b>FGNQSW</b>
02360624	APO-OLANZAPINE ODT	APX	<b>FGNQSW</b>
02406632	JAMP-OLANZAPINE ODT	JPC	<b>FGNQSW</b>
02436973	MINT-OLANZAPINE ODT	MNT	<b>FGNQSW</b>
02448734	AURO-OLANZAPINE ODT	ARO	<b>FGNQSW</b>
15MG ORALLY DISINTEGRATING TABLET			
02243088	ZYPREXA ZYDIS	LIL	<b>FNQSW</b>
02303213	PMS-OLANZAPINE ODT	PMS	<b>FGNQSW</b>
02327791	SANDOZ-OLANZAPINE ODT	SDZ	<b>FGNQSW</b>

02343681	OLANZAPINE ODT	SIV	<b>FGNQSW</b>
02352990	OLANZAPINE ODT	SNS	<b>FGNQSW</b>
02360632	APO-OLANZAPINE ODT	APX	<b>FGNQSW</b>
02406640	JAMP-OLANZAPINE ODT	JPC	<b>FGNQSW</b>
02436981	MINT-OLANZAPINE ODT	MNT	<b>FGNQSW</b>
02448742	AURO-OLANZAPINE ODT	ARO	<b>FGNQSW</b>

20MG ORALLY DISINTEGRATING TABLET			
02243089	ZYPREXA ZYDIS	LIL	<b>FNQSW</b>
02327805	SANDOZ-OLANZAPINE ODT	SDZ	<b>FGNQSW</b>
02343703	OLANZAPINE ODT	SIV	<b>FGNQSW</b>
02360640	APO-OLANZAPINE ODT	APX	<b>FGNQSW</b>
02406659	JAMP-OLANZAPINE ODT	JPC	<b>FGNQSW</b>
02448750	AURO-OLANZAPINE ODT	ARO	<b>FGNQSW</b>

**⑤PALIPERIDONE**

[SEE APPENDIX A FOR SA CRITERIA \(COMMUNITY MENTAL HEALTH DRUG PROGRAM](#)

DOES NOT REQUIRE A SA)

50MG/0.5ML INJECTION

02354217	INVEGA SUSTENNA (SA)	JAN	<b>BFNQSW</b>
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75MG/0.75ML INJECTION

02354225	INVEGA SUSTENNA (SA)	JAN	<b>BFNQSW</b>
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100MG/ML INJECTION

02354233	INVEGA SUSTENNA (SA)	JAN	<b>BFNQSW</b>
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150MG/1.5ML INJECTION

02354241	INVEGA SUSTENNA (SA)	JAN	<b>FNQSW</b>
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[SEE APPENDIX A](#) FOR SA CRITERIA (COMMUNITY MENTAL HEALTH DRUG PROGRAM

DOES NOT REQUIRE A SA)

175MG/0.875ML PREFILLED SYRINGE

02455943	INVEGA TRINZA (SA)	JAN	<b>BFNQSW</b>
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263MG/1.315ML PREFILLED SYRINGE

02455986	INVEGA TRINZA (SA)	JAN	<b>BFNQSW</b>
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350MG/1.75ML PREFILLED SYRINGE

02455994	INVEGA TRINZA (SA)	JAN	<b>BFNQSW</b>
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525MG/2.625ML PREFILLED SYRINGE

02456001	INVEGA TRINZA (SA)	JAN	<b>BFNQSW</b>
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**⑤PERICYAZINE**

5MG CAPSULE

01926780	NEULEPTIL	ERF	<b>FNQSW</b>
10MG CAPSULE			
01926772	NEULEPTIL	ERF	<b>FNQSW</b>
10MG/ML ORAL DROPS			
01926756	NEULEPTIL	ERF	<b>FNQSW</b>
<b>⑤PERPHENAZINE</b>			
2MG TABLET			
00335134	PERPHENAZINE	AAA	<b>FGNQSW</b>
4MG TABLET			
00335126	PERPHENAZINE	AAA	<b>FGNQSW</b>
8MG TABLET			
00335118	PERPHENAZINE	AAA	<b>FGNQSW</b>
16MG TABLET			
00335096	PERPHENAZINE	AAA	<b>FGNQSW</b>
<b>⑤PIMOZIDE</b>			
2MG TABLET			
02245432	PIMOZIDE	AAA	<b>FGNQSW</b>
4MG TABLET			
02245433	PIMOZIDE	AAA	<b>FGNQSW</b>
<b>⑤PROCHLORPERAZINE</b>			
5MG TABLET			
00886440	PROCHLORAZINE	AAA	<b>FGNQSW</b>
10MG TABLET			
00886432	PROCHLORAZINE	AAA	<b>FGNQSW</b>
<b>⑤QUETIAPINE</b>			
25MG TABLET			
02236951	SEROQUEL	AZE	<b>FNQSW</b>
02296551	PMS-QUETIAPINE	PMS	<b>FGNQSW</b>
02316080	ACT-QUETIAPINE	TEV	<b>FGNQSW</b>
02317893	QUETIAPINE	SIV	<b>FGNQSW</b>
02330415	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>
02353164	QUETIAPINE	SNS	<b>FGNQSW</b>
02387794	QUETIAPINE	ACH	<b>FGNQSW</b>
02390140	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>
02399822	MAR-QUETIAPINE	MAR	<b>FGNQSW</b>

02390205	AURO-QUETIAPINE	ARO	<b>FGNQSW</b>
02438003	MINT-QUETIAPINE	MNT	<b>FGNQSW</b>
02439158	NAT-QUETIAPINE	NAT	<b>FGNQSW</b>
02475979	AG-QUETIAPINE	ANG	<b>FGNQSW</b>
02486237	NRA-QUETIAPINE	NRA	<b>FGNQSW</b>
02501635	APO-QUETIAPINE FUMARATE	APX	<b>FGNQSW</b>

100MG TABLET

02236952	SEROQUEL	AZE	<b>FNQSW</b>
02296578	PMS-QUETIAPINE	PMS	<b>FGNQSW</b>
02316099	ACT-QUETIAPINE	TEV	<b>FGNQSW</b>
02317907	QUETIAPINE	SIV	<b>FGNQSW</b>
02330423	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>
02353172	QUETIAPINE	SNS	<b>FGNQSW</b>
02387808	QUETIAPINE	ACH	<b>FGNQSW</b>
02390159	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>
02399830	MAR-QUETIAPINE	MAR	<b>FGNQSW</b>
02390213	AURO-QUETIAPINE	ARO	<b>FGNQSW</b>
02438011	MINT-QUETIAPINE	MNT	<b>FGNQSW</b>
02439166	NAT-QUETIAPINE	NAT	<b>FGNQSW</b>
02501643	APO-QUETIAPINE FUMARATE	APX	<b>FGNQSW</b>

150MG TABLET

02439174	NAT-QUETIAPINE	NAT	<b>FGNQSW</b>
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200MG TABLET

02236953	SEROQUEL	AZE	<b>FNQSW</b>
02296594	PMS-QUETIAPINE	PMS	<b>FGNQSW</b>
02316110	ACT-QUETIAPINE	TEV	<b>FGNQSW</b>
02317923	QUETIAPINE	SIV	<b>FGNQSW</b>
02330458	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>
02353199	QUETIAPINE	SNS	<b>FGNQSW</b>
02387824	QUETIAPINE	ACH	<b>FGNQSW</b>
02390167	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>
02399849	MAR-QUETIAPINE	MAR	<b>FGNQSW</b>
02390248	AURO-QUETIAPINE	ARO	<b>FGNQSW</b>
02438046	MINT-QUETIAPINE	MNT	<b>FGNQSW</b>
02439182	NAT-QUETIAPINE	NAT	<b>FGNQSW</b>
02501651	APO-QUETIAPINE FUMARATE	APX	<b>FGNQSW</b>

300MG TABLET

02244107	SEROQUEL	AZE	<b>FNQSW</b>
02296608	PMS-QUETIAPINE	PMS	<b>FGNQSW</b>
02316129	ACT-QUETIAPINE	TEV	<b>FGNQSW</b>
02317931	QUETIAPINE	SIV	<b>FGNQSW</b>
02330466	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>



02353202	QUETIAPINE	SNS	<b>FGNQSW</b>
02387832	QUETIAPINE	ACH	<b>FGNQSW</b>
02390175	JAMP-QUETIAPINE	JPC	<b>FGNQSW</b>
02399857	MAR-QUETIAPINE	MAR	<b>FGNQSW</b>
02390256	AURO-QUETIAPINE	ARO	<b>FGNQSW</b>
02438054	MINT-QUETIAPINE	MNT	<b>FGNQSW</b>
02439190	NAT-QUETIAPINE	NAT	<b>FGNQSW</b>
02501678	APO-QUETIAPINE FUMARATE	APX	<b>FGNQSW</b>

**⑤RISPERIDONE**

**0.25MG TABLET**

02252007	PMS-RISPERIDONE	PMS	<b>FGNQSW</b>
02282119	APO-RISPERIDONE	APX	<b>FGNQSW</b>
02282690	TEVA-RISPERIDONE	TEV	<b>FGNQSW</b>
02303655	SANDOZ-RISPERIDONE	SDZ	<b>FGNQSW</b>
02328305	RAN-RISPERIDONE	RAN	<b>FGNQSW</b>
02356880	RISPERIDONE	SNS	<b>FGNQSW</b>
02359529	JAMP-RISPERIDONE	JPC	<b>FGNQSW</b>
02359790	MINT-RISPERIDONE	MNT	<b>FGNQSW</b>
02371766	MAR-RISPERIDONE	MAR	<b>FGNQSW</b>
02533804	RISPERIDONE	SIV	<b>FGNQSW</b>

**0.5MG TABLET**

02252015	PMS-RISPERIDONE	PMS	<b>FGNQSW</b>
02264188	TEVA-RISPERIDONE	TEV	<b>FGNQSW</b>
02282127	APO-RISPERIDONE	APX	<b>FGNQSW</b>
02303663	SANDOZ-RISPERIDONE	SDZ	<b>FGNQSW</b>
02328313	RAN-RISPERIDONE	RAN	<b>FGNQSW</b>
02356899	RISPERIDONE	SNS	<b>FGNQSW</b>
02359537	JAMP-RISPERIDONE	JPC	<b>FGNQSW</b>
02359804	MINT-RISPERIDONE	MNT	<b>FGNQSW</b>
02371774	MAR-RISPERIDONE	MAR	<b>FGNQSW</b>
02533928	RISPERIDONE	SIV	<b>FGNQSW</b>

**1MG TABLET**

02252023	PMS-RISPERIDONE	PMS	<b>FGNQSW</b>
02264196	TEVA-RISPERIDONE	TEV	<b>FGNQSW</b>
02279800	SANDOZ-RISPERIDONE	SDZ	<b>FGNQSW</b>
02282135	APO-RISPERIDONE	APX	<b>FGNQSW</b>
02328321	RAN-RISPERIDONE	RAN	<b>FGNQSW</b>
02356902	RISPERIDONE	SNS	<b>FGNQSW</b>
02359545	JAMP-RISPERIDONE	JPC	<b>FGNQSW</b>
02359812	MINT-RISPERIDONE	MNT	<b>FGNQSW</b>
02371782	MAR-RISPERIDONE	MAR	<b>FGNQSW</b>
02533936	RISPERIDONE	SIV	<b>FGNQSW</b>

2MG TABLET

02252031	PMS-RISPERIDONE	PMS	<b>FGNQSW</b>
02264218	TEVA-RISPERIDONE	TEV	<b>FGNQSW</b>
02279819	SANDOZ-RISPERIDONE	SDZ	<b>FGNQSW</b>
02282143	APO-RISPERIDONE	APX	<b>FGNQSW</b>
02328348	RAN-RISPERIDONE	RAN	<b>FGNQSW</b>
02356910	RISPERIDONE	SNS	<b>FGNQSW</b>
02359553	JAMP-RISPERIDONE	JPC	<b>FGNQSW</b>
02359820	MINT-RISPERIDONE	MNT	<b>FGNQSW</b>
02371790	MAR-RISPERIDONE	MAR	<b>FGNQSW</b>
02533944	RISPERIDONE	SIV	<b>FGNQSW</b>

3MG TABLET

02252058	PMS-RISPERIDONE	PMS	<b>FGNQSW</b>
02264226	TEVA-RISPERIDONE	TEV	<b>FGNQSW</b>
02279827	SANDOZ RISPERIDONE	SDZ	<b>FGNQSW</b>
02282151	APO-RISPERIDONE	APX	<b>FGNQSW</b>
02328364	RAN-RISPERIDONE	RAN	<b>FGNQSW</b>
02356929	RISPERIDONE	SNS	<b>FGNQSW</b>
02359561	JAMP-RISPERIDONE	JPC	<b>FGNQSW</b>
02359839	MINT-RISPERIDONE	MNT	<b>FGNQSW</b>
02371804	MAR-RISPERIDONE	MAR	<b>FGNQSW</b>
02533952	RISPERIDONE	SIV	<b>FGNQSW</b>

4MG TABLET

02252066	PMS-RISPERIDONE	PMS	<b>FGNQSW</b>
02264234	TEVA-RISPERIDONE	TEV	<b>FGNQSW</b>
02279835	SANDOZ-RISPERIDONE	SDZ	<b>FGNQSW</b>
02282178	APO-RISPERIDONE	APX	<b>FGNQSW</b>
02328372	RAN-RISPERIDONE	RBX	<b>FGNQSW</b>
02356937	RISPERIDONE	SNS	<b>FGNQSW</b>
02359588	JAMP-RISPERIDONE	JPC	<b>FGNQSW</b>
02359847	MINT-RISPERIDONE	MNT	<b>FGNQSW</b>
02371812	MAR-RISPERIDONE	MAR	<b>FGNQSW</b>
02533960	RISPERIDONE	SIV	<b>FGNQSW</b>

1MG/ML ORAL SOLUTION

02236950	RISPERDAL	JAN	<b>FNQSW</b>
02279266	PMS-RISPERIDONE	PMS	<b>FGNQSW</b>
02454319	JAMP-RISPERIDONE	JPC	<b>FGNQSW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA (COMMUNITY MENTAL HEALTH DRUG PROGRAM DOES NOT REQUIRE A SA)

12.5MG PROLONGED RELEASE INJECTION

02298465	RISPERDAL CONSTA (SA)	JAN	<b>BFNQSW</b>
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25MG PROLONGED RELEASE INJECTION  
02255707 RISPERDAL CONSTA (SA) JAN **BFNQSW**

37.5MG PROLONGED RELEASE INJECTION  
02255723 RISPERDAL CONSTA (SA) JAN **BFNQSW**

50MG PROLONGED RELEASE INJECTION  
02255758 RISPERDAL CONSTA (SA) JAN **BFNQSW**

**⑤ TRIFLUOPERAZINE**

1MG TABLET  
00345539 TRIFLUOPERAZINE AAA **FGNQSW**

2MG TABLET  
00312754 TRIFLUOPERAZINE AAA **FGNQSW**

5MG TABLET  
00312746 TRIFLUOPERAZINE AAA **FGNQSW**

10MG TABLET  
00326836 TRIFLUOPERAZINE AAA **FGNQSW**

**⑤ ZIPRASIDONE HYDROCHLORIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG CAPSULE  
02298597 ZELDOX (SA) UJC **FNQSW**  
02449544 AURO-ZIPRASIDONE (SA) ARO **FGNQSW**

40MG CAPSULE  
02298600 ZELDOX (SA) UJC **FNQSW**  
02449552 AURO-ZIPRASIDONE (SA) ARO **FGNQSW**

60MG CAPSULE  
02298619 ZELDOX (SA) UJC **FNQSW**  
02449560 AURO-ZIPRASIDONE (SA) ARO **FGNQSW**

80MG CAPSULE  
02298627 ZELDOX (SA) UJC **FNQSW**  
02449579 AURO-ZIPRASIDONE (SA) ARO **FGNQSW**

**ZUCLOPENTHIXOL DECANOATE**

200MG INJECTION  
02230406 CLOPIXOL LUD **B**

**⑤ ZUCLOPENTHIXOL HCL**

10MG TABLET

02230402	CLOPIXOL	LUD	FNQSW
25MG TABLET			
02230403	CLOPIXOL	LUD	FNQSW

**28:20.00 RESPIRATORY AND CEREBRAL STIMULANTS**

**⑤ DEXTROAMPHETAMINE/AMPHETAMINE**

**5MG CAPSULE**

02248808	ADDERALL XR	SHR	FQW
02439239	TEVA-AMPHETAMINE XR	TEV	FQW
02445492	APO-AMPHETAMINE XR	APX	FQW
02457288	SANDOZ-AMPHETAMINE XR	SDZ	FQW

**10MG CAPSULE**

02248809	ADDERALL XR	SHR	FQW
02439247	TEVA-AMPHETAMINE XR	TEV	FQW
02445506	APO-AMPHETAMINE XR	APX	FQW
02457296	SANDOZ-AMPHETAMINE XR	SDZ	FQW

**15MG CAPSULE**

02248810	ADDERALL XR	SHR	FQW
02439255	TEVA-AMPHETAMINE XR	TEV	FQW
02445514	APO-AMPHETAMINE XR	APX	FQW
02457318	SANDOZ-AMPHETAMINE XR	SDZ	FQW

**20MG CAPSULE**

02248811	ADDERALL XR	SHR	FQW
02439263	TEVA-AMPHETAMINE XR	TEV	FQW
02445522	APO-AMPHETAMINE XR	APX	FQW
02457326	SANDOZ-AMPHETAMINE XR	SDZ	FQW

**25MG CAPSULE**

02248812	ADDERALL XR	SHR	FQW
02439271	TEVA-AMPHETAMINE XR	TEV	FQW
02445530	APO-AMPHETAMINE XR	APX	FQW
02457334	SANDOZ-AMPHETAMINE XR	SDZ	FQW

**30MG CAPSULE**

02248813	ADDERALL XR	SHR	FQW
02439298	TEVA-AMPHETAMINE XR	TEV	FQW
02445549	APO-AMPHETAMINE XR	APX	FQW
02457342	SANDOZ-AMPHETAMINE XR	SDZ	FQW

**⑤ DEXTROAMPHETAMINE SULFATE**

**5MG TABLET**

01924516	DEXEDRINE	PAL	<b>FQW</b>
02443236	DEXTROAMPHETAMINE	AAA	<b>FQW</b>

**10MG SUSTAINED RELEASE CAPSULE**

01924559	DEXEDRINE	PAL	<b>FQW</b>
02448319	ACT-DEXTROAMPHETAMINE	TEV	<b>FQW</b>

**15MG SUSTAINED RELEASE CAPSULE**

01924567	DEXEDRINE	PAL	<b>FQW</b>
02448327	ACT-DEXTROAMPHETAMINE SR	TEV	<b>FQW</b>

**⑤ LISDEXAMFETAMINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**10MG CAPSULE**

02439603	VYVANSE (SA)	TAK	<b>FQW</b>
02545861	TEVA-LISDEXAMFETAMINE (SA)	TEV	<b>FQW</b>
02546248	SANDOZ-LISDEXAMFETAMINE (SA)	SDZ	<b>FQW</b>

**10MG CHEWABLE TABLET**

02490226	VYVANSE (SA)	TAK	<b>FQW</b>
02533340	TARO-LISDEXAMFETAMINE (SA)	TAR	<b>FQW</b>

**20MG CAPSULE**

02347156	VYVANSE (SA)	TAK	<b>FQW</b>
02545888	TEVA-LISDEXAMFETAMINE (SA)	TEV	<b>FQW</b>
02546256	SANDOZ-LISDEXAMFETAMINE (SA)	SDZ	<b>FQW</b>

**20MG CHEWABLE TABLET**

02490234	VYVANSE (SA)	TAK	<b>FQW</b>
02533359	TARO-LISDEXAMFETAMINE (SA)	TAR	<b>FQW</b>

**30MG CAPSULE**

02322951	VYVANSE (SA)	TAK	<b>FQW</b>
02545896	TEVA-LISDEXAMFETAMINE (SA)	TEV	<b>FQW</b>

**30MG CHEWABLE TABLET**

02490242	VYVANSE (SA)	TAK	<b>FQW</b>
02533367	TARO-LISDEXAMFETAMINE (SA)	TAR	<b>FQW</b>

**40MG CAPSULE**

02347164	VYVANSE (SA)	TAK	<b>FQW</b>
02545918	TEVA-LISDEXAMFETAMINE (SA)	TEV	<b>FQW</b>
02546272	SANDOZ-LISDEXAMFETAMINE (SA)	SDZ	<b>FQW</b>

40MG CHEWABLE TABLET			
02490250	VYVANSE (SA)	TAK	<b>FQW</b>
02533375	TARO-LISDEXAMFETAMINE (SA)	TAR	<b>FQW</b>

50MG CAPSULE			
02322978	VYVANSE (SA)	TAK	<b>FQW</b>
02545926	TEVA-LISDEXAMFETAMINE (SA)	TEV	<b>FQW</b>
02546280	SANDOZ-LISDEXAMFETAMINE (SA)	SDZ	<b>FQW</b>

50MG CHEWABLE TABLET			
02490269	VYVANSE (SA)	TAK	<b>FQW</b>
02533383	TARO-LISDEXAMFETAMINE (SA)	TAR	<b>FQW</b>

60MG CAPSULE			
02347172	VYVANSE (SA)	TAK	<b>FQW</b>
02545934	TEVA-LISDEXAMFETAMINE (SA)	TEV	<b>FQW</b>
02546299	SANDOZ-LISDEXAMFETAMINE (SA)	SDZ	<b>FQW</b>

60MG CHEWABLE TABLET			
02490277	VYVANSE (SA)	TAK	<b>FQW</b>
02533391	TARO-LISDEXAMFETAMINE (SA)	TAR	<b>FQW</b>

**⑤ METHYLPHENIDATE HCL**

5MG TABLET			
02234749	PMS-METHYLPHENIDATE	PMS	<b>FQW</b>
02273950	APO-METHYLPHENIDATE	APX	<b>FQW</b>

10MG TABLET			
00584991	PMS-METHYLPHENIDATE	PMS	<b>FQW</b>
02249324	APO-METHYLPHENIDATE	APX	<b>FQW</b>

20MG TABLET			
00585009	PMS-METHYLPHENIDATE	PMS	<b>FQW</b>
02249332	APO-METHYLPHENIDATE	APX	<b>FQW</b>

20MG SUSTAINED RELEASE TABLET			
02266687	APO-METHYLPHENIDATE SR	APX	<b>FQW</b>

18MG EXTENDED RELEASE TABLET			
02247732	CONCERTA	JAN	<b>FQW</b>
02441934	ACT-METHYLPHENIDATE ER	TEV	<b>FQW</b>
02452731	APO-METHYLPHENIDATE ER	APX	<b>FQW</b>

27MG EXTENDED RELEASE TABLET			
02250241	CONCERTA	JAN	<b>FQW</b>

02441942	ACT-METHYLPHENIDATE ER	TEV	<b>FQW</b>
02452758	APO-METHYLPHENIDATE ER	APX	<b>FQW</b>
36MG EXTENDED RELEASE TABLET			
02247733	CONCERTA	JAN	<b>FQW</b>
02441950	ACT-METHYLPHENIDATE ER	TEV	<b>FQW</b>
02452766	APO-METHYLPHENIDATE ER	APX	<b>FQW</b>
54MG EXTENDED RELEASE TABLET			
02247734	CONCERTA	JAN	<b>FQW</b>
02330377	APO-METHYLPHENIDATE ER	APX	<b>FQW</b>
02441969	ACT-METHYLPHENIDATE ER	TEV	<b>FQW</b>
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
10MG CONTROLLED RELEASE CAPSULE			
02277166	BIPHENTIN (SA)	ELV	<b>FQW</b>
02536943	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>
15MG CONTROLLED RELEASE CAPSULE			
02277131	BIPHENTIN (SA)	ELV	<b>FQW</b>
02536951	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>
20MG CONTROLLED RELEASE CAPSULE			
02277158	BIPHENTIN (SA)	ELV	<b>FQW</b>
02536978	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>
30MG CONTROLLED RELEASE CAPSULE			
02277174	BIPHENTIN (SA)	ELV	<b>FQW</b>
02536986	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>
40MG CONTROLLED RELEASE CAPSULE			
02277182	BIPHENTIN (SA)	ELV	<b>FQW</b>
02536994	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>
50MG CONTROLLED RELEASE CAPSULE			
02277190	BIPHENTIN (SA)	ELV	<b>FQW</b>
02537001	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>
60MG CONTROLLED RELEASE CAPSULE			
02277204	BIPHENTIN (SA)	ELV	<b>FQW</b>
02537028	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>
80MG CONTROLLED RELEASE CAPSULE			
02277212	BIPHENTIN (SA)	ELV	<b>FQW</b>
02537036	PMS-METHYLPHENIDATE CR	PMS	<b>FQW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA  
25MG EXTENDED RELEASE CAPSULE  
02470292 FOQUEST (SA)

ELV FQW

35MG EXTENDED RELEASE CAPSULE  
02470306 FOQUEST (SA)

ELV FQW

45MG EXTENDED RELEASE CAPSULE  
02470314 FOQUEST (SA)

ELV FQW

55MG EXTENDED RELEASE CAPSULE  
02470322 FOQUEST (SA)

ELV FQW

70MG EXTENDED RELEASE CAPSULE  
02470330 FOQUEST (SA)

ELV FQW

85MG EXTENDED RELEASE CAPSULE  
02470349 FOQUEST (SA)

ELV FQW

100MG EXTENDED RELEASE CAPSULE  
02470357 FOQUEST (SA)

ELV FQW

**⑤MODAFINIL**

[SEE APPENDIX A](#) FOR SA CRITERIA  
100MG TABLET

02239665 ALERTEC (SA)  
02285398 APO-MODAFINIL (SA)  
02420260 TEVA-MODAFINIL (SA)  
02430487 AURO-MODAFINIL (SA)  
02432560 MAR-MODAFINIL (SA)  
02503727 JAMP-MODAFINIL (SA)  
02530244 MODAFINIL (SA)

TEV FNQSW  
APX FGNQSW  
TEV FGNQSW  
ARO FGNQSW  
MAR FGNQSW  
JPC FGNQSW  
SNS FGNQSW

**28:24.08 ANXIOLYTICS, SEDATIVES, HYPNOTICS (BENZODIAZEPINES)**

**ALPRAZOLAM**

0.25MG TABLET

00548359 XANAX  
00865397 APO-ALPRAZ  
01913484 TEVA-ALPRAZOLAM

UJC FNQSW  
APX FNQSW  
TEV FNQSW

0.5MG TABLET

00548367 XANAX

UJC FNQSW



00865400	APO-ALPRAZ	APX	<b>FNQSW</b>
01913492	TEVA-ALPRAZOLAM	TEV	<b>FNQSW</b>
<b>BROMAZEPAM</b>			
1.5MG TABLET			
02177153	APO-BROMAZEPAM	APX	<b>FNQSW</b>
3MG TABLET			
02177161	APO-BROMAZEPAM	APX	<b>FNQSW</b>
02230584	TEVA-BROMAZEPAM	TEV	<b>FNQSW</b>
6MG TABLET			
02177188	APO-BROMAZEPAM	APX	<b>FNQSW</b>
02230585	TEVA-BROMAZEPAM	TEV	<b>FNQSW</b>
<b>CHLORDIAZEPOXIDE</b>			
5MG CAPSULE			
00522724	CHLORDIAZEPOXIDE	AAA	<b>FNQSW</b>
10MG CAPSULE			
00522988	CHLORDIAZEPOXIDE	AAA	<b>FNQSW</b>
25MG CAPSULE			
00522996	CHLORDIAZEPOXIDE	AAA	<b>FNQSW</b>
<b>CLORAZEPATE DIPOTASSIUM</b>			
3.75MG CAPSULE			
00860689	CLORAZEPATE	AAA	<b>FNQSW</b>
7.5MG CAPSULE			
00860700	CLORAZEPATE	AAA	<b>FNQSW</b>
15MG CAPSULE			
00860697	CLORAZEPATE	AAA	<b>FNQSW</b>
<b>DIAZEPAM</b>			
2MG TABLET			
00405329	DIAZEPAM	AAA	<b>FNQSW</b>
5MG TABLET			
00013285	VALIUM	SLP	<b>FNQSW</b>
00362158	DIAZEPAM	AAA	<b>FNQSW</b>
10MG TABLET			
00405337	DIAZEPAM	AAA	<b>FNQSW</b>

**FLURAZEPAM**

15MG CAPSULE

00521698 FLURAZEPAM

AAA **FNQSW**

30MG CAPSULE

00521701 FLURAZEPAM

AAA **FNQSW****LORAZEPAM**

0.5MG TABLET

00655740 APO-LORAZEPAM  
00711101 TEVA-LORAZEPAM  
00728187 PMS-LORAZEPAM  
02041413 ATIVANAPX **FNQSW**  
TEV **FNQSW**  
PMS **FNQSW**  
PFI **FNQSW**

1MG TABLET

00637742 TEVA-LORAZEPAM  
00655759 APO-LORAZEPAM  
00728195 PMS-LORAZEPAM  
02041421 ATIVANTEV **FNQSW**  
APX **FNQSW**  
PMS **FNQSW**  
PFI **FNQSW**

2MG TABLET

00637750 TEVA-LORAZEPAM  
00655767 APO-LORAZEPAM  
00728209 PMS-LORAZEPAM  
02041448 ATIVANTEV **FNQSW**  
APX **FNQSW**  
PMS **FNQSW**  
PFI **FNQSW****MIDAZOLAM**

5MG/ML INJECTION SOLUTION (2ML)

02240286 MIDAZOLAM

SDZ **NQ****NITRAZEPAM**

5MG TABLET

00511528 MOGADON

AAA **FNQSW**

10MG TABLET

00511536 MOGADON

AAA **FNQSW****OXAZEPAM**

10MG TABLET

00402680 APO-OXAZEPAM

APX **FNQSW**

15MG TABLET

00402745 APO-OXAZEPAM

APX **FNQSW**

30MG TABLET

00402737 APO-OXAZEPAM

APX **FNQSW**

**TEMAZEPAM**

15MG CAPSULE

00604453

RESTORIL

AAA **FNQSW**

30MG CAPSULE

00604461

RESTORIL

AAA **FNQSW****TRIAZOLAM**

Note: Treatment with Triazolam should usually not exceed 7 to 10 consecutive days. Use for more than 2 to 3 consecutive weeks requires a complete re-evaluation of the patient.

0.25MG TABLET

00808571

TRIAZOLAM

AAA **FW****28:24.92 MISCELLANEOUS ANXIOLYTICS, SEDATIVES, HYPNOTICS****⑤BUSPIRONE**

10MG TABLET

02211076

APO-BUSPIRONE

APX **FGNQSW**

02230942

PMS-BUSPIRONE

PMS **FGNQSW**

02231492

TEVA-BUSPIRONE

TEV **FGNQSW**

02447851

BUSPIRONE

SNS **FGNQSW**

02500213

AURO-BUSPIRONE

ARO **FGNQSW**

02509911

JAMP-BUSPIRONE

JPC **FGNQSW**

02519054

MINT-BUSPIRONE

MNT **FGNQSW****HYDROXYZINE HCL**

10 MG CAPSULE

00646059

HYDROXYZINE

AAA **FGNQSW**

25 MG CAPSULE

00646024

HYDROXYZINE

AAA **FGNQSW**

50 MG CAPSULE

00646016

HYDROXYZINE

AAA **FGNQSW**

2 MG/ML SYRUP

00024694

ATARAX

ERF **FNQSW****ZOPICLONE**

3.75 MG TABLET

02458543

PMS-ZOPICLONE

PMS **FNQW**

5 MG TABLET

02243426	PMS-ZOPICLONE	PMS	<b>FNQW</b>
02245077	APO-ZOPICLONE	APX	<b>FNQW</b>
02246534	TEVA-ZOPICLONE	TEV	<b>FNQW</b>
02267918	RAN-ZOPICLONE	RAN	<b>FNQW</b>
02344122	ZOPICLONE	SNS	<b>FNQW</b>
02385821	ZOPICLONE	SIV	<b>FNQW</b>
02386771	MAR-ZOPICLONE	MAR	<b>FNQW</b>
02391716	MINT-ZOPICLONE	MNT	<b>FNQW</b>
02406969	JAMP-ZOPICLONE	JPC	<b>FNQW</b>
02467941	M-ZOPICLONE	MRA	<b>FNQW</b>
02475839	AG-ZOPICLONE	ANG	<b>FNQW</b>
02477378	NRA-ZOPICLONE	NRA	<b>FNQW</b>

7.5 MG TABLET

01926799	IMOVANE	AVN	<b>FNQW</b>
02218313	APO-ZOPICLONE	APX	<b>FNQW</b>
02240606	PMS-ZOPICLONE	PMS	<b>FNQW</b>
02242481	TEVA-ZOPICLONE	TEV	<b>FNQW</b>
02267926	RAN-ZOPICLONE	RAN	<b>FNQW</b>
02282445	ZOPICLONE	SNS	<b>FNQW</b>
02385848	ZOPICLONE	SIV	<b>FNQW</b>
02386798	MAR-ZOPICLONE	MAR	<b>FNQW</b>
02391724	MINT-ZOPICLONE	MNT	<b>FNQW</b>
02406977	JAMP-ZOPICLONE	JPC	<b>FNQW</b>
02467968	M-ZOPICLONE	MRA	<b>FNQW</b>
02475847	AG-ZOPICLONE	ANG	<b>FNQW</b>
02477386	NRA-ZOPICLONE	NRA	<b>FNQW</b>

**28:28.00 ANTIMANIC AGENTS**

**⑤ LITHIUM CARBONATE**

150MG CAPSULE

00461733	CARBOLITH	VAL	<b>FNQSW</b>
02216132	PMS-LITHIUM CARBONATE	PMS	<b>FGNQSW</b>
02242837	APO-LITHIUM CARBONATE	APX	<b>FGNQSW</b>

150MG CAPSULE

02013231	LITHANE	ERF	<b>FNQSW</b>
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300MG CAPSULE

00236683	CARBOLITH	VAL	<b>FNQSW</b>
02216140	PMS-LITHIUM CARBONATE	PMS	<b>FGNQSW</b>

02242838	APO-LITHIUM CARBONATE	APX	FGNQSW
300MG CAPSULE			
00406775	LITHANE	ERF	FNQSW
600MG CAPSULE			
02011239	CARBOLITH	VAL	FNQSW
02216159	PMS-LITHIUM CARBONATE	PMS	FGNQSW
300MG SUSTAINED RELEASE TABLET			
02266695	LITHMAX	AAA	FNQSW

## **28:32.00 MISCELLANEOUS ANTIMIGRAINE AGENTS**

### **ALMOTRIPTAN**

12.5MG TABLET

02398443	MYLAN-ALMOTRIPTAN	MYL	FGNQSW
02405334	SANDOZ-ALMOTRIPTAN	SDZ	FGNQSW
02405806	APO-ALMOTRIPTAN	APX	FGNQSW
02434849	TEVA-ALMOTRIPTAN	TEV	FGNQSW
02466821	ALMOTRIPTAN	SNS	FGNQSW

Note: Coverage is limited to 6 tablets per 30 day period

### **ATOGEPAANT**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET

02533979	QULIPTA (SA)	ABV	FNQSW
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30MG TABLET

02533987	QULIPTA (SA)	ABV	FNQSW
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### **EPTINEZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG/1.0ML VIAL

02510839	VYEPTI (SA)	LUD	FNQW
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### **FREMANEZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

225MG/1.5ML

02497859	AJOVY PREFILLED SYRINGE (SA)	TEV	FNQSW
02509474	AJOVY AUTOINJECTOR (SA)	TEV	FNQSW

**GALCANEZUMAB**[SEE APPENDIX A](#) FOR SA CRITERIA

120MG/ML PREFILLED PEN

02491087            EMGALITY (SA)

LIL    **FNQSW**

120MG/ML PREFILLED SYRINGE

02491060            EMGALITY (SA)

LIL    **FNQSW****NARATRIPTAN HCL**[SEE APPENDIX A](#) FOR SA CRITERIA

1MG TABLET

02237820            AMERGE (SA)

GSK    **FNQSW**

02314290            TEVA-NARATRIPTAN (SA)

TEV    **FGNQSW**

2.5MG TABLET

02237821            AMERGE (SA)

GSK    **FNQSW**

02314304            TEVA-NARATRIPTAN (SA)

TEV    **FGNQSW**

02322323            SANDOZ-NARATRIPTAN (SA)

SDZ    **FGNQSW**

Note: Coverage is limited to 6 tablets per 30 day period.

**PIZOTYLINE**

1MG TABLET

00511552            SANDOMIGRAN DS

PAL    **FNQSW****RIZATRIPTAN**

5MG TABLET

02393468            APO-RIZATRIPTAN

APX    **FGNQSW**

02429233            JAMP-RIZATRIPTAN IR

JPC    **FGNQSW**

10MG TABLET

02240521            MAXALT

MSD    **FNQSW**

02379678            MAR-RIZATRIPTAN

MAR    **FGNQSW**

02380463            JAMP-RIZATRIPTAN

JPC    **FGNQSW**

02381702            ACT-RIZATRIPTAN

TEV    **FGNQSW**

02393476            APO-RIZATRIPTAN

APX    **FGNQSW**

02429241            JAMP-RIZATRIPTAN IR

JPC    **FGNQSW**

02441144            AURO-RIZATRIPTAN

ARO    **FGNQSW**

02516756            RIZATRIPTAN

SNS    **FGNQSW**

5MG ORALLY DISINTEGRATING TABLET

02240518            MAXALT RPD

MSD    **FNQSW**

02351870            SANDOZ-RIZATRIPTAN ODT

SDZ    **FGNQSW**

02379198            MYLAN-RIZATRIPTAN ODT

MYL    **FGNQSW**

02393360            PMS-RIZATRIPTAN ODT

PMS    **FGNQSW**

02396661            TEVA-RIZATRIPTAN ODT

TEV    **FGNQSW**

02436604            NAT-RIZATRIPTAN ODT

NAT    **FGNQSW**

02442906	RIZATRIPTAN ODT	SNS	FGNQSW
02446111	RIZATRIPTAN ODT	SIV	FGNQSW
02462788	MAR-RIZATRIPTAN ODT	MAR	FGNQSW
02465086	JAMP-RIZATRIPTAN ODT	JPC	FGNQSW

10MG ORALLY DISINTEGRATING TABLET

02240519	MAXALT RPD	MSD	FNQSW
02351889	SANDOZ-RIZATRIPTAN ODT	SDZ	FGNQSW
02379201	MYLAN-RIZATRIPTAN ODT	MYL	FGNQSW
02393379	PMS-RIZATRIPTAN ODT	PMS	FGNQSW
02396688	TEVA-RIZATRIPTAN ODT	TEV	FGNQSW
02436612	NAT-RIZATRIPTAN ODT	NAT	FGNQSW
02442914	RIZATRIPTAN ODT	SNS	FGNQSW
02446138	RIZATRIPTAN ODT	SIV	FGNQSW
02462796	MAR-RIZATRIPTAN ODT	MAR	FGNQSW
02465094	JAMP-RIZATRIPIAN ODT	JPC	FGNQSW
02492490	AG-RIZATRIPTAN ODT	ANG	FGNQSW

Note: Coverage is limited to 6 tablets per 30 day period.

**SUMATRIPTAN**

50MG TABLET

02212153	IMITREX DF	GSK	FNQSW
02256436	PMS-SUMATRIPTAN	PMS	FGNQSW
02268388	APO-SUMATRIPTAN	APX	FGNQSW
02268914	MYLAN-SUMATRIPTAN	MYL	FGNQSW
02286823	TEVA-SUMATRIPTAN DF	TEV	FGNQSW
02286521	SUMATRIPTAN	SNS	FGNQSW
02385570	SUMATRIPTAN DF	SIV	FGNQSW
02546035	SUMATRIPTAN	SIV	FGNQSW

100MG TABLET

02212161	IMITREX DF	GSK	FNQSW
02239367	TEVA-SUMATRIPTAN	TEV	FGNQSW
02256444	PMS-SUMATRIPTAN	PMS	FGNQSW
02268396	APO-SUMATRIPTAN	APX	FGNQSW
02268922	MYLAN-SUMATRIPTAN	MYL	FGNQSW
02286831	TEVA-SUMATRIPTAN DF	TEV	FGNQSW
02286548	SUMATRIPTAN	SNS	FGNQSW
02385589	SUMATRIPTAN DF	SIV	FGNQSW
02546043	SUMATRIPTAN	SIV	FGNQSW

[SEE APPENDIX A](#) FOR SA CRITERIA

6MG/0.5ML INJECTION SOLUTION

02212188	IMITREX (SA)	GSK	FNQSW
02361698	TARO-SUMATRIPTAN (SA)	TAR	FGNQSW

[SEE APPENDIX A](#) FOR SA CRITERIA

**5MG NASAL SPRAY**

02230418	IMITREX (SA)	GSK	FNQSW
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**20MG NASAL SPRAY**

02230420	IMITREX (SA)	GSK	FNQSW
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Note: Coverage is limited to 6 tablets or 6 sprays or 6 syringes per 30 day period.

**ZOLMITRIPTAN**

**2.5MG TABLET**

02238660	ZOMIG	XPI	FNQSW
02313960	TEVA-ZOLMITRIPTAN	TEV	FGNQSW
02362988	SANDOZ-ZOLMITRIPTAN	SDZ	FGNQSW
02380951	APO-ZOLMITRIPTAN	APX	FGNQSW
02399458	MAR-ZOLMITRIPTAN	MAR	FGNQSW
02419521	MINT-ZOLMITRIPTAN	MNT	FGNQSW
02421534	NAT-ZOLMITRIPTAN	NAT	FGNQSW
02421623	JAMP-ZOLMITRIPTAN	JPC	FGNQSW
02442655	ZOLMITRIPTAN	SNS	FGNQSW
02477106	JAMP-ZOLMITRIPTAN	JPC	FGNQSW
02481030	AURO-ZOLMITRIPTAN	ARO	FGNQSW

**2.5MG ORALLY DISINTEGRATING TABLET**

02243045	ZOMIG RAPIMELT	XPI	FNQSW
02342545	TEVA-ZOLMITRIPTAN ODT	TEV	FGNQSW
02362996	SANDOZ-ZOLMITRIPTAN ODT	SDZ	FGNQSW
02428237	JAMP-ZOLMITRIPTAN ODT	JPC	FGNQSW
02442671	ZOLMITRIPTAN ODT	SNS	FGNQSW

Note: Coverage is limited to 6 tablets per 30 day period.

**28:36.00 ANTI PARKINSONIAN AGENTS**

**BROMOCRIPTINE**

**2.5MG TABLET**

02087324	BROMOCRIPTINE	AAA	FGNQSW
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**5MG CAPSULE**

02230454	BROMOCRIPTINE	AAA	FGNQSW
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**CABERGOLINE**

**0.5MG TABLET**

02455897	APO-CABERGOLINE	APX	FGNQSW
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**CARBIDOPA & LEVODOPA & ENTACAPONE**[SEE APPENDIX A](#) FOR SA CRITERIA

12.5/50/200MG TABLET

02305933 STALEVO 50 (SA) SDZ **FNQSW**

18.75/75/200MG TABLET

02337827 STALEVO 75 (SA) SDZ **FNQSW**

25/100/200MG TABLET

02305941 STALEVO 100 (SA) SDZ **FNQSW**

31.25/125/200MG TABLET

02337835 STALEVO 125 (SA) SDZ **FNQSW**

37.5/150/200MG TABLET

02305968 STALEVO 150 (SA) SDZ **FNQSW****ENTACAPONE**

200MG TABLET

02243763 COMTAN SDZ **FNQSW**02375559 TEVA-ENTACAPONE TEV **FGNQSW**02380005 SANDOZ-ENTACAPONE SDZ **FGNQSW**02535939 MINT-ENTACAPONE MNT **FGNQSW****LEVODOPA & CARBIDOPA**

100MG &amp; 10MG TABLET

02195933 APO-LEVOCARB APX **FGNQSW**02244494 TEVA-LEVOCARBIDOPA TEV **FGNQSW**02457954 MINT-LEVOCARB MNT **FGNQSW**02531593 AURO-LEVOCARB ARO **FGNQSW**

100MG &amp; 25MG TABLET

02195941 APO-LEVOCARB APX **FGNQSW**02244495 TEVA-LEVOCARBIDOPA TEV **FGNQSW**02457962 MINT-LEVOCARB MNT **FGNQSW**02531607 AURO-LEVOCARB ARO **FGNQSW**

250MG &amp; 25MG TABLET

02195968 APO-LEVOCARB APX **FGNQSW**02244496 TEVA-LEVOCARBIDOPA TEV **FGNQSW**02457970 MINT-LEVOCARB MNT **FGNQSW**02531615 AURO-LEVOCARB ARO **FGNQSW**

100MG &amp; 25MG CONTROLLED RELEASE TABLET

02272873 AA-LEVOCARB CR AAA **FGNQSW**

200MG & 50MG CONTROLLED RELEASE TABLET  
02245211 AA-LEVOCARB CR

AAA FGNQSW

[SEE APPENDIX A](#) FOR SA CRITERIA  
20MG/ML & 5MG INTESTINAL GEL CASSETTE  
02292165 DUODOPA (SA)

ABV NMQW

**PRAMIPEXOLE DIHYDROCHLORIDE**

0.25MG TABLET

02237145 MIRAPEX  
02292378 APO-PRAMIPEXOLE  
02297302 ACT-PRAMIPEXOLE  
02309122 PRAMIPEXOLE  
02315262 SANDOZ PRAMIPEXOLE  
02367602 PRAMIPEXOLE  
02424061 AURO-PRAMIPEXOLE

BOE FNQSW  
APX FGNQSW  
TEV FGNQSW  
SIV FGNQSW  
SDZ FGNQSW  
SNS FGNQSW  
ARO FGNQSW

1MG TABLET

02292394 APO-PRAMIPEXOLE  
02297329 ACT-PRAMIPEXOLE  
02309149 PRAMIPEXOLE  
02315289 SANDOZ-PRAMIPEXOLE  
02367629 PRAMIPEXOLE  
02424096 AURO-PRAMIPEXOLE

APX FGNQSW  
TEV FGNQSW  
SIV FGNQSW  
SDZ FGNQSW  
SNS FGNQSW  
ARO FGNQSW

1.5MG TABLET

02292408 APO-PRAMIPEXOLE  
02297337 ACT-PRAMIPEXOLE  
02309157 PRAMIPEXOLE  
02315297 SANDOZ-PRAMIPEXOLE  
02367645 PRAMIPEXOLE  
02424118 AURO-PRAMIPEXOLE

APX FGNQSW  
TEV FGNQSW  
SIV FGNQSW  
SDZ FGNQSW  
SNS FGNQSW  
ARO FGNQSW

**ROPINIROLE HCL**

0.25MG TABLET

02314037 RAN-ROPINIROLE  
02316846 TEVA-ROPINIROLE  
02352338 JAMP-ROPINIROLE

RAN FGNQSW  
TEV FGNQSW  
JPC FGNQSW

1MG TABLET

02314053 RAN-ROPINIROLE  
02316854 TEVA-ROPINIROLE  
02352346 JAMP-ROPINIROLE

RAN FGNQSW  
TEV FGNQSW  
JPC FGNQSW

2MG TABLET

02314061 RAN-ROPINIROLE

RAN FGNQSW

02316862	TEVA-ROPINIROLE	TEV	<b>FGNQSW</b>
02352354	JAMP-ROPINIROLE	JPC	<b>FGNQSW</b>

5MG TABLET			
02314088	RAN-ROPINIROLE	RAN	<b>FGNQSW</b>
02316870	TEVA-ROPINIROLE	TEV	<b>FGNQSW</b>

**ROTIGOTINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

2MG TRANSDERMAL PATCH			
02403900	NEUPRO (SA)	UCB	<b>FNQSW</b>

4MG TRANSDERMAL PATCH			
02403927	NEUPRO (SA)	UCB	<b>FNQSW</b>

6MG TRANSDERMAL PATCH			
02403935	NEUPRO (SA)	UCB	<b>FNQSW</b>

8MG TRANSDERMAL PATCH			
02403943	NEUPRO (SA)	UCB	<b>FNQSW</b>

**SELEGILINE HCL**

5MG TABLET			
02068087	TEVA-SELEGILINE	TEV	<b>FGNQSW</b>
02230641	SELEGILINE	AAA	<b>FGNQSW</b>

**28:36.08 ANTICHOLINERIC AGENTS**

**⑤BENZTROPINE MESYLATE**

1MG TABLET			
00706531	PDP-BENZTROPINE	PEN	<b>FGNQSW</b>

1MG/ML INJECTION SOLUTION (2ML)			
02238903	BENZTROPINE OMEGA	OMG	<b>NQ</b>

**PROCYCLIDINE HCL**

5MG TABLET			
00587354	PDP-PROCYCLIDINE	PEN	<b>FGNQSW</b>

**TRIHEXYPHENIDYL HCL**

2MG TABLET			
00545058	TRIHEXYPHENIDYL	AAA	<b>FGNQSW</b>

5MG TABLET  
00545074

TRIHENYDYL

AAA FGNQSW

**28:92.00 MISCELLANEOUS CENTRAL NERVOUS SYSTEM AGENTS**

**⑤ ACAMPROSATE**

[SEE APPENDIX A](#) FOR SA CRITERIA (FOR SUBSTANCE USE HARM REDUCTION DRUG PROGRAM, NO SA IS REQUIRED)

333MG DELAYED RELEASE TABLET

02293269

CAMPRAL (SA)

MYL FLNQSW

**⑤ ATOMOXETINE**

10MG CAPSULE

02262800

STRATTERA

LIL FQW

02314541

TEVA-ATOMOXETINE

TEV FGQW

02318024

APO-ATOMOXETINE

APX FGQW

02381028

PMS-ATOMOXETINE

PMS FGQW

02386410

SANDOZ-ATOMOXETINE

SDZ FGQW

02445883

ATOMOXETINE

SIV FGQW

02467747

ATOMOXETINE

SNS FGQW

02471485

AURO-ATOMOXETINE

ARO FGQW

02506807

JAMP-ATOMOXETINE

JPC FGQW

18MG CAPSULE

02262819

STRATTERA

LIL FQW

02314568

TEVA-ATOMOXETINE

TEV FGQW

02318032

APO-ATOMOXETINE

APX FGQW

02381036

PMS-ATOMOXETINE

PMS FGQW

02386429

SANDOZ-ATOMOXETINE

SDZ FGQW

02445905

ATOMOXETINE

SIV FGQW

02467755

ATOMOXETINE

SNS FGQW

02471493

AURO-ATOMOXETINE

ARO FGQW

02506815

JAMP-ATOMOXETINE

JPC FGQW

25MG CAPSULE

02262827

STRATTERA

LIL FQW

02314576

TEVA-ATOMOXETINE

TEV FGQW

02318040

APO-ATOMOXETINE

APX FGQW

02381044

PMS-ATOMOXETINE

PMS FGQW

02386437

SANDOZ-ATOMOXETINE

SDZ FGQW

02445913

ATOMOXETINE

SIV FGQW

02467763

ATOMOXETINE

SNS FGQW

02471507

AURO-ATOMOXETINE

ARO FGQW

02506823

JAMP-ATOMOXETINE

JPC FGQW

**40MG CAPSULE**

02262835	STRATTERA	LIL	FQW
02314584	TEVA-ATOMOXETINE	TEV	FGQW
02318059	APO-ATOMOXETINE	APX	FGQW
02381052	PMS-ATOMOXETINE	PMS	FGQW
02386445	SANDOZ-ATOMOXETINE	SDZ	FGQW
02445948	ATOMOXETINE	SIV	FGQW
02467771	ATOMOXETINE	SNS	FGQW
02471515	AURO-ATOMOXETINE	ARO	FGQW
02506831	JAMP-ATOMOXETINE	JPC	FGQW

**60MG CAPSULE**

02262843	STRATTERA	LIL	FQW
02314592	TEVA-ATOMOXETINE	TEV	FGQW
02318067	APO-ATOMOXETINE	APX	FGQW
02381060	PMS-ATOMOXETINE	PMS	FGQW
02386453	SANDOZ-ATOMOXETINE	SDZ	FGQW
02445956	ATOMOXETINE	SIV	FGQW
02467798	ATOMOXETINE	SNS	FGQW
02471523	AURO-ATOMOXETINE	ARO	FGQW
02506858	JAMP-ATOMOXETINE	JPC	FGQW

**80MG CAPSULE**

02279347	STRATTERA	LIL	FQW
02318075	APO-ATOMOXETINE	APX	FGQW
02362511	TEVA-ATOMOXETINE	TEV	FGQW
02386461	SANDOZ-ATOMOXETINE	SDZ	FGQW
02467801	ATOMOXETINE	SNS	FGQW
02471531	AURO-ATOMOXETINE	ARO	FGQW
02506866	JAMP-ATOMOXETINE	JPC	FGQW

**100MG CAPSULE**

02279355	STRATTERA	LIL	FQW
02318083	APO-ATOMOXETINE	APX	FGQW
02386488	SANDOZ-ATOMOXETINE	SDZ	FGQW
02467828	ATOMOXETINE	SNS	FGQW
02471558	AURO-ATOMOXETINE	ARO	FGQW
02506874	JAMP-ATOMOXETINE	JPC	FGQW

**EDARAVONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.3MG/ML SOLUTION FOR INJECTION

02475472	RADICAVA (SA)	BMT	NMQW
00904538	RADICAVA (SA)*		NMQW

\*use when drug cost in excess of CPHA maximum

105MG/5ML ORAL SOLUTION  
 02532611 RADICAVA (SA)  
 00904996 RADICAVA (SA)\*  
 \*use when drug cost in excess of CPHA maximum

BMT **NMQW**  
**NMQW**

**RILUZOLE**

50MG TABLET  
 02242763 RILUTEK  
 02352583 APO-RILUZOLE  
 02390299 MYLAN-RILUZOLE

AVN **FNQSW**  
 APX **FGNQSW**  
 MYL **FGNQSW**

**36:26.00 DIABETES MELLITUS**

**NOTE: THE DRUG IDENTIFICATION NUMBERS LISTED IN THIS SECTION ARE FOR BILLING PURPOSES ONLY.**

**BLOOD GLUCOSE TEST STRIP**

97799814 ACCU-CHEK AVIVA (100)  
 97799962 ACCU-CHEK COMPACT (102)  
 97799177 ACCU-CHEK GUIDE (100)  
 97799497 ACCU-CHEK MOBILE (100)  
 97799702 ASCENSIA CONTOUR (100)  
 97799459 CONTOUR NEXT (100)  
 97799564 EZ HEALTH ORACLE (100)  
 97799597 FREESTYLE LITE (100)  
 97799373 GE200 (100)  
 97799403 MEDISURE (100)  
 97799985 ONE TOUCH ULTRA (100)  
 97799475 ONE TOUCH VERIO (100)  
 97799840 PRECISION FREESTYLE/XTRA (100)  
 97799532 TRUE TEST (100)  
 97799602 TRUE TRACK (100)

ROC **DQ**  
 ROC **DQ**  
 ROC **DQ**  
 ROC **DQ**  
 BDD **DQ**  
 BDD **DQ**  
 THI **DQ**  
 ABC **DQ**  
 BIN **DQ**  
 MSR **DQ**  
 LSN **DQ**  
 LSN **DQ**  
 ABC **DQ**  
 TRI **DQ**  
 TRI **DQ**

**URINE GLUCOSE TEST STRIP**

STRIP  
 00977160 DIASTIX

BAY **DQW**

**URINE KETONE TEST STRIP**

STRIP  
 00977322 KETOSTIX

BAY **DQW**

### **36:60.00 THYROID FUNCTION**

#### **THYROTROPIN ALFA**

[SEE APPENDIX A](#) FOR SA CRITERIA

1.1MG INJECTION

02246016 THYROGEN (SA)

GZY FNQSW

### **36:84.00 TUBERCULOSIS**

#### **TUBERCULIN PURIFIED PROTEIN DERIVATIVE**

5TUB/0.5ML INJECTION SOLUTION

00317268 TUBERSOL

AVN I

### **40:08.00 ALKALINIZING AGENTS**

#### **SODIUM BICARBONATE**

500MG TABLET

80022194 SODIUM BICARBONATE  
80030520 JAMP-SODIUM BICARBONATE

SDZ NW  
JPC NW

50MMOL INJECTION SOLUTION (50ML SYRINGE)

00261998 SODIUM BICARBONATE INJECTION

PFI NQ

### **40:12.00 REPLACEMENT AGENTS**

#### **CALCIUM CARBONATE**

250MG TABLET

00999910 CALCIUM CARBONATE

Note: The Drug Identification Number listed is for billing purposes only.

NW

500MG TABLET

00999919 CALCIUM CARBONATE

Note: The Drug Identification Number listed is for billing purposes only.

NW

#### **DEXTROSE**

50% INJECTION SOLUTION (50ML SYRINGE)

00037974	DEXTROSE 50%	HOS	<b>NQ</b>
<b>MAGNESIUM GLUCOHEPTONATE</b>			
100MG/ML ORAL SOLUTION			
00026697	ROUGIER-MAGNESIUM	ROG	<b>FNQSW</b>
80004109	MAGNESIUM-ODAN	ODN	<b>FNQSW</b>
80009357	JAMP-MAGNESIUM	JPC	<b>FNQSW</b>
<b>POTASSIUM CHLORIDE</b>			
2MMOL/ML INJECTION SOLUTION (10ML)			
00037869	POTASSIUM CHLORIDE	PFI	<b>NQ</b>
8MMOL EXTENDED RELEASE TABLET			
80013005	JAMP-K8	JPC	<b>NW</b>
80108882	PRZ-K8	PRZ	<b>NW</b>
20MMOL/15ML ORAL SOLUTION			
02238604	PMS-POTASSIUM CHLORIDE	PMS	<b>NW</b>
80024835	JAMP-POTASSIUM CHLORIDE	JPC	<b>NW</b>
80046782	ODAN POTASSIUM CHLORIDE	ODN	<b>NW</b>
20MMOL EXTENDED RELEASE TABLET			
80004415	ODAN-K 20	ODN	<b>NW</b>
80107649	PRZ-K 20	PRZ	<b>NW</b>
<b>POTASSIUM CITRATE</b>			
25MMOL EFFERVESCENT TABLET			
02085992	K-LYTE	WES	<b>NW</b>
80033602	JAMP-K EFFERVESCENT	JPC	<b>NW</b>
<b>SODIUM CHLORIDE</b>			
0.9% INJECTION SOLUTION (10ML)			
00037796	SODIUM CHLORIDE	PFI	<b>NQ</b>
02304341	SODIUM CHLORIDE	TLG	<b>NQ</b>
0.9% IRRIGATION SOLUTION (1000ML)			
00786160	SODIUM CHLORIDE	BAX	<b>CNQW</b>
<b>STERILE WATER</b>			
INJECTION SOLUTION (10ML)			
02142546	STERILE WATER FOR INJECTION	PFI	<b>CNQ</b>
02299186	STERILE WATER FOR INJECTION	TLG	<b>CNQ</b>



## **40:18.00 ION-REMOVING RESINS**

### **SEVELAMER CARBONATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

800MG TABLET

02461501 ACCEL-SEVELAMER (SA) ACC **FGNQSW**

### **SEVELAMER HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

800MG TABLET

02244310 RENAGEL (SA) AVN **FNQSW**

### **SODIUM POLYSTYRENE SULFONATE**

ORAL POWDER (1G BINDS WITH APPROXIMATELY 1MMOL K+ IN VIVO)

00755338 SOLYSTAT PEN **FGNQSW**

02026961 KAYEXALATE AVN **FNQSW**

02473941 ODAN-SODIUM POLYSTYRENE SULFONATE ODN **FGNQSW**

02497557 JAMP-SODIUM POLYSTYRENE SULFONATE JPC **FGNQSW**

15G/60ML ORAL POWDER

00769541 SOLYSTAT PEN **FGNQSW**

02473968 ODAN-SODIUM POLYSTYRENE SULFONATE ODN **FGNQSW**

### **SUCROFERRIC OXYHYDROXIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

500MG CHEWABLE TABLET

02471574 VELPHORO (SA) VFM **FNQSW**

## **40:20.00 CALORIC AGENTS**

### **TRihePTANOIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

8.3KCAL/ML

02512556 DOJOLVI (SA) UGX **NMQW**

00900021 DOJOLVI (SA)\* **NMQW**

\*use when drug cost in excess of CPHA maximum

## **40:28.00 DIURETICS**

### **⑤CHLORTHALIDONE**

50MG TABLET

00360279	APO-CHLORTHALIDONE	APX	<b>FGNQSW</b>
02523817	JAMP-CHLORTHALIDONE	JPC	<b>FGNQSW</b>

### **ETHACRYNIC ACID**

25MG TABLET

02258528	EDECRIN	BLO	<b>FNQSW</b>
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### **⑤FUROSEMIDE**

20MG TABLET

00337730	TEVA-FUROSEMIDE	TEV	<b>FGNQSW</b>
00396788	APO-FUROSEMIDE	APX	<b>FGNQSW</b>
02351420	FUROSEMIDE	SNS	<b>FGNQSW</b>
02466759	MINT-FUROSEMIDE	MNT	<b>FGNQSW</b>

40MG TABLET

00337749	TEVA-FUROSEMIDE	TEV	<b>FGNQSW</b>
00362166	APO-FUROSEMIDE	APX	<b>FGNQSW</b>
02351439	FUROSEMIDE	SNS	<b>FGNQSW</b>
02466767	MINT-FUROSEMIDE	MNT	<b>FGNQSW</b>

80MG TABLET

00707570	APO-FUROSEMIDE	APX	<b>FGNQSW</b>
00765953	TEVA-FUROSEMIDE	TEV	<b>FGNQSW</b>
02351447	FUROSEMIDE	SNS	<b>FGNQSW</b>
02466775	MINT-FUROSEMIDE	MNT	<b>FGNQSW</b>

10MG/ML ORAL SOLUTION

02224720	LASIX	AVN	<b>FNQSW</b>
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10MG/ML INJECTION

02382539	FUROSEMIDE (2ML AMP)	SDZ	<b>NQ</b>
02527502	FUROSEMIDE (2ML VIAL)	JPC	<b>NQ</b>

10MG/ML INJECTION

00527033	FUROSEMIDE (2ML VIAL)	SDZ	<b>NQ</b>
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### **⑤HYDROCHLOROTHIAZIDE**

12.5MG TABLET

02327856	APO-HYDRO	APX	<b>FGNQSW</b>
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02425947	MINT-HYDROCHLOROTHIAZIDE	MNT	FGNQSW
25MG TABLET			
00021474	TEVA-HYDROCHLOROTHIAZIDE	TEV	FGNQSW
00326844	APO-HYDRO 25	APX	FGNQSW
02247386	PMS-HYDROCHLOROTHIAZIDE	PMS	FGNQSW
02360594	HYDROCHLOROTHIAZIDE	SNS	FGNQSW
02426196	MINT-HYDROCHLOROTHIAZIDE	MNT	FGNQSW

50MG TABLET			
00021482	TEVA-HYDROCHLOROTHIAZIDE	TEV	FGNQSW
00312800	APO-HYDRO 50	APX	FGNQSW
02247387	PMS-HYDROCHLOROTHIAZIDE	PMS	FGNQSW
02360608	HYDROCHLOROTHIAZIDE	SNS	FGNQSW

**⑤INDAPAMIDE HEMIHYDRATE**

1.25MG TABLET			
02240067	MYLAN-INDAPAMIDE	MYL	FGNQSW
02245246	APO-INDAPAMIDE	APX	FGNQSW

2.5MG TABLET			
02153483	MYLAN-INDAPAMIDE	MYL	FGNQSW
02223678	APO-INDAPAMIDE	APX	FGNQSW

**⑤METOLAZONE**

2.5MG TABLET			
00888400	ZAROXOLYN	AVN	FNQSW

**40:28.10 DIURETICS (POTASSIUM SPARING)**

**⑤AMILORIDE HCL & HYDROCHLOROTHIAZIDE**

5MG & 50MG TABLET			
00784400	AA-AMILZIDE	AAA	FGNQSW

**⑤SPIRONOLACTONE**

25MG TABLET			
00028606	ALDACTONE	PFI	FNQSW
00613215	TEVA-SPIRONOLACTONE	TEV	FGNQSW
02488140	MINT-SPIRONOLACTONE	MNT	FGNQSW
02518821	JAMP-SPIRONOLACTONE	JPC	FGNQSW

100MG TABLET			
00285455	ALDACTONE	PFI	FNQSW

00613223	TEVA-SPIRONOLACTONE	TEV	<b>FGNQSW</b>
02488159	MINT-SPIRONOLACTONE	MNT	<b>FGNQSW</b>
02518848	JAMP-SPIRONOLACTONE	JPC	<b>FGNQSW</b>

**⑤ SPIRONOLACTONE & HYDROCHLOROTHIAZIDE**

25MG & 25MG TABLET

00613231	TEVA-SPIRONOLACTONE/HCTZ	TEV	<b>FGNQSW</b>
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50MG & 50MG TABLET

00657182	TEVA-SPIRONOLACTONE/HCTZ	TEV	<b>FGNQSW</b>
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**⑤ TRIAMTERENE & HYDROCHLOROTHIAZIDE**

50MG & 25MG TABLET

00441775	APO-TRIAZIDE	APX	<b>FGNQSW</b>
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00532657	TEVA-TRIAMTERENE/HCTZ	TEV	<b>FGNQSW</b>
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**44:00.00 ENZYMES**

**AGALSIDASE ALFA**

[SEE APPENDIX A](#) FOR SA CRITERIA

1MG/ML VIAL

02249057	REPLAGAL (SA)	TAK	<b>NMQW</b>
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**AGALSIDASE BETA**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG VIAL

02248965	FABRAZYME (SA)	AVN	<b>NMQW</b>
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35MG VIAL

02248966	FABRAZYME (SA)	AVN	<b>NMQW</b>
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**DORNASE ALFA**

[SEE APPENDIX A](#) FOR SA CRITERIA

1MG/ML INHALATION SOLUTION

02046733	PULMOZYME (SA)	HLR	<b>C</b>
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**VELAGLUCERASE ALFA**

[SEE APPENDIX A](#) FOR SA CRITERIA

400 UNIT VIAL

02357119	VPRIV (SA)	SHR	<b>NMQW</b>
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00904378	VPRIV (SA)*		<b>NMQW</b>
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00904379	VPRIV (SA)*		<b>NMQW</b>
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00904380	VPRIV (SA)*		<b>NMQW</b>
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\*use when drug cost in excess of CPHA maximum

### **48:08.00 ANTITUSSIVES**

#### **CODEINE & GUAIFENESIN & PHENIRAMINE**

2MG & 20MG & 1.5MG PER ML SYRUP  
01934740 ROBITUSSIN AC

PFI W

#### **DEXTROMETHORPHAN HBR**

3MG/ML SYRUP  
01944738 BENYLIN DM (SUCROSE & ALCOHOL FREE)

MCL NW

#### **HYDROCODONE**

1MG/ML SYRUP  
02324253 PDP-HYDROCODONE

PEN N

### **48:24.00 MYCOLYTIC AGENTS**

#### **ACETYLCYSTEINE**

200MG/ML VIAL  
02243098 ACETYLCYSTEINE

SDZ CFGNQSW

### **12:12.20 INTERLEUKIN ANTAGONISTS**

#### **BENRALIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

30MG/ML SYRINGE  
02473232 FASENRA (SA)

AZN NMQW

30MG/ML AUTOINJECTOR  
02496135 FASENRA (SA)

AZN NMQW

#### **MEPOLIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG/ML AUTOINJECTOR  
02492989 NUCALA (SA)

GSK NMQW

100MG/ML SYRINGE  
02492997 NUCALA (SA)

GSK NMQW

**48:14.12 CYSTIC FIBROSIS TRANSMEMBRANE CONDUCTANCE  
REGULATOR POTENTIATORS**

**ELEXACAFTOR & TEZACAFTOR & IVACAFTOR & IVACAFTOR**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG & 25MG & 37.5MG TABLET & 75MG TABLET

02526670 TRIKAFTA (SA) VTX C

100MG & 50MG & 75MG TABLET & 150MG TABLET

02517140 TRIKAFTA (SA) VTX C

80MG & 40MG & 60MG & 59.5MG GRANULES

02542285 TRIKAFTA (SA) VTX C

100MG & 50MG & 75MG & 75MG GRANULES

02542277 TRIKAFTA (SA) VTX C

**IVACAFTOR**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG TABLET

02397412 KALYDECO (SA) VTX C

**48:16.00 EXPECTORANTS**

**GUAIFENESIN**

40MG/ML ORAL LIQUID

01931032 ROBITUSSIN (SUCROSE & ALCOHOL FREE) PFI NW

02142783 ROBITUSSIN MUCUS AND PHLEGM PFI NW

02320940 BENYLIN MUCOUS & PHLEGM RELIEF MCL NW

**48:92.00 RESPIRATORY AGENTS, MISCELLANEOUS**

**OMALIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG VIAL

02260565 XOLAIR (SA) NVR NMQW

150MG PREFILLED SYRINGE

02459795 XOLAIR (SA) NVR NMQW

**TEZEPELUMAB**[SEE APPENDIX A](#) FOR SA CRITERIA

210MG/1.91ML PREFILLED SYRINGE

02529548           TEZSPIRE (SA)

AZE   **NMQW**

210MG/1.91ML PREFILLED PEN

02529556           TEZSPIRE (SA)

AZE   **NMQW****52:02.00 ANTIALLERGIC AGENTS****KETOTIFEN**

0.025% OPHTHALMIC DROPS

02242324           ZADITOR

LTH   **FNQSW****OLOPATADINE**

0.1% OPHTHALMIC DROPS

02233143           PATANOL

NVR   **FNQSW**

02305054           APO-OLOPATADINE

APX   **FGNQSW**

02358913           SANDOZ-OLOPATADINE

SDZ   **FGNQSW**

02422727           MINT-OLOPATADINE

MNT   **FGNQSW**

02458411           JAMP-OLOPATADINE

JPC   **FGNQSW****OLOPATADINE**

0.2% OPHTHALMIC DROPS

02362171           PATADAY

NVR   **FNQSW**

02402823           APO-OLOPATADINE

APX   **FGNQSW**

02420171           SANDOZ-OLOPATADINE

SDZ   **FGNQSW**

02508605           MINT-OLOPATADINE

MNT   **FGNQSW****52:04.04 ANTI INFECTIVES (ANTIBIOTICS)****CIPROFLOXACIN**[SEE APPENDIX A](#) FOR SA CRITERIA

0.3% OPHTHALMIC OINTMENT (3.5G)

02200864           CILOXAN (SA)

ALC   **FNQSW**

0.3% OPHTHALMIC SOLUTION

01945270           CILOXAN (SA)

ALC   **FNQSW**

02387131           SANDOZ-CIPROFLOXACIN (SA)

SDZ   **FGNQSW**

**CIPROFLOXACIN & DEXAMETHASONE**[SEE APPENDIX A](#) FOR SA CRITERIA

0.3% &amp; 0.1% OTIC SUSPENSION

02252716	CIPRODEX (SA)	ALC	<b>FNQSW</b>
02481901	TARO-CIPROFLOXACIN/DEXAMETHASONE (SA)	TAR	<b>FGNQSW</b>
02506882	SANDOZ-CIPROFLOXACIN/DEXAMETHASONE(SA)	SDZ	<b>FGNQSW</b>

**ERYTHROMYCIN BASE**

0.5% OPHTHALMIC OINTMENT (3.5G)

01912755	PDP-ERYTHROMYCIN	PEN	<b>FGNQSW</b>
02141574	ERYTHROMYCIN	PSL	<b>FGNQSW</b>

**GATIFLOXACIN**[SEE APPENDIX A](#) FOR SA CRITERIA

0.3% OPHTHALMIC DROPS

02257270	ZYMAR (SA)	ALL	<b>FNQSW</b>
02327260	APO-GATIFLOXACIN (SA)	APX	<b>FGNQSW</b>

**MOXIFLOXACIN**[SEE APPENDIX A](#) FOR SA CRITERIA

0.5% OPHTHALMIC DROPS

02252260	VIGAMOX (SA)	ALC	<b>FNQSW</b>
02406373	APO-MOXIFLOXACIN (SA)	APX	<b>FGNQSW</b>
02411520	SANDOZ-MOXIFLOXACIN (SA)	SDZ	<b>FGNQSW</b>
02432218	PMS-MOXIFLOXACIN (SA)	PMS	<b>FGNQSW</b>
02472120	JAMP-MOXIFLOXACIN (SA)	JPC	<b>FGNQSW</b>
02484757	AG-MOXIFLOXACIN (SA)	ANG	<b>FGNQSW</b>
02529076	MOXIFLOXACIN (SA)	SNS	<b>FGNQSW</b>

**OFLOXACIN**[SEE APPENDIX A](#) FOR SA CRITERIA

0.3% OPHTHALMIC SOLUTION

02143291	OCUFLOX (SA)	ALL	<b>FNQSW</b>
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**POLYMYXIN B & GRAMICIDIN**

10,000U &amp; 0.025MG/ML OPHTHALMIC/OTIC SOLUTION

02239156	POLYSPORIN	JJM	<b>NW</b>
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**TOBRAMYCIN**

0.3% OPHTHALMIC OINTMENT (3.5G)

00614254	TOBREX	ALC	<b>FNQSW</b>
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0.3% OPHTHALMIC SOLUTION

00513962	TOBREX	ALC	<b>FNQSW</b>
02241755	SANDOZ TOBRAMYCIN	SDZ	<b>FGNQSW</b>



**52:04.06 ANTI INFECTIVES (ANTIVIRALS)**

**TRIFLURIDINE**

1% OPHTHALMIC SOLUTION  
00687456 VIROPTIC

VAL **FNQSW**

**52:04.92 MISCELLANEOUS ANTI INFECTIVES**

**CHLORHEXIDINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.12% ORAL RINSE

02237452 PERIDEX (SA)  
02240433 PERICHLOR (SA)

MDA **N**  
PMS **N**

**52:08.00 ANTI INFLAMMATORY AGENTS**

**BECLOMETHASONE DIPROPIONATE**

50MCG/DOSE AQUEOUS NASAL SPRAY

02172712 MYLAN-BECLO AQ.  
02238796 APO-BECLOMETHASONE

MYL **FGNQSW**  
APX **FGNQSW**

**BUDESONIDE**

64MCG/DOSE NASAL SPRAY

02241003 MYLAN-BUDESONIDE AQ

MYL **FGNQSW**

100MCG/DOSE NASAL SPRAY

02230648 MYLAN-BUDESONIDE AQ

MYL **FGNQSW**

**CYCLOSPORINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.1% OPHTHALMIC EMULSION

02484137 VERKAZIA (SA)

SNN **FQW**

**DEXAMETHASONE**

0.1% OPHTHALMIC OINTMENT (3.5G)

00042579 MAXIDEX

ALC **FNQSW**

0.1% OPHTHALMIC SUSPENSION

00042560	MAXIDEX	ALC	<b>FNQSW</b>
<b>DICLOFENAC SODIUM</b>			
0.1% OPHTHALMIC SOLUTION			
01940414	VOLTAREN OPHTHA	ALC	<b>FNQSW</b>
02441020	APO-DICLOFENAC	APX	<b>FGNQSW</b>
02454807	SANDOZ-DICLOFENAC OPHTHA	SDZ	<b>FGNQSW</b>
02475065	DICLOFENAC	PSL	<b>FGNQSW</b>
02475197	MINT-DICLOFENAC	MNT	<b>FGNQSW</b>
02534525	JAMP-DICLOFENAC	JPC	<b>FGNQSW</b>
<b>FLUOROMETHOLONE</b>			
0.1% OPHTHALMIC SUSPENSION			
00247855	FML 0.1%	ALL	<b>FNQSW</b>
00432814	SANDOZ-FLUOROMETHOLONE	SDZ	<b>FGNQSW</b>
<b>FLUOROMETHOLONE ACETATE</b>			
0.1% OPHTHALMIC SUSPENSION			
00756784	FLAREX	ALC	<b>FNQSW</b>
<b>FLUTICASONE PROPIONATE</b>			
50MCG/DOSE AQUEOUS NASAL SPRAY			
02294745	APO-FLUTICASONE	APX	<b>FGNQSW</b>
<b>KETOROLAC TROMETHAMINE</b>			
0.45% OPHTHALMIC SOLUTION			
02369362	ACUVAIL	ABV	<b>FNQSW</b>
0.5% OPHTHALMIC SOLUTION			
01968300	ACULAR	ALL	<b>FNQSW</b>
02245821	KETOROLAC	AAA	<b>FGNQSW</b>
<b>MOMETASONE</b>			
50MCG/DOSE NASAL SPRAY			
02238465	NASONEX	MSD	<b>FNQSW</b>
02403587	APO-MOMETASONE	APX	<b>FGNQSW</b>
02449811	SANDOZ-MOMETASONE	SDZ	<b>FGNQSW</b>
02475863	TEVA-MOMETASONE	TEV	<b>FGNQSW</b>
02519127	MOMETASONE	SNS	<b>FGNQSW</b>
<b>PREDNISOLONE ACETATE</b>			
1% OPHTHALMIC SUSPENSION			
00301175	PRED FORTE	ALL	<b>FNQSW</b>
00700401	TEVA-PREDNISOLONE	TEV	<b>FGNQSW</b>
01916203	SANDOZ-PREDNISOLONE	SDZ	<b>FGNQSW</b>

**TRIAMCINOLONE**

55MCG/DOSE NASAL SPRAY

02213834

NASACORT AQ

02437635

APO-TRIAMCINOLONE AQ

SNC

FNQSW

APX

FGNQSW

**52:08.08 COMBINATION ANTI-INFECTIVE / ANTI INFLAMMATORY AGENTS**

**CLIOQUINOL & FLUMETHASONE PIVALATE**

1% & 0.02% OTIC SOLUTION

00074454

LOCACORTEN-VIOFORM

PAL

FNQSW

**FRAMYCETIN SULFATE & GRAMICIDIN & DEXAMETHASONE**

5MG & 50MCG & 0.5MG/ML OPHTHALMIC/OTIC SOLUTION

02224623

SOFACORT

AVN

FNQSW

**TOBRAMYCIN & DEXAMETHASONE**

0.3% & 0.1% OPHTHALMIC OINTMENT

00778915

TOBRADEX

ALC

FNQSW

0.3% & 0.1% OPHTHALMIC SUSPENSION

00778907

TOBRADEX

ALC

FNQSW

**52:10.00 CARBONIC ANHYDRASE INHIBITORS**

**ACETAZOLAMIDE**

250MG TABLET

00545015

ACETAZOLAMIDE

AAA

FGNQSW

**BRINZOLAMIDE**

02238873

AZOPT

ALC

FNQSW

**DORZOLAMIDE HCL**

2% OPHTHALMIC SOLUTION

02216205

TRUSOPT

ELV

FNQSW

02316307

SANDOZ-DORZOLAMIDE

SDZ

FGNQSW

02453347

JAMP-DORZOLAMIDE

JPC

FGNQSW

02457210

MED-DORZOLAMIDE

GMP

FGNQSW

02522373

DORZOLAMIDE

JPC

FGNQSW

**METHAZOLAMIDE**

50MG TABLET

02245882

METHAZOLAMIDE

AAA FGNQSW

**52:24.00 MYDRIATICS**

**ATROPINE SULFATE**

1% OPHTHALMIC SOLUTION

00035017 ISOPTO ATROPINE

ALC FNQSW

02023695 ATROPINE

PSL FGNQSW

**CYCLOPENTOLATE**

1% OPHTHALMIC SOLUTION

00252506 CYCLOGYL

ALC FNQSW

**PHENYLEPHRINE HCL**

2.5% OPHTHALMIC SOLUTION

00465763 MYDFRIN

ALC FNQSW

**52:28.00 MOUTHWASHES AND GARGLES**

**BENZYDAMINE HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.15% ORAL RINSE

02239537 PMS-BENZYDAMINE (SA)

PMS FGNQSW

02463105 ODAN-BENZYDAMINE (SA)

ODN FGNQSW

**52:32.00 VASOCONSTRICTORS**

**XYLOMETAZOLINE**

0.1% NASAL SPRAY

00653330 OTRIVIN

NVR N

**52:40.00 ALPHA AND BETA ADRENERGIC AGENTS AND PROSTAGLANDIN ANALOGS**

**BETAXOLOL HCL**

0.25% OPHTHALMIC SUSPENSION

01908448	BETOPTIC S	ALC	FNQSW
<b>BIMATOPROST</b>			
0.1 MG/ML OPHTHALMIC SOLUTION			
02324997	LUMIGAN	ALL	FNQSW
<b>BRIMONIDINE TARTRATE</b>			
0.15% OPHTHALMIC SOLUTION			
02248151	ALPHAGAN P	ALL	FNQSW
02301334	BRIMONIDINE P	AAA	FGNQSW
0.2% OPHTHALMIC SOLUTION			
02236876	ALPHAGAN	ALL	FNQSW
02260077	APO-BRIMONIDINE	APX	FGNQSW
02305429	SANDOZ BRIMONIDINE	SDZ	FGNQSW
02449226	JAMP-BRIMONIDINE	JPC	FGNQSW
02507811	MED-BRIMONIDINE	GMP	FGNQSW
02515377	BRIMONIDINE TARTRATE	TEL	FGNQSW
<b>BRIMONIDINE &amp; TIMOLOL</b>			
0.2% & 0.5% OPHTHALMIC SOLUTION			
02248347	COMBIGAN	ALL	FNQSW
02375311	APO-BRIMONIDINE-TIMOP	APX	FGNQSW
02531704	JAMP-BRIMONIDINE-TIMOLOL	JP	FGNQSW
<b>BRINZOLAMIDE &amp; BRIMONIDINE</b>			
1% & 0.2% OPHTHALMIC SUSPENSION			
02435411	SIMBRINZA	ALC	FNQSW
<b>BRINZOLAMIDE &amp; TIMOLOL</b>			
1% & 0.5% OPHTHALMIC SUSPENSION			
02331624	AZARGA	ALC	FNQSW
<b>DORZOLAMIDE &amp; TIMOLOL</b>			
2% & 0.5% OPHTHALMIC SOLUTION			
02240113	COSOPT	ELV	FNQSW
02299615	APO-DORZO-TIMOP	APX	FGNQSW
02437686	MED-DORZOLAMIDE-TIMOLOL	GMP	FGNQSW
02344351	SANDOZ-DORZOLAMIDE/TIMOLOL	SDZ	FGNQSW
02441659	RIVA-DORZOLAMIDE/TIMOLOL	RIV	FGNQSW
02457539	JAMP-DORZOLAMIDE/TIMOLOL	JPC	FGNQSW
02489635	DORZOLAMIDE AND TIMOLOL	TEL	FGNQSW
02522020	DORZOLAMIDE-TIMOLOL	JPC	FGNQSW
02537796	M-DORZOLAMIDE-TIMOLOL	MRA	FGNQSW
<b>LATANOPROST</b>			

50MCG/ML OPHTHALMIC SOLUTION

02231493	XALATAN	UJC	FNQSW
02254786	TEVA-LATANOPROST	TEV	FGNQSW
02296527	APO-LATANOPROST	APX	FGNQSW
02317125	PMS-LANANOPROST	PMS	FGNQSW
02367335	SANDOZ-LATANOPROST	SDZ	FGNQSW
02373041	GD-LATANOPROST	UJC	FGNQSW
02426935	MED-LATANOPROST	GMP	FGNQSW
02453355	JAMP-LATANOPROST	JPC	FGNQSW
02489570	LATANOPROST	TLG	FGNQSW
02513285	M-LATANOPROST	MRA	FGNQSW

**LATANOPROST & TIMOLOL**

50MCG & 5MG PER ML OPHTHALMIC SOLUTION

02246619	XALACOM	UJC	FNQSW
02373068	GD-LATANOPROST/TIMOLOL	UJC	FGNQSW
02436256	TEVA-LATANOPROST/TIMOLOL	TEV	FGNQSW
02453770	JAMP-LATANOPROST/TIMOLOL	JPC	FGNQSW
02454505	MED-LATANOPROST/TIMOLOL	MED	FGNQSW
02514516	M-LATANOPROST/TIMOLOL	MRA	FGNQSW

**LATANOPROSTENE BUNOD**

0.024% OPHTHALMIC SOLUTION

02484218	VYZULTA	BLO	FNQSW
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**TIMOLOL MALEATE**

0.25% OPHTHALMIC SOLUTION

02166712	SANDOZ-TIMOLOL	SDZ	FGNQSW
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0.5% OPHTHALMIC SOLUTION

00451207	TIMOPTIC	ELV	FNQSW
00755834	APO-TIMOP	APX	FGNQSW
02166720	SANDOZ-TIMOLOL MALEATE	SDZ	FGNQSW
02447800	JAMP-TIMOLOL	JPC	FGNQSW

0.25% GEL FORMING SOLUTION

02242275	TIMOLOL MALEATE-EX	SDZ	FGNQSW
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0.5% GEL FORMING SOLUTION

02171899	TIMOPTIC-XE	PFR	FNQSW
02242276	TIMOLOL MALEATE-EX	SDZ	FGNQSW

**TRAVOPROST**

0.003% OPHTHALMIC SOLUTION

02457997	IZBA	NVR	FNQSW
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0.004% OPHTHALMIC SOLUTION		ALC	<b>FNQSW</b>
02318008	TRAVATAN Z	SDZ	<b>FGNQSW</b>
02413167	SANDOZ-TRAVOPROST		

**TRAVOPROST & TIMOLOL**

0.004% & 0.5% OPHTHALMIC SOLUTION		ALC	<b>FNQSW</b>
02278251	DUOTRAV	APX	<b>FGNQSW</b>
02415305	APO-TRAVOPROST-TIMOLOL		

**52:40.20 MIOTICS**

**PILOCARPINE HCL**

2% OPHTHALMIC SOLUTION		ALC	<b>FNQSW</b>
00000868	ISOPTO CARPINE		

4% OPHTHALMIC SOLUTION		ALC	<b>FNQSW</b>
00000884	ISOPTO CARPINE		

**52:92.00 EYE, EAR, NOSE, AND THROAT DRUGS, MISCELLANEOUS**

**AFLIBERCEPT**

[SEE APPENDIX A](#) FOR SA CRITERIA

2MG/0.05ML VIAL		BAY	<b>NMQW</b>
02415992	EYLEA (SA)		

**APRACLONIDINE HCL**

0.5% OPHTHALMIC SOLUTION		EPM	<b>FNQSW</b>
02076306	IOPIDINE		

**ARTIFICIAL TEARS**

0.5% OPHTHALMIC SOLUTION		ALC	<b>NW</b>
00000809	ALCON TEARS		

1% OPHTHALMIC SOLUTION		ALC	<b>NW</b>
00000817	ALCON TEARS		

5% OPHTHALMIC OINTMENT		BLO	<b>NW</b>
00750816	MURO-128		

**BROLUCIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

6 MG/0.05 ML PREFILLED SYRINGE  
02496976 BEOVU (SA) NVR **NMQW**

**LANOLIN & MINERAL OIL & PETROLATUM**

3 % & 3 % & 94 % OINTMENT  
02444062 SYSTANE ALC **NW**

**RANIBIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

2.3MG/0.23ML VIAL  
02296810 LUCENTIS (SA) NVR **NMQW**

02525852 BYOOVIZ (SA) BGH **NMQW**

02542250 RANOPTO (SA) TEV **NMQW**

0.5MG/0.5ML PREFILLED SYRINGE

02425629 LUCENTIS (SA) NVR **NMQW**

**CROMOLYN SODIUM**

2% OPHTALMIC SOLUTION

02009277 CROMOLYN PEN **FNSW**

**FARICIMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

6MG & 0.05ML VIAL

02527618 VABYSMO (SA) HLR **NMQW**

**56:04.00 ANTACIDS AND ADSORBENTS**

**ALGINIC ACID & ALUMINIUM HYDROXIDE**

50MG & 20MG/ML ORAL SUSPENSION

02159775 GAVISCON GSK **NW**

**ALGINIC ACID & MAGNESIUM CARBONATE**

200MG & 40MG TABLET

02159791 GAVISCON HEARTBURN RELIEF GSK **NW**

**MAGNESIUM HYDROXIDE & ALUMINIUM HYDROXIDE**

40MG & 33MG/ML ORAL SUSPENSION

01966529 DIOVOL CDC **NW**

**MAGNESIUM HYDROXIDE & ALUMINIUM HYDROXIDE & SIMETHICONE**

200MG & 200MG & 25MG TABLET



00116882

DIOVOL PLUS

CDC NW

**56:08.00 ANTIDIARRHEA AGENTS**

**DIPHENOXYLATE HCL/ATROPINE SULFATE**

2.5MG/0.025MG TABLET

00036323

LOMOTIL

PFI FNQSW

**LOPERAMIDE**

2MG CAPLET

02132591

TEVA-LOPERAMIDE

TEV FNQSW

02183862

IMODIUM

MCL FNQSW

02212005

APO-LOPERAMIDE

APX FNQSW

02228351

PMS-LOPERAMIDE

PMS FNQSW

02229552

DIARR-EZE

PMS FNQSW

0.2MG/ML ORAL SOLUTION

02016095

PMS-LOPERAMIDE HCL

PMS FNQSW

**56:10.00 ANTIFLATULENTS**

**SIMETHICONE**

80MG TABLET

00292990

OVOL

CDC NW

**56:12.00 CATHARTICS AND LAXATIVES**

Note: Cathartics and laxatives should only be used after failure of simpler measures. A high fiber diet, adequate hydration, and a review of potentially constipating medications is often effective in relieving constipation.

**BISACODYL**

5MG ENTERIC COATED TABLET

00254142

DULCOLAX

SNC NW

02273411

ODAN-BISACODYL

ODN NW

10MG RECTAL SUPPOSITORY

00003875

DULCOLAX

SNC NW

02361450

JAMP-BISACODYL

JPC NW

02520478	AMB-BISACODYL	AMB	<b>NW</b>
10MG RECTAL SUPPOSITORY			
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
02241091	MAGIC BULLET (SA)	D&C	<b>FNSW</b>

**GLYCERIN**

90%/2.6G SUPPOSITORY			
02020394	GLYCERIN	ROG	<b>N</b>

**LACTULOSE**

[SEE APPENDIX A](#) FOR SA CRITERIA

667MG/ML SYRUP

00703486	PMS-LACTULOSE (SA)	PMS	<b>NW</b>
00854409	RATIO-LACTULOSE (SA)	TEV	<b>NW</b>
02295881	JAMP-LACTULOSE (SA)	JPC	<b>NW</b>
02412268	LACTULOSE (SA)	SNS	<b>NW</b>
02469391	PMS-LACTULOSE-PHARMA (SA)	PMS	<b>NW</b>

**MAGNESIUM CITRATE**

50MG/ML ORAL SOLUTION

00262609	CITRO-MAG	ROG	<b>NW</b>
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**MAGNESIUM HYDROXIDE & MINERAL OIL**

60MG & 0.25ML PER ML ORAL EMULSION

00202045	MAGNOLAX	PEN	<b>N</b>
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**POLYETHYLENE GLYCOL 3350**

ORAL POWDER

09991054	POLYETHYLENE GLYCOL 3350		<b>NW</b>
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Note: The Drug Identification Number listed is for billing purposes only.

**PSYLLIUM MUCILLOID**

ORAL POWDER

02174782	METAMUCIL SUGAR FREE	PGA	<b>NW</b>
02174812	METAMUCIL	PGA	<b>NW</b>

**SENNOSIDES A&B**

8.6MG TABLET

00026158	SENOKOT	ANB	<b>NW</b>
00896411	PMS-SENNOSIDES	PMS	<b>NW</b>
80009595	JAMP-SENNA	JPC	<b>NW</b>

1.7MG/ML ORAL LIQUID

00367729	SENOKOT	PFR	<b>N</b>
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**SODIUM PHOSPHATES**

220MG/ML ENEMA (130ML)  
00009911 FLEET

JJM NW

**56:14.00 CHOLELITHOLYTIC AGENTS**

**URSODIOL**

250MG TABLET

02273497	PMS-URSODIOL C	PMS	<b>FGNQSW</b>
02426900	GLN-URSODIOL	GLM	<b>FGNQSW</b>
02472392	JAMP-URSODIOL	JPC	<b>FGNQSW</b>
02505363	AG-URSODIOL	ANG	<b>FGNQSW</b>
02515520	URSODIOL	SNS	<b>FGNQSW</b>

500MG TABLET

02273500	PMS-URSODIOL C	PMS	<b>FGNQSW</b>
02426919	GLN-URSODIOL	GLM	<b>FGNQSW</b>
02472406	JAMP-URSODIOL	JPC	<b>FGNQSW</b>
02505371	AG-URSODIOL	ANG	<b>FGNQSW</b>
02515539	URSODIOL	SNS	<b>FGNQSW</b>

**56:16.00 DIGESTANTS**

**LIPASE & PROTEASE & AMYLASE**

5,000 & 320 & 5,100 UNIT GRANULES

02445158 CREON MINIMICROSPHERES MICRO BGP **CFNQSW**

10,000 & 730 & 11,200 UNIT CAPSULE

02200104 CREON 10 MINIMICROSPHERES BGP **CFNQSW**

25,000 & 1,600 & 25,500 UNIT CAPSULE

01985205 CREON 25 MINIMICROSPHERES BGP **CFNQSW**

35,000 & 2,240 & 35,700 UNIT CAPSULE

02494639 CREON 35 MINIMICROSPHERES BGP **CFNQSW**

**PANCRELIPASE EQUIVALENT TO LIPASE & PROTEASE & AMYLASE**

10,000 & 35,000 & 40,000USP U CAPSULE

00263818 COTAZYM MSD **CFNQSW**

10,800 & 45,000 & 42,000	USP U CAPSULE (ENTERIC COATED PARTICLES)		
00502790	COTAZYM ECS 8	MSD	<b>CFNQSW</b>
25,000 & 100,000 & 100,000	USP U CAPSULE (ENTERIC COATED PARTICLES)		
00821373	COTAZYM ECS 20	MSD	<b>CFNQSW</b>
10,440 & 57,100 & 56,400	USP U TABLET		
02230019	VIOKACE	ARN	<b>CFNQSW</b>
20,880 & 112,500 & 113,400	USP U TABLET		
02241933	VIOKACE	ARN	<b>CFNQSW</b>

**56:22.00 ANTIEMETICS**

**APREPITANT**

[SEE APPENDIX A](#) FOR SA CRITERIA

80MG CAPSULE			
02298791	EMEND (SA)	MSD	<b>FNQSW</b>
125MG CAPSULE			
02298805	EMEND (SA)	MSD	<b>FNQSW</b>
80MG & 80MG & 125MG CAPSULE (PACKAGE)			
02298813	EMEND TRI-PACK (SA)	MSD	<b>FNQSW</b>

**DIMENHYDRINATE**

50MG TABLET			
00999972	DIMENHYDRINATE		<b>NW</b>

Note: The Drug Identification Number listed is for billing purposes only.

50MG RECTAL SUPPOSITORY			
00392553	SANDOZ-DIMENHYDRINATE	SDZ	<b>NW</b>

100MG RECTAL SUPPOSITORY			
00013609	GRAVOL	CDC	<b>NW</b>
00392545	SANDOZ-DIMENHYDRINATE	SDZ	<b>NW</b>

50MG/ML INTRAMUSCULAR INJECTION SOLUTION (5ML)			
00392537	DIMENHYDRINATE IM	SDZ	<b>NW</b>

**DOXYLAMINE SUCCINATE & PYRIDOXINE HCL**

10MG & 10MG DELAYED RELEASE TABLET			
00609129	DICLECTIN	DUI	<b>FQW</b>

02406187	PMS-DOXYLAMINE-PYRIDOXINE	PMS	<b>FGQW</b>
02413248	APO-DOXYLAMINE/B6	APX	<b>FGQW</b>

**GRANISETRON**

[SEE APPENDIX A](#) FOR SA CRITERIA

1MG TABLET

02308894	APO-GRANISETRON (SA)	APX	<b>FGNQSW</b>
02452359	NAT-GRANISETRON (SA)	NAT	<b>FGNQSW</b>
02472686	JAMP GRANISETRON (SA)	JPC	<b>FGNQSW</b>

**NABILONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.5MG CAPSULE

02256193	CESAMET (SA)	VAL	<b>FNQSW</b>
02380900	PMS-NABILONE (SA)	PMS	<b>FNQSW</b>
02384884	TEVA-NABILONE (SA)	TEV	<b>FNQSW</b>

1MG CAPSULE

00548375	CESAMET (SA)	VAL	<b>FNQSW</b>
02380919	PMS-NABILONE (SA)	PMS	<b>FNQSW</b>
02384892	TEVA-NABILONE (SA)	TEV	<b>FNQSW</b>

**NETUPITANT & PALONOSETRON HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

300MG/0.5MG CAPSULE

02468735	AKYNZEO (SA)	ELV	<b>FNQSW</b>
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**ONDANSETRON**

[SEE APPENDIX A](#) FOR SA CRITERIA

4MG MEDICATED FILM

02389983	ONDISSOLVE ODT(SA)	TAK	<b>FNQSW</b>
02541351	JAMP-ONDANSETRON ODT (SA)	JPC	<b>FGNQSW</b>

8MG MEDICATED FILM

02389991	ONDISSOLVE ODT(SA)	TAK	<b>FNQSW</b>
02541378	JAMP-ONDANSETRON ODT (SA)	JPC	<b>FGNQSW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA

4MG ORAL DISINTEGRATING TABLET

02481723	ONDANSETRON ODT (SA)	SDZ	<b>FGNQSW</b>
02487330	MINT-ONDANSETRON ODT (SA)	MNT	<b>FGNQSW</b>
02511282	AURO-ONDANSETRON ODT (SA)	ARO	<b>FGNQSW</b>
02514966	MAR-ONDANSETRON ODT (SA)	MAR	<b>FGNQSW</b>
02519232	ONDANSETRON ODT (SA)	JPC	<b>FGNQSW</b>
02519445	PMS-ONDANSETRON ODT (SA)	PMS	<b>FGNQSW</b>
02524279	ONDANSETRON ODT (SA)	SNS	<b>FGNQSW</b>

8MG ORAL DISINTEGRATING TABLET

02481731	ONDANSETRON ODT (SA)	SDZ	<b>FGNQSW</b>
02487349	MINT-ONDANSETRON ODT (SA)	MNT	<b>FGNQSW</b>
02511290	AURO-ONDANSETRON ODT (SA)	ARO	<b>FGNQSW</b>
02514974	MAR-ONDANSETRON ODT (SA)	MAR	<b>FGNQSW</b>
02519240	ONDANSETRON ODT (SA)	JPC	<b>FGNQSW</b>
02519453	PMS-ONDANSETRON ODT (SA)	PMS	<b>FGNQSW</b>
02524287	ONDANSETRON ODT(SA)	SNS	<b>FGNQSW</b>

**ONDANSETRON HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

4MG TABLET

02213567	ZOFRAN (SA)	NVR	<b>FNQSW</b>
02258188	PMS-ONDANSETRON (SA)	PMS	<b>FGNQSW</b>
02274310	SANDOZ-ONDANSETRON (SA)	SDZ	<b>FGNQSW</b>
02288184	APO-ONDANSETRON (SA)	APX	<b>FGNQSW</b>
02296349	TEVA-ONDANSETRON (SA)	TEV	<b>FGNQSW</b>
02297868	MYLAN-ONDANSETRON (SA)	MYL	<b>FGNQSW</b>
02305259	MINT-ONDANSETRON (SA)	MNT	<b>FGNQSW</b>
02312247	RAN-ONDANSETRON (SA)	RAN	<b>FGNQSW</b>
02313685	JAMP-ONDANSETRON (SA)	JPC	<b>FGNQSW</b>
02371731	MAR-ONDANSETRON (SA)	MAR	<b>FGNQSW</b>
02417839	NAT-ONDANSETRON (SA)	NAT	<b>FGNQSW</b>
02421402	ONDANSETRON (SA)	SNS	<b>FGNQSW</b>
02541424	ONDANSETRON (SA)	SIV	<b>FGNQSW</b>

8MG TABLET

02213575	ZOFRAN (SA)	NVR	<b>FNQSW</b>
02258196	PMS-ONDANSETRON (SA)	PMS	<b>FGNQSW</b>
02274329	SANDOZ-ONDANSETRON (SA)	SDZ	<b>FGNQSW</b>
02288192	APO-ONDANSETRON (SA)	APX	<b>FGNQSW</b>
02296357	TEVA-ONDANSETRON (SA)	TEV	<b>FGNQSW</b>
02297876	MYLAN-ONDANSETRON (SA)	MYL	<b>FGNQSW</b>
02305267	MINT-ONDANSETRON (SA)	MNT	<b>FGNQSW</b>
02312255	RAN-ONDANSETRON (SA)	RAN	<b>FGNQSW</b>
02313693	JAMP-ONDANSETRON (SA)	JPC	<b>FGNQSW</b>
02371758	MAR-ONDANSETRON (SA)	MAR	<b>FGNQSW</b>
02417847	NAT-ONDANSETRON (SA)	NAT	<b>FGNQSW</b>
02421410	ONDANSETRON (SA)	SNS	<b>FGNQSW</b>
02541432	ONDANSETRON (SA)	SIV	<b>FGNQSW</b>

0.8MG/ML ORAL SOLUTION

02229639	ZOFRAN (SA)	NVR	<b>FNQSW</b>
02291967	APO-ONDANSETRON (SA)	APX	<b>FGNQSW</b>
02490617	JAMP-ONDANSETRON (SA)	JPC	<b>FGNQSW</b>

## **56:28.12 HISTAMINE H2 ANTAGONISTS**

### **CIMETIDINE**

200MG TABLET  
00584215

CIMETIDINE

AAA **FGNQSW**

300MG TABLET  
00487872

CIMETIDINE

AAA **FGNQSW**

### **FAMOTIDINE**

20MG TABLET

02022133

TEVA-FAMOTIDINE

TEV **FGNQSW**

02507749

JAMP-FAMOTIDINE

JPC **FGNQSW**

02538628

MINT-FAMOTIDINE

MNT **FGNQSW**

40MG TABLET

02022141

TEVA-FAMOTIDINE

TEV **FGNQSW**

02507757

JAMP-FAMOTIDINE

JPC **FGNQSW**

02538636

MINT-FAMOTIDINE

MNT **FGNQSW**

### **NIZATIDINE**

150MG CAPSULE

00778338

AXID

PEN **FNQSW**

### **RANITIDINE HCL**

150MG TABLET

00733059

APO-RANITIDINE

APX **FGNQSW**

02242453

PMS-RANITIDINE

PMS **FGNQSW**

02336480

RAN-RANITIDINE

RAN **FGNQSW**

02353016

RANITIDINE

SNS **FGNQSW**

02443708

MAR-RANITIDINE

MAR **FGNQSW**

02463717

JAMP-RANITIDINE

JPC **FGNQSW**

02526379

MINT-RANITIDINE

MNT **FGNQSW**

300MG TABLET

00733067

APO-RANITIDINE

APX **FGNQSW**

02242454

PMS-RANITIDINE

PMS **FGNQSW**

02336502

RAN-RANITIDINE

RAN **FGNQSW**

02353024

RANITIDINE

SNS **FGNQSW**

02443716

MAR-RANITIDINE

MAR **FGNQSW**

02463725

JAMP-RANITIDINE

JPC **FGNQSW**

02526387

MINT-RANITIDINE

MNT **FGNQSW**

15MG/ML ORAL SOLUTION			
02280833	APO-RANITIDINE	APX	<b>FGNQSW</b>

**56:28.28 PROSTAGLANDINS**

**MISOPROSTOL**

100MCG TABLET			
02244022	MISOPROSTOL	AAA	<b>FGNQSW</b>

200MCG TABLET			
02244023	MISOPROSTOL	AAA	<b>FGNQSW</b>

**56:28.32 PROTECTANTS**

**SUCRALFATE**

1G TABLET			
02045702	TEVA-SUCRALFATE	TEV	<b>FGNQSW</b>
02100622	SULCRATE	ALL	<b>FNQSW</b>
02125250	APO-SUCRALFATE	APX	<b>FGNQSW</b>

200MG/ML ORAL SUSPENSION			
02103567	SULCRATE PLUS	AVN	<b>FNQSW</b>

**56:28.36 PROTON PUMP INHIBITORS**

**LANSOPRAZOLE**

SEE PROTON PUMP INHIBITORS IN APPENDIX A FOR [SA CRITERIA](#)

15MG DELAYED RELEASE CAPSULE			
02165503	PREVACID (SA)	ABB	<b>FNQSW</b>
02280515	TEVA-LANSOPRAZOLE (SA)	TEV	<b>FGNQSW</b>
02293811	APO-LANSOPRAZOLE (SA)	APX	<b>FGNQSW</b>
02353830	MYLAN-LANSOPRAZOLE (SA)	MYL	<b>FGNQSW</b>
02357682	LANSOPRAZOLE (SA)	SNS	<b>FGNQSW</b>
02385643	SANDOZ-LANSOPRAZOLE (SA)	SDZ	<b>FGNQSW</b>
02385767	LANSOPRAZOLE DR (SA)	SIV	<b>FGNQSW</b>
02395258	PMS-LANSOPRAZOLE (SA)	PMS	<b>FGNQSW</b>
02402610	TARO-LANSOPRAZOLE (SA)	SUN	<b>FGNQSW</b>
02433001	LANSOPRAZOLE (SA)	PMS	<b>FGNQSW</b>



**30MG DELAYED RELEASE CAPSULE**

02165511	PREVACID (SA)	ABB	<b>FNQSW</b>
02280523	TEVA-LANSOPRAZOLE (SA)	TEV	<b>FGNQSW</b>
02293838	APO-LANSOPRAZOLE (SA)	APX	<b>FGNQSW</b>
02353849	MYLAN-LANSOPRAZOLE (SA)	MYL	<b>FGNQSW</b>
02357690	LANSOPRAZOLE (SA)	SNS	<b>FGNQSW</b>
02385651	SANDOZ-LANSOPRAZOLE (SA)	SDZ	<b>FGNQSW</b>
02395266	PMS-LANSOPRAZOLE (SA)	PMS	<b>FGNQSW</b>
02402629	TARO-LANSOPRAZOLE (SA)	SUN	<b>FGNQSW</b>
02410389	LANSOPRAZOLE (SA)	SIV	<b>FGNQSW</b>
02433028	LANSOPRAZOLE (SA)	PMS	<b>FGNQSW</b>

**15MG DELAYED RELEASE TABLET**

02249464	PREVACID FASTAB (SA)	ABB	<b>FNQSW</b>
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**30MG DELAYED RELEASE TABLET**

02249472	PREVACID FASTAB (SA)	ABB	<b>FNQSW</b>
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**LANSOPRAZOLE & CLARITHROMYCIN & AMOXICILLIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

**30MG & 500MG & 500MG 7-DAY PACKAGE**

02470780	AA-LANSOPRAZOLE-AMOXICILLIN-CLARITHROMYCIN	AAA	<b>FGNQSW</b>
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**OMEPRAZOLE**

[SEE PROTON PUMP INHIBITORS IN APPENDIX A](#) FOR SA CRITERIA **FOR DOSAGES GREATER THAN 20MG PER DAY.**

**10MG CAPSULE**

02295407	TEVA-OMEPRAZOLE	TEV	<b>FGNQSW</b>
02296438	SANDOZ-OMEPRAZOLE	SDZ	<b>FGNQSW</b>

**20MG CAPSULE**

00846503	LOSEC	XPI	<b>FNQSW</b>
02245058	APO-OMEPRAZOLE	APX	<b>FGNQSW</b>
02296446	SANDOZ-OMEPRAZOLE	SDZ	<b>FGNQSW</b>
02348691	OMEPRAZOLE	SNS	<b>FGNQSW</b>
02411857	OMEPRAZOLE-20	SIV	<b>FGNQSW</b>

**20MG DELAYED RELEASE TABLET**

02295415	TEVA-OMEPRAZOLE	TEV	<b>FGNQSW</b>
02416549	OMEPRAZOLE MAGNESIUM	ACH	<b>FGNQSW</b>
02420198	JAMP-OMEPRAZOLE	JPC	<b>FGNQSW</b>
02439549	NAT-OMEPRAZOLE	NAT	<b>FGNQSW</b>
02501880	NRA-OMEPRAZOLE MAGNESIUM	NRA	<b>FGNQSW</b>
02504294	OMEPRAZOLE MAGNESIUM	SNS	<b>FGNQSW</b>

**PANTOPRAZOLE MAGNESIUM**

[SEE PROTON PUMP INHIBITORS IN APPENDIX A](#) FOR SA CRITERIA **FOR DOSAGES GREATER THAN 40 MG/DAY**

40MG ENTERIC TABLET

02267233	TECTA	TAK	FNQSW
02408570	MYLAN-PANTOPRAZOLE T	MYL	FGNQSW
02440628	TEVA-PANTOPRAZOLE MAGNESIUM	TEV	FGNQSW
02441853	PANTOPRAZOLE MAGNESIUM	ALH	FGNQSW
02466147	PANTOPRAZOLE T	SNS	FGNQSW
02519534	PANTOPRAZOLE T	SIV	FGNQSW

**PANTOPRAZOLE SODIUM**

[SEE PROTON PUMP INHIBITORS IN APPENDIX A](#) FOR SA CRITERIA **FOR DOSAGES GREATER THAN ONE UNIT/DAY**

20MG ENTERIC TABLET

02241804	PANTOLOC	TAK	FNQSW
02285479	TEVA-PANTOPRAZOLE	TEV	FGNQSW
02292912	APO-PANTOPRAZOLE	APX	FGNQSW
02301075	SANDOZ-PANTOPRAZOLE	SDZ	FGNQSW
02392615	JAMP-PANTOPRAZOLE SODIUM	JPC	FGNQSW
02408414	JAMP-PANTOPRAZOLE EC	JPC	FGNQSW
02428172	PANTOPRAZOLE	SIV	FGNQSW
02536137	PANTOPRAZOLE	SNS	FGNQSW

40MG ENTERIC TABLET

02229453	PANTOLOC	TAK	FNQSW
02285487	TEVA-PANTOPRAZOLE	TEV	FGNQSW
02292920	APO-PANTOPRAZOLE	APX	FGNQSW
02301083	SANDOZ-PANTOPRAZOLE	SDZ	FGNQSW
02305046	RAN-PANTOPRAZOLE	RAN	FGNQSW
02307871	PMS-PANTOPRAZOLE	PMS	FGNQSW
02357054	JAMP-PANTOPRAZOLE EC	JPC	FGNQSW
02370808	PANTOPRAZOLE	SNS	FGNQSW
02392623	JAMP-PANTOPRAZOLE SODIUM	JPC	FGNQSW
02415208	AURO-PANTOPRAZOLE	ARO	FGNQSW
02416565	MAR-PANTOPRAZOLE	MAR	FGNQSW
02417448	MINT-PANTOPRAZOLE	MNT	FGNQSW
02428180	PANTOPRAZOLE	SIV	FGNQSW
02467372	M-PANTOPRAZOLE SODIUM	MRA	FGNQSW
02471825	NRA-PANTOPRAZOLE	NRA	FGNQSW
02481588	AG-PANTOPRAZOLE SODIUM	ANG	FGNQSW

**RABEPRAZOLE SODIUM**

[SEE PROTON PUMP INHIBITORS IN APPENDIX A](#) FOR SA CRITERIA **FOR DOSAGES GREATER THAN 20MG/DAY**

10MG ENTERIC COATED TABLET

02243796	PARIET	JAN	FNQSW
02314177	SANDOZ-RABEPRAZOLE	SDZ	FGNQSW
02345579	APO-RABEPRAZOLE	APX	FGNQSW
02356511	RABEPRAZOLE	SNS	FGNQSW
02385449	RABEPRAZOLE	SIV	FGNQSW
02415283	JAMP-RABEPRAZOLE	JPC	FGNQSW

20MG ENTERIC COATED TABLET

02243797	PARIET	JAN	FNQSW
02314185	SANDOZ-RABEPRAZOLE	SDZ	FGNQSW
02356538	RABEPRAZOLE	SNS	FGNQSW
02385457	RABEPRAZOLE	SIV	FGNQSW
02415291	JAMP-RABEPRAZOLE	JPC	FGNQSW

**56:32.00 MISCELLANEOUS G.I. DRUGS**

**DOMPERIDONE MALEATE**

10MG TABLET

01912070	TEVA-DOMPERIDONE	TEV	FGNQSW
02103613	APO-DOMPERIDONE	APX	FGNQSW
02238341	DOMPERIDONE	SIV	FGNQSW
02268078	RAN-DOMPERIDONE	RAN	FGNQSW
02350440	DOMPERIDONE	SNS	FGNQSW
02369206	JAMP-DOMPERIDONE	JPC	FGNQSW
02403870	MAR-DOMPERIDONE	MAR	FGNQSW
02462834	PRZ-DOMPERIDONE	PRZ	FGNQSW

**METOCLOPRAMIDE HCL**

5MG TABLET

02230431	PMS-METOCLOPRAMIDE	PMS	FGNQSW
02517795	MAR-METOCLOPRAMIDE	MAR	FGNQSW

10MG TABLET

02230432	PMS-METOCLOPRAMIDE	PMS	FGNQSW
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1MG/ML ORAL SOLUTION

02230433	PMS-METOCLOPRAMIDE	PMS	FGNQSW
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5MG/ML INJECTION SOLUTION (2ML)

02185431	METOCLOPRAMIDE HCL	SDZ	NQ
02243563	METOCLOPRAMIDE	OMG	NQ
02537397	METOCLOPRAMIDE HCL	JPC	NQ

**OBETICHOLIC**SEE [APPENDIX A](#) FOR SA CRITERIA

5MG TABLET

02463121 OCALIVA (SA) INT NMQW

10MG TABLET

02463148 OCALIVA (SA) INT NMQW

**SULFASALAZINE**

500MG ENTERIC COATED TABLET

00598488 PMS-SULFASALAZINE E.C. PMS FGNQSW  
02064472 SALAZOPYRIN PFI FNQSW

500MG TABLET

00598461 PMS-SULFASALAZINE PMS FGNQSW  
02064480 SALAZOPYRIN PFI FNQSW**TRIMEBUTINE MALEATE**

100MG TABLET

02245663 APO-TRIMEBUTINE APX FGNQSW  
02538202 MINT-TRIMEBUTINE MNT FGNQSW

200MG TABLET

02245664 APO-TRIMEBUTINE APX FGNQSW  
02538210 MINT-TRIMEBUTINE MNT FGNQSW**56:36.00 ANTI-INFLAMMATORY AGENTS**

5-AMINOSALICYLIC ACID (MESALAZINE)

400MG ENTERIC COATED TABLET

02171929 TEVA-5 ASA TEV FGNQSW

500MG ENTERIC COATED TABLET

02112787 SALOFALK ALL FNQSW

500MG DELAYED RELEASE TABLET

02099683 PENTASA FEI FNQSW

1G EXTENDED RELEASE TABLET

02399466 PENTASA FEI FNQSW

500MG RECTAL SUPPOSITORY

02112760	SALOFALK	APT	<b>FNQSW</b>
1G/ACTUATION RECTAL FOAM			
02474026	MEZERA	AVI	<b>FNQSW</b>
1G RECTAL SUPPOSITORY			
02242146	SALOFALK	APT	<b>FNQSW</b>
02153564	PENTASA	FEI	<b>FNQSW</b>
02474018	MEZERA	AVI	<b>FNQSW</b>
1G/100ML RECTAL ENEMA			
02153521	PENTASA	FEI	<b>FNQSW</b>
4G/100ML RECTAL ENEMA			
02153556	PENTASA	FEI	<b>FNQSW</b>
2G/60G RETENTION ENEMA (60G)			
02112795	SALOFALK	APT	<b>FNQSW</b>
4G/60G RETENTION ENEMA (60G)			
02112809	SALOFALK	APT	<b>FNQSW</b>
<b>BETAMETHASONE DISODIUM PHOSPHATE</b>			
5MG/100ML ENEMA (100ML)			
02060884	BETNESOL	PAL	<b>FNQSW</b>
<b>MESALAMINE</b>			
500MG TABLET			
02524481	MEZARA	AVI	<b>FNQSW</b>
1.2GM DELAYED RELEASE TABLET			
02297558	MEZAVANT	SHI	<b>FNQSW</b>
<b>OLSALAZINE SODIUM</b>			
250MG CAPSULE			
02063808	DIPENTUM	ATN	<b>FNQSW</b>

## **64:00.00 HEAVY METAL ANTAGONISTS**

### **DEFERASIROX**

[SEE APPENDIX A](#) FOR SA CRITERIA

90MG TABLET

02485265	APO-DEFERASIROX (TYPE J)	APX	<b>NMQW</b>
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02489899	SANDOZ-DEFERASIROX (TYPE J)	SDZ	<b>NMQW</b>
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02507315	TARO-DEFERASIROX (TYPE J)	TAR	NMQW
02528290	PMS-DEFERASIROX (TYPE J)	PMS	NMQW
180MG TABLET			
02485273	APO-DEFERASIROX (TYPE J)	APX	NMQW
02489902	SANDOZ-DEFERASIROX (TYPE J)	SDZ	NMQW
02507323	TARO-DEFERASIROX (TYPE J)	TAR	NMQW
02528304	PMS-DEFERASIROX (TYPE J)	PMS	NMQW
360MG TABLET			
02485281	APO-DEFERASIROX (TYPE J)	APX	NMQW
02489910	SANDOZ-DEFERASIROX (TYPE J)	SDZ	NMQW
02507331	TARO-DEFERASIROX (TYPE J)	TAR	NMQW
02528312	PMS-DEFERASIROX (TYPE J)	PMS	NMQW
125MG DISPERSIBLE TABLETS			
02287420	EXJADE (SA)	NVR	NMQW
02461544	APO-DEFERASIROX (SA)	APX	NMQW
02464454	SANDOZ-DEFERASIROX (SA)	SDZ	NMQW
250MG DISPERSIBLE TABLETS			
02287439	EXJADE (SA)	NVR	NMQW
02461552	APO-DEFERASIROX (SA)	APX	NMQW
02464462	SANDOZ-DEFERASIROX (SA)	SDZ	NMQW
500MG DISPERSIBLE TABLETS			
02287447	EXJADE (SA)	NVR	NMQW
02461560	APO-DEFERASIROX (SA)	APX	NMQW
02464470	SANDOZ-DEFERASIROX (SA)	SDZ	NMQW
<b>PENICILLAMINE</b>			
250MG CAPSULE			
00016055	CUPRIMINE	VAL	FNQSW

## **68:04.00 CORTICOSTEROIDS**

### **BECLOMETHASONE DIPROPIONATE**

50MCG/INHALER			
02242029	QVAR	VAL	FNQSW
100MCG/INHALER			
02242030	QVAR	VAL	FNQSW

**BETAMETHASONE**

6MG/ML VIAL

00028096

CELESTONE SOLUSPAN

ORG **FNQSW****BUDESONIDE**

100MCG/DOSE INHALER POWDER (200 DOSE)

00852074

PULMICORT TURBUHALER

AZE **FQW**

200MCG/DOSE INHALER POWDER (200 DOSE)

00851752

PULMICORT TURBUHALER

AZE **CFNQSW**

400MCG/DOSE INHALER POWDER (200 DOSE)

00851760

PULMICORT TURBUHALER

AZE **CFNQSW**

[SEE APPENDIX A](#) FOR SA CRITERIA (NURSING HOME PROGRAM AND CYSTIC FIBROSIS PROGRAM DOES NOT REQUIRE AN SA REQUEST)

0.125MG/ML INHALATION SOLUTION (2ML)

02229099

PULMICORT NEBUAMP (SA)

AZE **CFNQSW**

02465949

TEVA-BUDESONIDE (SA)

TEV **CFGNQSW**

02494264

TARO-BUDESONIDE (SA)

TAR **CFGNQSW**

0.25MG/ML INHALATION SOLUTION (2ML)

01978918

PULMICORT NEBUAMP (SA)

AZE **CFNQSW**

02494272

TARO-BUDESONIDE (SA)

TAR **CFGNQSW**

0.5MG/ML INHALATION SOLUTION (2ML)

01978926

PULMICORT NEBUAMP (SA)

AZE **CFNQSW**

02465957

TEVA-BUDESONIDE (SA)

TEV **CFGNQSW**

02494280

TARO-BUDESONDE (SA)

TAR **CFGNQSW**

2MG/ACTUATION RECTAL FOAM

02498057

UCERIS

BLO **FNQSW**

3MG DELAYED AND EXTENDED RELEASE CAPSULE

02229293

ENTOCORT

TPG **FNQSW**

0.02MG/ML RECTAL ENEMA

02052431

ENTOCORT

TPG **FNQSW****CICLESONIDE**

100MCG/DOSE INHALATION AEROSOL

02285606

ALVESCO

COV **FGNQSW**

200MCG/DOSE INHALATION AEROSOL

02285614

ALVESCO

COV **FGNQSW**

**CORTISONE ACETATE**

25MG TABLET

00280437 CORTISONE

VAL **CFNQSW****DEXAMETHASONE**

0.5MG TABLET

01964976 PMS-DEXAMETHASONE

PMS **CFGNQSW**

02261081 APO-DEXAMETHASONE

APX **CFGNQSW**

0.75MG TABLET

01964968 PMS-DEXAMETHASONE

PMS **FGNQSW**

2MG TABLET

02279363 PMS-DEXAMETHASONE

PMS **FGNQSW**

4MG TABLET

01964070 PMS-DEXAMETHASONE

PMS **CFGNQSW**

02250055 APO-DEXAMETHASONE

APX **CFGNQSW****DEXAMETHASONE 21 PHOSPHATE**

4MG/ML INJECTION SOLUTION (5ML)

00664227 DEXAMETHASONE

SDZ **FGNQSW**

01977547 DEXAMETHASONE

STE **FGNQSW****FLUDROCORTISONE ACETATE**

0.1MG TABLET

02086026 FLORINEF

PAL **FNQSW****FLUTICASONE FUROATE**

100MCG POWDER FOR INHALATION

02446561 ARNUITY ELLIPTA

GSK **FNQSW**

200MCG POWDER FOR INHALATION

02446588 ARNUITY ELLIPTA

GSK **FNQSW****FLUTICASONE PROPIONATE**

55MCG/DOSE AEROSOL POWDER

02467895 AERMONY RESPICLICK

TEV **FGNQSW**

113MCG/DOSE AEROSOL POWDER

02467909 AERMONY RESPICLICK

TEV **FGNQSW**

232MCG/DOSE AEROSOL POWDER

02467917 AERMONY RESPICLICK

TEV **FGNQSW**

50MCG/DOSE INHALATION AEROSOL HYDROFLUOROALKANE (HFA)



02244291	FLOVENT HFA	GSK	CFNQSW
125MCG/DOSE INHALATION AEROSOL HYDROFLUOROALKANE (HFA)			
02244292	FLOVENT HFA	GSK	CFNQSW
02503123	PMS-FLUTICASONE HFA	PMS	CFGNQSW
02526557	APO-FLUTICASONE HFA	APX	CFGNQSW
250MCG/DOSE INHALATION AEROSOL HYDROFLUOROALKANE (HFA)			
02244293	FLOVENT HFA	GSK	CFNQSW
02503131	PMS-FLUTICASONE HFA	PMS	CFGNQSW
02510987	APO-FLUTICASONE HFA	APX	CFGNQSW
100MCG/DOSE AEROSOL POWDER DISK (60)			
02237245	FLOVENT DISKUS	GSK	FNQSW
250MCG/DOSE AEROSOL POWDER DISK (60)			
02237246	FLOVENT DISKUS	GSK	FNQSW
500MCG/DOSE AEROSOL POWDER DISK (60)			
02237247	FLOVENT DISKUS	GSK	FNQSW
<b>HYDROCORTISONE</b>			
10MG TABLET			
00030910	CORTEF	PFI	CFNQSW
02524465	AURO-HYDROCORTISONE	ARO	CFGNQSW
20MG TABLET			
00030929	CORTEF	PFI	CFNQSW
02524473	AURO-HYDROCORTISONE	ARO	CFGNQSW
<b>HYDROCORTISONE SODIUM SUCCINATE</b>			
250MG INJECTION POWDER			
00030619	SOLU-CORTEF	PFI	NQ
<b>METHYLPREDNISOLONE</b>			
4MG TABLET			
00030988	MEDROL	PFI	CFNQSW
16MG TABLET			
00036129	MEDROL	PFI	FNQSW
<b>METHYLPREDNISOLONE ACETATE</b>			
40MG/ML INJECTION SUSPENSION (1ML)			
00030759	DEPO MEDROL	PFI	FNQSW
40MG/ML INJECTION SUSPENSION (2ML)			

01934333	DEPO MEDROL	PFI	<b>FNQSW</b>
80MG/ML INJECTION SUSPENSION (1ML)			
00030767	DEPO MEDROL	PFI	<b>FNQSW</b>
<b>MOMETASONE FUROATE</b>			
100MCG DRY POWDER INHALER			
02438690	ASMANEX	ORG	<b>FNQSW</b>
200MCG DRY POWDER INHALER			
02243595	ASMANEX	ORG	<b>FNQSW</b>
400MCG DRY POWDER INHALER			
02243596	ASMANEX	ORG	<b>FNQSW</b>
<b>PREDNISOLONE SODIUM PHOSPHATE</b>			
1MG/ML ORAL LIQUID			
02245532	PMS-PREDNISOLONE	PMS	<b>CFGNQSW</b>
<b>PREDNISONE</b>			
1MG TABLET			
00271373	WINPRED	AAA	<b>CFNQSW</b>
5MG TABLET			
00021695	TEVA-PREDNISONE	TEV	<b>CFGNQSW</b>
00312770	APO-PREDNISONE	APX	<b>CFGNQSW</b>
50MG TABLET			
00232378	TEVA-PREDNISONE	TEV	<b>CFGNQSW</b>
00550957	APO-PREDNISONE	APX	<b>CFGNQSW</b>
<b>TRIAMCINOLONE ACETONIDE</b>			
10MG/ML VIAL			
01999761	KENALOG-10	BMS	<b>FNQSW</b>
40MG/ML VIAL			
01999869	KENALOG-40	BMS	<b>FNQSW</b>
01977563	TRIAMCINOLONE ACETONIDE	STE	<b>FNQSW</b>
<b>TRIAMCINOLONE HEXACETONIDE</b>			
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
20MG/ML AMPULE			
02470632	TRISPAN (SA)	MED	<b>FWQ</b>

## **68:08.00 ANDROGENS**

### **DANAZOL**

50MG CAPSULE			
02018144	CYCLOMEN	AVN	<b>FQW</b>
100MG CAPSULE			
02018152	CYCLOMEN	AVN	<b>FQW</b>
200MG CAPSULE			
02018160	CYCLOMEN	AVN	<b>FQW</b>

### **TESTOSTERONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

25MG/2.5GM TRANSDERMAL GEL			
02245345	ANDROGEL (SA)	BGP	<b>FNQSW</b>
02463792	TARO-TESTOSTERONE (SA)	TAR	<b>FNQSW</b>
50MG/5GM TRANSDERMAL GEL			
02245346	ANDROGEL (SA)	BGP	<b>FNQSW</b>
02463806	TARO-TESTOSTERONE (SA)	TAR	<b>FNQSW</b>
50MG/5GM TRANSDERMAL GEL			
02245346	ANDROGEL (SA)	BGP	<b>FNQSW</b>
02280248	TESTIM (SA)	PAL	<b>FNQSW</b>

### **TESTOSTERONE CYPIONATE**

100MG/ML VIAL			
00030783	DEPO-TESTOSTERONE	PFI	<b>FQW</b>
02496003	TARO-TESTOSTERONE	TAR	<b>FQW</b>

### **TESTOSTERONE UNDECANOATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

40MG CAPSULE			
02322498	PMS-TESTOSTERONE (SA)	PMS	<b>FNQSW</b>
02421186	TARO-TESTOSTERONE (SA)	TAR	<b>FNQSW</b>

## **68:12.00 CONTRACEPTIVES**

### **ESTRADIOL & ETONOGESTREL**

2MG & 11.4MG VAGINAL INSERT

02253186	NUVARING	MSD	FQW
02520028	HALOETTE	SLP	FGQW

**ETHINYL ESTRADIOL & DESOGESTREL**

0.025MG & 0.10MG TABLET (21 DAY)			
02272903	LINESSA	ASN	FQW

0.025MG & 0.10MG TABLET (28 DAY)			
02257238	LINESSA	ASN	FQW

0.03MG & 0.15MG TABLET (21 DAY)			
02042487	MARVELON	MSD	FQW
02317192	APRI 21	TEV	FGQW
02396491	FREYA 21	MYL	FGQW
02410249	MIRVALA 21	APX	FGQW

0.03MG & 0.15MG TABLET (28 DAY)			
02042479	MARVELON	MSD	FQW
02317206	APRI 28	TEV	FGQW
02396610	FREYA 21	MYL	FGQW
02410257	MIRVALA 28	APX	FGQW

**ETHINYL ESTRADIOL & DROSPIRENONE**

3.0MG & 0.03MG TABLETS (21 DAY)			
02261723	YASMIN 21	BAY	FQW
02410788	ZAMINE 21	APX	FGQW
02421437	DROSPIRENONE-ETHINYL ESTRADIOL	GLN	FGQW

3.0MG & 0.03MG TABLETS (28 DAY)			
02261731	YASMIN 28	BAY	FQW
02410796	ZAMINE 28	APX	FGQW
02421445	DROSPIRENONE-ETHINYL ESTRADIOL	GLN	FGQW

**ETHINYL ESTRADIOL & L-NORGESTREL**

0.2MG & 0.1MG TABLET (21 DAY)			
02236974	ALESSE	PFI	FQW
02298538	AVIANE 21	TEV	FGQW
02387875	ALYSENA	APX	FGQW
02532174	AUDRINA (21)	JPC	FGQW

0.2MG & 0.1MG TABLET (28 DAY)			
02236975	ALESSE	PFI	FQW
02298546	AVIANE 28	TEV	FGQW
02387883	ALYSENA	APX	FGQW
02532182	AUDRINA (28)	JPC	FGQW

0.03MG & 0.05MG (6); 0.04MG & 0.075MG (5); 0.03MG & 0.125MG (10) TABLET (21 DAY)  
00707600 TRIQUILAR BAY **FQW**

0.03MG & 0.05MG (6); 0.04MG & 0.075MG (5); 0.03MG & 0.125MG (10); INERT TABLETS  
(7) TABLET (28 DAY)  
00707503 TRIQUILAR BAY **FQW**

0.03MG & 0.15MG TABLET (21 DAY)  
02042320 MIN-OVRAL PFI **FQW**  
02295946 PORTIA TEV **FGQW**  
02387085 OVIMA APX **FGQW**

0.03MG & 0.15MG TABLET (28 DAY)  
02042339 MIN-OVRAL PFI **FQW**  
02295954 PORTIA TEV **FGQW**  
02387093 OVIMA APX **FGQW**

**ETHINYL ESTRADIOL & NORETHINDRONE**

0.035MG & 0.5MG TABLET (21 DAY)  
02187086 BREVICON PFI **FQW**

0.035MG & 0.5MG TABLET (28 DAY)  
02187094 BREVICON PFI **FQW**

0.035MG & 0.5MG (7); 0.035MG & 1.0MG (7); 0.035MG & 0.5MG (7) TABLET (21 DAY)  
02187108 SYNPHASIC PFI **FQW**

0.035MG & 0.5MG (7); 0.035MG & 1.0MG (7); 0.035MG & 0.5MG (7); INERT TABLETS (7)  
TABLET (28 DAY)  
02187116 SYNPHASIC PFI **FQW**

0.035MG & 1.0MG TABLET (21 DAY)  
02189054 BREVICON 1/35 PFI **FQW**  
02197502 SELECT 1/35 PFI **FQW**

0.035MG & 1.0MG TABLET (28DAY)  
02189062 BREVICON 1/35 PFI **FQW**  
02199297 SELECT 1/35 PFI **FQW**

**ETHINYL ESTRADIOL & NORGESTIMATE**

0.025MG & 0.180MG (7); 0.025MG & 0.215MG (7); 0.025MG & 0.250MG (7) TABLET (21  
DAY)  
02401967 TRICIRA LO APX **FGQW**

0.025MG & 0.180MG (7); 0.025MG & 0.215MG (7); 0.025MG & 0.250MG (7); INERT  
TABLETS (7) TABLET (28 DAY)

02401975 TRICIRA LO APX FGQW

0.035MG & 0.180MG (7); 0.035MG & 0.215MG (7); 0.035MG & 0.250MG (7) TABLET  
(21 DAY)

02486296 TRI-JORDYNA 21 GLM FGQW  
02508087 TRI-CIRA 21 APX FGQW

0.035MG & 0.180MG (7); 0.035MG & 0.215MG (7); 0.035MG & 0.250MG (7); TABLET  
(28 DAY)

02486318 TRI-JORDYNA 28 GLM FGQW  
02508095 TRI-CIRA 28 APX FGQW

**ETONOGESTREL**

68MG SC IMPLANT

02499509 NEXPLANON ORG FQW

**LEVONORGESTROL**

1.5MG TABLET

02293854 PLAN B PAL FQW  
02433532 BACKUP PLAN ONESTEP APX FGQW

19.5MG INTRAUTERINE SYSTEM

02459523 KYLEENA BAY FQW

52MG INTRAUTERINE SYSTEM

02243005 MIRENA BAY FQW

**NORETHINDRONE**

0.35MG TABLET (28 DAY)

02410303 MOVISSE MYL FGQW  
02441306 JENCYCLA LUP FGQW

**68:16.00 ESTROGENS**

**CONJUGATED ESTROGENS**

0.3MG TABLET

02414678 PREMARIN PFI FNQSW

0.625MG TABLET

02414686 PREMARIN PFI FNQSW

1.25MG TABLET

02414694 PREMARIN PFI FNQSW

0.625MG/G VAGINAL CREAM				
02043440	PREMARIN		PFI	<b>FNQSW</b>
<b>ESTRADIOL</b>				
0.5MG TABLET				
02225190	ESTRACE		ACS	<b>FNQSW</b>
02449048	LUPIN-ESTRADIOL		LUP	<b>FGNQSW</b>
1MG TABLET				
02148587	ESTRACE		ACS	<b>FNQSW</b>
02449056	LUPIN-ESTRADIOL		LUP	<b>FGNQSW</b>
2MG TABLET				
02148595	ESTRACE		ACS	<b>FNQSW</b>
02449064	LUPIN-ESTRADIOL		LUP	<b>FGNQSW</b>
25MCG TRANSERMAL PATCH				
02245676	ESTRADOT		SDZ	<b>FNQSW</b>
50MCG TRANSDERMAL PATCH				
02244000	ESTRADOT		SDZ	<b>FNQSW</b>
02246967	SANDOZ-ESTRADIOL DERM		SDZ	<b>FGNQSW</b>
75MCG TRANSDERMAL PATCH				
02244001	ESTRADOT		SDZ	<b>FNQSW</b>
02246968	SANDOZ-ESTRADIOL DERM		SDZ	<b>FGNQSW</b>
100MCG TRANSDERMAL PATCH				
02244002	ESTRADOT		SDZ	<b>FNQSW</b>
02246969	SANDOZ-ESTRADIOL DERM		SDZ	<b>FGNQSW</b>
7.5MCG/24 HOUR VAGINAL RING				
02168898	ESTRING		PAL	<b>FNQSW</b>
4MCG VAGINAL INSERT				
02503689	IMVEXXY		KNI	<b>FNQSW</b>
10MCG VAGINAL INSERT				
02503697	IMVEXXY		KNI	<b>FNQSW</b>
10MCG VAGINAL TABLET				
02325462	VAGIFEM		NNO	<b>FNQSW</b>
0.06% TOPICAL GEL				
02238704	ESTROGEL		ORG	<b>FNQSW</b>

**ESTRADIOL & NORETHINRONE ACETATE**

50MCG & 140MCG TRANSDERMAL PATCH  
02241835            ESTALIS

SDZ    **FNQSW**

50MCG & 250MCG TRANSDERMAL PATCH  
02241837            ESTALIS

SDZ    **FNQSW**

**ESTRADIOL & PROGESTERONE**

1 MG & 100 MG    CAPSULE  
02505223            BIJUVA

KNI    **FQW**

**ESTRONE**

0.1% VAGINAL CREAM  
00727369            ESTRAGYN

SLP    **FNQSW**

**68:18.00 GONADOTROPINS**

**GOSERELIN ACETATE**

3.6MG DEPOT INJECTION  
02049325            ZOLADEX

TRT    **FNQSW**

10.8MG DEPOT INJECTION  
02225905            ZOLADEX LA

TRT    **FNQSW**

**68:20.00 ANTIDIABETIC DRUGS (ORAL HYPOGLYCEMICS)**

**⑤ACARBOSE**

50MG TABLET  
02493780            ACARBOSE  
02494078            MAR-ACARBOSE

STR    **DNQW**  
MAR    **DNQW**

100MG TABLET  
02493799            ACARBOSE  
02494086            MAR-ACARBOSE

STR    **DNQW**  
MAR    **DNQW**

**⑤CANAGLIFLOZIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG TABLET  
02425483            INVOKANA (SA)

JAN    **DNQW**

300MG TABLET



02425491 INVOKANA (SA) JAN DNQW

**⑤DAPAGLIFLOZIN**

**(Note: \$5 copay only applies to use in diabetes)**

5MG TABLET

02435462	FORXIGA	AZE	DFNQSW
02518732	SANDOZ-DAPAGLIFLOZIN	SDZ	DFGNQSW
02519852	GLN-DAPAGLIFLOZIN	GLN	DFGNQSW
02527189	APO-DAPAGLIFLOZIN	APX	DFGNQSW
02531364	JAMP-DAPAGLIFLOZIN	JPC	DFGNQSW
02531402	AURO-DAPAGLIFLOZIN	ARO	DFGNQSW
02531550	PMS-DAPAGLIFLOZIN	PMS	DFGNQSW
02535297	M-DAPAGLIFLOZIN	MRA	DFGNQSW

10MG TABLET

02435470	FORXIGA	AZE	DFNQSW
02518740	SANDOZ-DAPAGLIFLOZIN	SDZ	DFGNQSW
02519860	GLN-DAPAGLIFLOZIN	GLN	DFGNQSW
02527197	APO-DAPAGLIFLOZIN	APX	DFGNQSW
02531372	JAMP-DAPAGLIFLOZIN	JPC	DFGNQSW
02531410	AURO-DAPAGLIFLOZIN	ARO	DFGNQSW
02531569	PMS-DAPAGLIFLOZIN	PMS	DFGNQSW
02535300	M-DAPAGLIFLOZIN	MRA	DFGNQSW

**⑤DAPAGLIFLOZIN & METFORMIN HYDROCHLORIDE**

5MG/850MG TABLET

02449935	XIGDUO	AZE	DNQW
02533073	AURO-DAPAGLIFLOZIN-METFORMIN	ARO	DNQW
02536153	APO-DAPAGLIFLOZIN-METFORMIN	APX	DNQW

5MG/1000MG TABLET

02449943	XIGDUO	AZE	DNQW
02533081	AURO-DAPAGLIFLOZIN-METFORMIN	ARO	DNQW
02536161	APO-DAPAGLIFLOZIN-METFORMIN	APX	DNQW

**⑤EMPAGLIFLOZIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET

02443937	JARDIANCE (SA)	BOE	DNQW
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25MG TABLET

02443945	JARDIANCE (SA)	BOE	DNQW
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**⑤EMPAGLIFLOZIN & METFORMIN HCL**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG & 500MG TABLET

02456575	SYNJARDY (SA)	BOE	<b>DNQW</b>
5MG & 850MG TABLET			
02456583	SYNJARDY (SA)	BOE	<b>DNQW</b>
5MG & 1000MG TABLET			
02456591	SYNJARDY (SA)	BOE	<b>DNQW</b>
12.5MG & 500MG TABLET			
02456605	SYNJARDY (SA)	BOE	<b>DNQW</b>
12.5MG & 850MG TABLET			
02456613	SYNJARDY (SA)	BOE	<b>DNQW</b>
12.5MG & 1000MG TABLET			
02456621	SYNJARDY (SA)	BOE	<b>DNQW</b>

**⑤ GLICLAZIDE**

30MG MODIFIED RELEASE TABLET			
02242987	DIAMICRON MR	SEV	<b>DNQW</b>
02297795	APO-GLICLAZIDE MR	APX	<b>DNQW</b>
02423286	MINT-GLICLAZIDE MR	MNT	<b>DNQW</b>
02438658	MYLAN-GLICLAZIDE MR	MYL	<b>DNQW</b>
02461323	SANDOZ-GLICLAZIDE MR	SDZ	<b>DNQW</b>
02463571	TARO-GLICLAZIDE MR	SUN	<b>DNQW</b>
02524856	GLICLAZIDE MR	SNS	<b>DNQW</b>

80MG TABLET			
02238103	TEVA-GLICLAZIDE	TEV	<b>DNQW</b>
02245247	APO-GLICLAZIDE	APX	<b>DNQW</b>
02287072	GLICLAZIDE	SNS	<b>DNQW</b>

60MG EXTENDED RELEASE TABLET			
02356422	DIAMICRON MR	SEV	<b>DNQW</b>
02407124	APO-GLICLAZIDE MR	APZ	<b>DNQW</b>
02423294	MINT-GLICLAZIDE MR	MNT	<b>DNQW</b>
02439328	TARO-GLICLAZINE MR	SUN	<b>DNQW</b>
02461331	SANDOZ-GLICLAZIDE MR	SDZ	<b>DNQW</b>
02524864	GLICLAZIDE MR	SNS	<b>DNQW</b>

**GLIMEPIRIDE**

1MG TABLET			
02269589	SANDOZ-GLIMEPIRIDE	SDZ	<b>DNQW</b>
2MG TABLET			
02269597	SANDOZ-GLIMEPIRIDE	SDZ	<b>DNQW</b>

4MG TABLET			
02269619	SANDOZ-GLIMEPIRIDE	SDZ	DNQW

**⑤ GLYBURIDE**

2.5MG TABLET			
01913654	APO-GLYBURIDE	APX	DNQW
01913670	TEVA-GLYBURIDE	TEV	DNQW
02350459	GLYBURIDE	SNS	DNQW

5MG TABLET			
01913662	APO-GLYBURIDE	APX	DNQW
01913689	TEVA-GLYBURIDE	TEV	DNQW
02350467	GLYBURIDE	SNS	DNQW

**⑤ LINAGLIPTIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG TABLET			
02370921	TRAJENTA (SA)	BOE	DNQW

**⑤ LINAGLIPTIN & METFORMIN HYDROCHLORIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

2.5MG/500MG TABLET			
02403250	JENTADUETO (SA)	BOE	DNQW

2.5MG/850MG TABLET			
02403269	JENTADUETO (SA)	BOE	DNQW

2.5MG/1000MG TABLET			
02403277	JENTADUETO (SA)	BOE	DNQW

**⑤ METFORMIN**

500MG TABLET			
02099233	GLUCOPHAGE	AVN	DNQW
02223562	PMS-METFORMIN	PMS	DNQW
02246820	SANDOZ-METFORMIN FC	SDZ	DNQW
02257726	TEVA-METFORMIN	TEV	DNQW
02269031	RAN-METFORMIN	RAN	DNQW
02353377	METFORMIN	SNS	DNQW
02378620	MAR-METFORMIN	MAR	DNQW
02380196	JAMP-METFORMIN	JPC	DNQW
02385341	METFORMIN	SIV	DNQW
02388766	MINT-METFORMIN	MNT	DNQW
02438275	AURO-METFORMIN	ARO	DNQW
02520303	PMSC-METFORMIN	PMS	DNQW
02531895	PRZ-METFORMIN	PRZ	DNQW

850MG TABLET

02162849	GLUCOPHAGE	AVN	DNQW
02242589	PMS-METFORMIN	PMS	DNQW
02246821	SANDOZ-METFORMIN FC	SDZ	DNQW
02257734	TEVA-METFORMIN	TEV	DNQW
02269058	RAN-METFORMIN	RAN	DNQW
02353385	METFORMIN	SNS	DNQW
02378639	MAR-METFORMIN	MAR	DNQW
02380218	JAMP-METFORMIN	JPC	DNQW
02385368	METFORMIN	SIV	DNQW
02388774	MINT-METFORMIN	MNT	DNQW
02438283	AURO-METFORMIN	ARO	DNQW
02520311	PMSC-METFORMIN	PMS	DNQW
02531909	PRZ-METFORMIN	PRZ	DNQW

**PIOGLITAZONE HCL**

15MG TABLET

02297906	SANDOZ-PIOGLITAZONE	SDZ	DNQW
02302861	ACT-PIOGLITAZONE	TEV	DNQW
02302942	APO-PIOGLITAZONE	APX	DNQW
02326477	MINT-PIOGLICAZONE	MNT	DNQW
02375850	RAN-PIOGLITAZONE	RAN	DNQW
02391600	PIOGLITAZONE HYDROCHLORIDE	ACH	DNQW
02397307	JAMP-PIOGLITAZONE	JPC	DNQW

30MG TABLET

02297914	SANDOZ-PIOGLITAZONE	SDZ	DNQW
02302888	ACT-PIOGLITAZONE	TEV	DNQW
02302950	APO-PIOGLITAZONE	APX	DNQW
02326485	MINT-PIOGLITAZONE	MNT	DNQW
02339587	PIOGLITAZONE	ACH	DNQW
02365529	JAMP-PIOGLITAZONE	JPC	DNQW
02375869	RAN-PIOGLITAZONE	RAN	DNQW

45MG TABLET

02297922	SANDOZ-PIOGLITAZONE	SDZ	DNQW
02302896	ACT-PIOGLITAZONE	TEV	DNQW
02302977	APO-PIOGLITAZONE	APX	DNQW
02326493	MINT-PIOGLITAZONE	MNT	DNQW
02339595	PIOGLITAZONE	ACH	DNQW
02365537	JAMP-PIOGLITAZONE	JPC	DNQW
02375877	RAN-PIOGLITAZONE	RAN	DNQW

**⑤SAXAGLIPTIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

2.5MG TABLET

02375842	ONGLYZA (SA)	AZE	DNQW
02468603	SANDOZ-SAXAGLIPTIN (SA)	SDZ	DNQW
02507471	APO-SAXAGLIPTIN (SA)	APX	DNQW

5MG TABLET

02333554	ONGLYZA (SA)	AZE	DNQW
02468611	SANDOZ-SAXAGLIPTIN (SA)	SDZ	DNQW
02507498	APO-SAXAGLIPTIN (SA)	APX	DNQW

⑤ **SAXAGLIPTIN & METFORMIN HYDROCHLORIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

2.5MG/500MG TABLET

02389169	KOMBOGLYZE (SA)	AZE	DNQW
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2.5MG/850MG TABLET

02389177	KOMBOGLYZE (SA)	AZE	DNQW
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2.5MG/1000MG TABLET

02389185	KOMBOGLYZE (SA)	AZE	DNQW
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⑤ **SITAGLIPTIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

25MG TABLET

02388839	JANUVIA (SA)	MSD	DNQW
02504049	SANDOZ-SITAGLIPTIN (SA)	SDZ	DNQW
02508656	APO-SITAGLIPTIN MALATE (SA)	APX	DNQW
02512475	ACH-SITAGLIPTIN (SA)	ACH	DNQW
02522705	TEVA-SITAGLIPTIN MALATE (SA)	TEV	DNQW
02529033	SITAGLIPTIN (SA)	SIV	DNQW
02529866	AURO-SITAGLIPTIN HCL (SA)	ARO	DNQW
02531631	TARO-SITAGLIPTIN FUMERATE (SA)	TAR	DNQW
02534134	JAMP SITAGLIPTIN (SA)	JPC	DNQW

50MG TABLET

02388847	JANUVIA (SA)	MSD	DNQW
02504057	SANDOZ-SITAGLIPTIN (SA)	SDZ	DNQW
02508664	APO-SITAGLIPTIN MALATE (SA)	APX	DNQW
02512483	ACH-SITAGLIPTIN (SA)	ACH	DNQW
02522713	TEVA-SITAGLIPTIN MALATE (SA)	TEV	DNQW
02529041	SITAGLIPTIN (SA)	SIV	DNQW
02529874	AURO-SITAGLIPTIN HCL (SA)	ARO	DNQW
02531658	TARO-SITAGLIPTIN FUMERATE (SA)	TAR	DNQW
02534142	JAMP SITAGLIPTIN (SA)	JPC	DNQW

100MG TABLET

02303922	JANUVIA (SA)	MSD	<b>DNQW</b>
02504065	SANDOZ-SITAGLIPTIN (SA)	SDZ	<b>DNQW</b>
02508672	APO-SITAGLIPTIN MALATE (SA)	APX	<b>DNQW</b>
02512491	ACH-SITAGLIPTIN (SA)	ACH	<b>DNQW</b>
02522721	TEVA-SITAGLIPTIN MALATE (SA)	TEV	<b>DNQW</b>
02529068	SITAGLIPTIN (SA)	SIV	<b>DNQW</b>
02529882	AURO-SITAGLIPTIN HCL (SA)	ARO	<b>DNQW</b>
02531666	TARO-SITAGLIPTIN FUMERATE (SA)	TAR	<b>DNQW</b>
02534150	JAMP SITAGLIPTIN (SA)	JPC	<b>DNQW</b>

**⑤ SITAGLIPTIN & METFORMIN HYDROCHLORIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**50MG/500MG TABLET**

02333856	JANUMET (SA)	MSD	<b>DNQW</b>
02503956	SANDOZ SITAGLIPTIN-METFORMIN (SA)	SDZ	<b>DNQW</b>
02509415	APO-SITAGLIPTIN MAL-METFORMIN (SA)	APX	<b>DNQW</b>

**50MG/850MG TABLET.**

02333864	JANUMET (SA)	MSD	<b>DNQW</b>
02503964	SANDOZ SITAGLIPTIN-METFORMIN (SA)	SDZ	<b>DNQW</b>
02509423	APO-SITAGLIPTIN MAL-METFORMIN (SA)	APX	<b>DNQW</b>

**50MG/1000MG TABLET**

02333872	JANUMET (SA)	MSD	<b>DNQW</b>
02503972	SANDOZ SITAGLIPTIN-METFORMIN (SA)	SDZ	<b>DNQW</b>
02509431	APO-SITAGLIPTIN MAL-METFORMIN (SA)	APX	<b>DNQW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA

**50MG/500MG EXTENDED RELEASE TABLET**

02416786	JANUMET XR (SA)	MSD	<b>DNQW</b>
02506270	APO-SITAGLIPTIN-METFORMIN XR	APX	<b>DNQW</b>
02529106	SANDOZ SITAGLIPTIN-METFORMN XR	SDZ	<b>DNQW</b>

**50MG/1000MG EXTENDED RELEASE TABLET**

02416794	JANUMET XR (SA)	MSD	<b>DNQW</b>
02506289	APO-SITAGLIPTIN-METFORMIN XR	APX	<b>DNQW</b>
02529114	SANDOZ SITAGLIPTIN-METFORMN XR	SDZ	<b>DNQW</b>

**100MG/1000MG EXTENDED RELEASE TABLET**

02416808	JANUMET XR (SA)	MSD	<b>DNQW</b>
02506297	APO-SITAGLIPTIN-METFORMIN XR	APX	<b>DNQW</b>
02529122	SANDOZ SITAGLIPTIN-METFORMN XR	SDZ	<b>DNQW</b>

**68:20.06 ANTIDIABETIC DRUGS (INCRETIN MIMETICS)**

**⑤ SEMAGLUTIDE**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.25-0.5MG PER DOSE (2MG/1.5ML) PEN INJECTOR  
02471477 OZEMPIC (SA) NNO **DNQW**

0.25-0.5MG PER DOSE (2MG/3ML) PEN INJECTOR  
02540258 OZEMPIC (SA) NNO **DNQW**

1MG/0.75ML (4MG/3ML) PEN INJECTOR  
02471469 OZEMPIC (SA) NNO **DNQW**

[SEE APPENDIX A](#) FOR SA CRITERIA

3MG TABLET  
02497581 RYBELSUS (SA) NNO **DNQW**

7MG TABLET  
02497603 RYBELSUS (SA) NNO **DNQW**

14MG TABLET  
02497611 RYBELSUS (SA) NNO **DNQW**

**68:20.08 ANTIDIABETIC DRUGS (INSULINS-HUMAN BIOSYNTHETIC)**

**⑤ INSULIN (DEGLUDEC)**

100 UNIT/ML PREFILLED PEN  
02467879 TRESIBA FLEXTOUCH NNO **DNQW**

100 UNIT/ML CARTRIDGE  
02467860 TRESIBA PENFILL NNO **DNQW**

200 UNIT/ML PREFILLED PEN  
02467887 TRESIBA FLEXTOUCH NNO **DNQW**

**INSULIN (DETEMIR)**

[SEE APPENDIX A](#) FOR SA CRITERIA

100 UNIT/ML CARTRIDGE  
02271842 LEVEMIR (SA) NNO **DNQW**

100 UNIT/ML PREFILLED PEN  
02412829 LEVEMIR FLEXTOUCH (SA) NNO **DNQW**

**⑤INSULIN (GLARGINE)**

100 UNIT/ML CARTRIDGE

02444844                BASAGLAR                                LIL    **DNQW**

100 UNIT/ML PREFILLED PEN (80 UNIT)

02461528                BASAGLAR                                LIL    **DNQW**

02526441                SEMGLEE                                BGP    **DNQW**

**SEE APPENDIX A FOR SA CRITERIA**

300 UNIT/ML PREFILLED PEN

02441829                TOUJEO SOLOSTAR (SA)                                AVN    **DNQW**

300 UNIT/ML PREFILLED PEN

02493373                TOUJEO DOUBLESTAR (SA)                                AVN    **DNQW**

**⑤INSULIN (GLULISINE)**

100 UNIT/ML CARTRIDGE

02279479                APIDRA    AVN    **DNQW**

100 UNIT/ML VIAL

02279460                APIDRA    AVN    **DNQW**

100 UNIT/ML PREFILLED PEN

02294346                APIDRA    AVN    **DNQW**

**⑤INSULIN (REGULAR) ASPART**

100IU/ML INJECTION SOLUTION (10ML)

02529254                TRURAPI\*    AVN    **DNQW**

\*Please ensure pump compatibility before switching to a biosimilar insulin

**SEE APPENDIX A FOR SA CRITERIA**

100IU/ML INJECTION SOLUTION (10ML)

02245397                NOVORAPID (SA)    NNO    **DNQW**

100IU/ML INJECTION SOLUTION (CARTRIDGE)

02506564                TRURAPI    AVN    **DNQW**

100IU/ML INJECTION SOLUTION (PEN)

02506572                TRURAPI SOLOSTAR    AVN    **DNQW**

02520974                KIRSTY    EMD    **DNQW**

**⑤INSULIN (ISOPHANE) HUMAN BIOSYNTHETIC**

100U/ML INJECTION SUSPENSION (10ML)

00587737                HUMULIN-N    LIL    **DNQW**

02024225                NOVOLIN GE NPH    NNO    **DNQW**



100U/ML INJECTION SUSPENSION (CARTRIDGE)			
01959239	HUMULIN-N CARTRIDGE	LIL	<b>DNQW</b>
02024268	NOVOLIN GE NPH PENFILL	NNO	<b>DNQW</b>

100U/ML INJECTION SUSPENSION (KWIKPEN)			
02403447	HUMULIN-N KWIKPEN	LIL	<b>DNQW</b>

**⑤INSULIN (REGULAR) HUMAN BIOSYNTHETIC**

100U/ML INJECTION SOLUTION (10ML)			
00586714	HUMULIN-R	LIL	<b>DNQW</b>
02024233	NOVOLIN GE TORONTO	NNO	<b>DNQW</b>

100U/ML INJECTION SOLUTION (CARTRIDGE)			
01959220	HUMULIN-R CARTRIDGE	LIL	<b>DNQW</b>
02024284	NOVOLIN GE TORONTO PENFILL	NNO	<b>DNQW</b>

[SEE APPENDIX A](#) FOR SA CRITERIA  
 500U/ML INJECTION SOLUTION (PEN)  
 02466864 ENTUZITY (SA)

LIL **DNQW**

**⑤INSULIN (REGULAR/ISOPHANE) HUMAN BIOSYNTHETIC**

100U/ML INJECTION SUSPENSION 30%/70% (10ML)			
00795879	HUMULIN 30/70	LIL	<b>DNQW</b>
02024217	NOVOLIN GE 30/70	NNO	<b>DNQW</b>

100U/ML INJECTION SUSPENSION 30%/70% (CARTRIDGE)			
01959212	HUMULIN 30/70 CARTRIDGE	LIL	<b>DNQW</b>
02025248	NOVOLIN GE 30/70 PENFILL	NNO	<b>DNQW</b>

100U/ML INJECTION SUSPENSION 40%/60% (CARTRIDGE)			
02024314	NOVOLIN GE 40/60 PENFILL	NNO	<b>DNQW</b>

100U/ML INJECTION SUSPENSION 50%/50% (CARTRIDGE)			
02024322	NOVOLIN GE 50/50 PENFILL	NNO	<b>DNQW</b>

**⑤INSULIN (REGULAR) LISPRO**

100U/ML INJECTION SOLUTION (10ML)			
02469901	ADMELOG	AVN	<b>DNQW</b>

100U/ML INJECTION SOLUTION (CARTRIDGE)			
02469898	ADMELOG	AVN	<b>DNQW</b>

100U/ML INJECTION SOLUTION (KWIKPEN)			
02469871	ADMELOG SOLOSTAR	AVN	<b>DNQW</b>

**⑤INSULIN (REGULAR/PROTAMINE) LISPRO**

100U/ML INJECTION SUSPENSION 25%/75% (CARTRIDGE)			
02240294	HUMALOG MIX 25 CARTRIDGE	LIL	DNQW

100U/ML INJECTION SUSPENSION 25%/75% (KWIKPEN)			
02403420	HUMALOG MIX 25 KWIKPEN	LIL	DNQW

**68:28.00 PITUITARY AGENTS**

**DESMOPRESSIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

10U/DOSE INTRANASAL SOLUTION (SPRAY PUMP)			
02242465	DEMOSPRESSIN (SA)	AAA	FGNQSW

0.1MG TABLET

00824305	DDAVP (SA)	FEI	FNQSW
02284030	APO-DESMOPRESSIN (SA)	APX	FGNQSW
02304368	PMS-DESMOPRESSIN (SA)	PMS	FGNQSW

0.2MG TABLET

00824143	DDAVP (SA)	FEI	FNQSW
02284049	APO-DESMOPRESSIN (SA)	APX	FGNQSW
02304376	PMS-DESMOPRESSIN (SA)	PMS	FGNQSW

60MCG ORAL DISINTEGRATING TABLET

02284995	DDAVP MELT (SA)	FEI	FNQSW
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120MCG ORAL DISINTEGRATING TABLET

02285002	DDVAP MELT (SA)	FEI	FNQSW
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240MCG ORAL DISINTEGRATING TABLET

02285010	DDVAP MELT (SA)	FEI	FNQSW
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**SOMATROGON**

[SEE APPENDIX A](#) FOR SA CRITERIA

24 MG/1.2 ML PREFILLED PEN

02521679	NGENLA (SA)	PFI	Y
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60 MG 1.2 ML PREFILLED PEN

02521687	NGENLA (SA)	PFI	Y
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**SOMATROPIN**

5.3 MG/ML SYRINGE

02401703	GENOTROPIN GOQUICK	PFI	Y
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12 MG/ML SYRINGE				
02401711	GENOTROPIN GOQUICK	PFI	Y	
0.6 MG/0.25 ML				
02401762	GENOTROPIN MINIQUICK	PFI	Y	
0.8 MG/0.25 ML				
02401770	GENOTROPIN MINIQUICK	PFI	Y	
1 MG/0.25 ML				
02401789	GENOTROPIN MINIQUICK	PFI	Y	
1.2 MG/0.25 ML				
02401797	GENOTROPIN MINIQUICK	PFI	Y	
1.4 MG/0.25 ML				
02401800	GENOTROPIN MINIQUICK	PFI	Y	
1.6 MG/0.25 ML				
02401819	GENOTROPIN MINIQUICK	PFI	Y	
1.8 MG/0.25 ML				
02401827	GENOTROPIN MINIQUICK	PFI	Y	
2 MG/0.25 ML				
02401835	GENOTROPIN MINIQUICK	PFI	Y	
<b>SOMATROPIN</b>				
6 MG INJECTION (CARTRIDGE)				
02243077	HUMATROPE CARTRIDGE	LIL	Y	
12 MG INJECTION (CARTRIDGE)				
02243078	HUMATROPE CARTRIDGE	LIL	Y	
24 MG INJECTION (CARTRIDGE)				
02243079	HUMATROPE CARTRIDGE	LIL	Y	
<b>SOMATROPIN</b>				
5 MG/1.5 ML CARTRIDGE				
02325063	OMNITROPE	SDZ	Y	
10 MG/1.5 ML CARTRIDGE				
02325071	OMNITROPE	SDZ	Y	
15 MG/1.5 ML CARTRIDGE				

02459647 OMNITROPE SDZ Y

**SOMATROPIN**

5 MG/1.5 ML PREFILLED PEN  
02334852 NORDITROPIN NORDIFLEX NNO Y

10 MG/1.5 ML PREFILLED PEN  
02334860 NORDITROPIN NORDIFLEX NNO Y

15 MG/1.5 ML PREFILLED PEN  
02334879 NORDITROPIN NORDIFLEX NNO Y

**SOMATROPIN**

5 MG/2 ML PEN INJECTOR  
02399091 NUTROPIN AQ HLR Y

10 MG/2 ML PEN INJECTOR  
02376393 NUTROPIN AQ HLR Y

20 MG/2 ML PEN INJECTOR  
02399083 NUTROPIN AQ HLR Y

**SOMATROPIN**

6 MG CARTRIDGE  
02350122 SAIZEN EMD Y

12 MG CARTRIDGE  
02350130 SAIZEN EMD Y

20 MG CARTRIDGE  
02350149 SAIZEN EMD Y

**68:30.04 SOMATOTROPIN AGONISTS**

[SEE APPENDIX A](#) FOR SA CRITERIA

**MECASERMIN**

10MG/ML VIAL  
02509733 INCRELEX (SA) IPS Y

## **68:32.00 PROGESTOGENS**

### **DIENOGEST**

[SEE APPENDIX A](#) FOR SA CRITERIA

#### **2MG TABLET**

02374900	VISANNE (SA)	BAY	<b>FQW</b>
02493055	ASPEN-DIENOGEST	ASN	<b>FGQW</b>
02498189	JAMP-DIENOGEST	JPC	<b>FGQW</b>
02543613	M-DIENOGEST	MRA	<b>FGQW</b>

### **MEDROXYPROGESTERONE ACETATE**

#### **2.5MG TABLET**

00708917	PROVERA	PFI	<b>FNQSW</b>
02221284	TEVA-MEDROXYPROGESTERONE	TEV	<b>FGNQSW</b>
02244726	AA-MEDROXY	AAA	<b>FGNQSW</b>

#### **5MG TABLET**

00030937	PROVERA	PFI	<b>FNQSW</b>
02221292	TEVA-MEDROXYPROGESTERONE	TEV	<b>FGNQSW</b>
02244727	AA-MEDROXY	AAA	<b>FGNQSW</b>

#### **10MG TABLET**

00729973	PROVERA	PFI	<b>FNQSW</b>
02221306	TEVA-MEDROXYPROGESTERONE	TEV	<b>FGNQSW</b>
02277298	AA-MEDROXY	AAA	<b>FGNQSW</b>

### **MEDROXYPROGESTERONE ACETATE**

150MG/ML INJECTION SUSPENSION (1ML)

02523493	DEPO-PROVERA	PFI	<b>FQW</b>
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### **PROGESTERONE**

#### **100MG CAPSULE**

02166704	PROMETRIUM	ORG	<b>FNQSW</b>
02439913	TEVA-PROGESTERONE	TEV	<b>FGNQSW</b>
02463113	REDDY-PROGESTERONE	RCH	<b>FGNQSW</b>
02476576	PMS-PROGESTERONE	PMS	<b>FGNQSW</b>
02493578	AURO-PROGESTERONE	ARO	<b>FGNQSW</b>
02516187	PROGRESTERONE	SNS	<b>FGNQSW</b>

## **68:36.04 THYROID AGENTS**

### **DESICCATED THYROID**

#### **30MG TABLET**

00023949	THYROID	ERF	<b>FNQSW</b>
60MG TABLET 00023957	THYROID	ERF	<b>FNQSW</b>
125MG TABLET 00023965	THYROID	ERF	<b>FNQSW</b>
<b>LEVOTHYROXINE SODIUM</b>			
25MCG TABLET 02172062	SYNTHROID	BGP	<b>FNQSW</b>
50MCG TABLET 02172070 02213192	SYNTHROID ELTROXIN	BGP ASN	<b>FNQSW FNQSW</b>
75MCG TABLET 02172089	SYNTHROID	BGP	<b>FNQSW</b>
88MCG TABLET 02172097	SYNTHROID	BGP	<b>FNQSW</b>
100MCG TABLET 02172100 02213206	SYNTHROID ELTROXIN	BGP ASN	<b>FNQSW FNQSW</b>
112MCG TABLET 02171228	SYNTHROID	BGP	<b>FNQSW</b>
125MCG TABLET 02172119	SYNTHROID	BGP	<b>FNQSW</b>
137MCG TABLET 02233852	SYNTHROID	BGP	<b>FNQSW</b>
150MCG TABLET 02172127 02213214	SYNTHROID ELTROXIN	BGP ASN	<b>FNQSW FNQSW</b>
175MCG TABLET 02172135	SYNTHROID	BGP	<b>FNQSW</b>
200MCG TABLET 02172143 02213222	SYNTHROID ELTROXIN	BGP ASN	<b>FNQSW FNQSW</b>

300MCG TABLET 02172151	SYNTHROID	BGP	FNQSW
<b>LIOTHYRONINE</b>			
5MCG TABLET 02494337	TEVA-LIOTHYRONINE	TEV	FGNQSW
25MCG TABLET 02494345	TEVA-LIOTHYRONINE	TEV	FGNQSW
<b>PROPYLTHIOURACIL</b>			
50MG TABLET 02521059	HALYCIL	ARN	FGNQSW
02523019	PROPYLTHIOURACIL	PCI	FGNQSW

**68:36.08 ANTI-THYROID AGENTS**

**METHIMAZOLE**

<b>5MG TABLET</b>			
00015741	TAPAZOLE	PAL	FNQSW
02480107	MAR-METHIMAZOLE	MAR	FGNQSW
02490625	JAMP-METHIMAZOLE	JPC	FGNQSW
<b>10MG TABLET</b>			
02296039	TAPAZOLE	PAL	FNQSW
02480115	MAR-METHIMAZOLE	MAR	FGNQSW
02490633	JAMP-METHIMAZOLE	JPC	FGNQSW

**72:00.00 LOCAL ANESTHETICS**

**LIDOCAINE HCL**

<b>1% INJECTION SOLUTION</b>			
00001732	XYLOCAINE	ASN	NQ
<b>2% INJECTION SOLUTION</b>			
00036641	XYLOCAINE	ASN	NQ
<b>2% ORAL SOLUTION</b>			
01968823	LIDODAN VISCOUS	ODN	FNQSW

**84:04.04 ANTI INFECTIVES (ANTIBIOTICS)**

**CLINDAMYCIN PHOSPHATE**

1% TOPICAL SOLUTION

02266938 TARO-CLINDAMYCIN  
02483769 CLINDAMYCIN

TAR FQW  
TLG FQW

**FRAMYCETIN SULFATE**

1% OINTMENT DRESSING (10CM X 10CM)

01988840 SOFRA TULLE

ERF FNQSW

**FUSIDIC ACID**

2% TOPICAL CREAM

00586668 FUCIDIN  
02528096 TARO-FUSIDIC ACID

LEO FNQSW  
TAR FGNQSW

**METRONIDAZOLE**

0.75% VAGINAL GEL

02125226 NIDAGEL

BLO FNQSW

1% TOPICAL CREAM

02156091 NORITATE

VAL FNQSW

**MUPIROCIN**

2% TOPICAL OINTMENT

02279983 TARO-MUPIROCIN

TAR FGNQSW

**POLYMYXIN B & BACITRACIN**

10,000U & 500U/G TOPICAL OINTMENT

02181908 POLYDERM  
02237227 POLYSPORIN  
02357569 JAMPOLYCIN

TAR N  
JJM N  
JPC N

**POLYMYXIN B & GRAMICIDIN**

10,000U & 250U/G TOPICAL CREAM

02230844 POLYSPORIN

JJM N

**SODIUM FUSIDATE**

2% TOPICAL OINTMENT

00586676 FUCIDIN

LEO FNQSW

**84:04.08 ANTI INFECTIVES (FUNGICIDES)**

**CICLOPIROX OLAMINE**

1% TOPICAL CREAM



02221802 LOPROX VAL **FNQSW**

1% TOPICAL LOTION

02221810 LOPROX VAL **FNSQW**

**CLOTRIMAZOLE**

1% TOPICAL CREAM

00812382 CLOTRIMADERM TAR **NSW**

02150867 CANESTEN BAY **NSW**

1% VAGINAL CREAM

00812366 CLOTRIMADERM TAR **NSW**

02150891 CANESTEN 6 BAY **NSW**

2% VAGINAL CREAM

00812374 CLOTRIMADERM TAR **NSW**

02150905 CANESTEN 3 BAY **NSW**

**KETOCONAZOLE**

2% TOPICAL CREAM

02245662 KETODERM TAR **FNQSW**

2% SHAMPOO

02182920 NIZORAL JJM **N**

**MICONAZOLE NITRATE**

2% TOPICAL CREAM

02085852 MICATIN WES **NSW**

02126567 MONISTAT DERM JJM **NSW**

2% VAGINAL CREAM

02084309 MONISTAT-7 JJM **NSW**

02231106 MICOZOLE TAR **NSW**

400MG VAGINAL OVULE

02126605 MONISTAT-3 JJM **NSW**

100MG VAGINAL SUPPOSITORY & 2% TOPICAL CREAM (COMBINATION PACKAGE)

02126257 MONISTAT-7 JJM **NSW**

400MG VAGINAL OVULE & 2% TOPICAL CREAM (COMBINATION PACKAGE)

02126249 MONISTAT-3 COMBINATION JJM **NSW**

**NYSTATIN**

100,000U/G TOPICAL CREAM

00716871 NYADERM TAR **GNSW**

25,000U/G VAGINAL CREAM  
00716901 NYADERM TAR GNSW

100,000U/G VAGINAL CREAM  
02194163 TEVA-NYSTATIN TEV GNSW

**TERBINAFINE HCL**

1% TOPICAL CREAM  
02031094 LAMISIL NVR FNQSW

1% TOPICAL SPRAY  
02238703 LAMISIL NVR FNQSW

**TOLNAFTATE**

1% TOPICAL CREAM  
00576034 TINACTIN BAY NW

1% TOPICAL POWDER  
00576042 TINACTIN BAY N

**84:04.12 ANTI-INFECTIVES, SCABICIDES, AND PEDICULICIDES**

**ISOPROPYL MYRISTATE**

50% TOPICAL LIQUID  
02279592 RESULTZ MFI NW

**PERMETHRIN**

1% CREME RINSE  
00771368 NIX CREME RINSE GSK NW  
02231480 KWELLADA-P CREME RINSE GSK NW

5% TOPICAL CREAM  
02219905 NIX DERMAL CREAM GSK NW

5% TOPICAL LOTION  
02231348 KWELLADA-P LOTION GSK NW

**84:04.16 ANTI INFECTIVES, OTHER ANTI INFECTIVES**

**METRONIDAZOLE**

1% TOPICAL GEL

02297809	METROGEL	GAC	FNQSW
10% VAGINAL CREAM			
01926861	FLAGYL	AVN	FNQSW
<b>SILVER SULFADIAZINE</b>			
1% TOPICAL CREAM			
00323098	FLAMAZINE	SNE	FNQSW

**84:06.00 ANTI INFLAMMATORY AGENTS**

APPROXIMATE RELATIVE POTENCIES OF TOPICAL STEROID PREPARATIONS

ULTRA HIGH POTENCY  
GROUP N

Betamethasone dipropionate 0.05% glycol cream, ointment, lotion  
Betamethasone dipropionate 0.05% & Salicylic Acid 3%, ointment  
Clobetasol propionate 0.05% cream, ointment, scalp lotion

HIGH POTENCY  
GROUP II

Amcinonide 0.1% ointment  
Betamethasone dipropionate 0.05% ointment  
Clobetasone butyrate 0.05% cream, ointment  
Desoximetasone 0.25% cream, ointment  
Desoximetasone 0.05% gel  
Fluocinonide 0.05% cream, ointment, gel

GROUP III

Betamethasone dipropionate 0.05% cream, lotion  
Betamethasone valerate 0.1% ointment  
Diflucortolone valerate 0.1% oily cream  
Triamcinolone acetonide 0.1% ointment

MID POTENCY

GROUP IV

Amcinonide 0.1% cream, lotion  
Beclomethasone dipropionate 0.025% cream, lotion (lotion d/c=d)  
Flucinolone acetonide 0.025% ointment  
Desoximetasone 0.05% cream  
Mometasone furoate 0.1% cream, ointment  
Triamcinolone acetonide 0.1% cream

GROUP V

Betamethasone valerate 0.1% cream, lotion, scalp lotion  
 Betamethasone valerate 0.05% cream, ointment, lotion  
 Triamcinolone acetonide 0.25% cream

**LOW POTENCY**

**GROUP VI**

Desonide 0.05% cream, ointment

**GROUP VII**

Hydrocortisone 0.5% cream, ointment, lotion

Hydrocortisone 1% cream, ointment, lotion

Hydrocortison 1% & Urea 10% cream, lotion

The classification of products in this table is based upon the >WHO Model Prescribing Information: Drugs Used in Dermatology (1995).

In general, ointments, as a result of their more occlusive property, tend to exhibit higher potency than creams containing the same concentration of the same anti-inflammatory agent. Cream formulations, in turn, appear to be more potent than lotions of the same strength.

**AMCINONIDE**

0.1% TOPICAL CREAM

02246714	TARO-AMCINONIDE	TAR	<b>FGNQSW</b>
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**BECLOMETHASONE DIPROPIONATE**

0.025% TOPICAL CREAM

02089602	PROPADERM	PAL	<b>FNQSW</b>
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**BETAMETHASONE DIPROPIONATE**

0.05% TOPICAL CREAM

00323071	DIPROSONE	MSD	<b>FNQSW</b>
01925350	TARO-SONE	TAR	<b>FGNQSW</b>
00804991	TEVA-TOPISONE	TEV	<b>FGNQSW</b>

0.05% TOPICAL LOTION

00417246	DIPROSONE	MSD	<b>FNQSW</b>
00809187	TEVA-TOPISONE	TEV	<b>FGNQSW</b>

0.05% TOPICAL OINTMENT

00344923	DIPROSONE	MSD	<b>FNQSW</b>
00805009	TEVA-TOPISONE	TEV	<b>FGNQSW</b>

0.05% TOPICAL GLYCOL BASE CREAM

00849650	TEVA-TOPILENE GLYCOL	TEV	<b>GFNQSW</b>
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0.05% TOPICAL GLYCOL BASE OINTMENT

00629367	DIPROLENE	ORG	<b>FNQSW</b>
00849669	TEVA-TOPILENE GLYCOL	TEV	<b>FGNQSW</b>

0.05% TOPICAL GLYCOL LOTION			
01927914	TEVA-TOPILENE	TEV	<b>FGNQSW</b>

**BETAMETHASONE DIPROPIONATE & CALCIPOTRIOL**

50MCG/0.5MG/GM TOPICAL GEL			
02319012	DOVOBET	LEO	<b>FNQSW</b>
02525178	TARO-CALCIPOTRIOL/BETAMETHASONE	TAR	<b>FGNQSW</b>

50MCG/0.5MG/GM TOPICAL OINTMENT			
02427419	TEVA-BETAMETHASONE/CALCIPOTRIOL	TEV	<b>FGNQSW</b>

50MCG/0.5MG/GM TOPICAL FOAM			
02457393	ENSTILAR	LEO	<b>FNQSW</b>

**BETAMETHASONE DIPROPIONATE & SALICYLIC ACID**

0.05% & 2% TOPICAL LOTION			
02245688	RATIO-TOPISALIC	TEV	<b>FGNQSW</b>

0.05% & 3% TOPICAL OINTMENT			
00578436	DIPROSALIC	MSD	<b>FNQSW</b>

**BETAMETHASONE VALERATE**

0.05% TOPICAL CREAM			
00535427	TEVA-ECTOSONE	TEV	<b>FGNQSW</b>
00716618	BETADERM	TAR	<b>FGNQSW</b>
02357860	CELESTODERM V/2	VAL	<b>FNQSW</b>

0.1% TOPICAL CREAM			
00535435	TEVA-ECTOSONE	TEV	<b>FGNQSW</b>
00716626	BETADERM	TAR	<b>FGNQSW</b>

0.05% TOPICAL OINTMENT			
00716642	BETADERM	TAR	<b>FGNQSW</b>
02357879	CELESTODERM V/2	VAL	<b>FNQSW</b>

0.1% TOPICAL OINTMENT			
00716650	BETADERM	TAR	<b>FGNQSW</b>
02357852	CELESTODERM V	VAL	<b>FNQSW</b>

0.05% TOPICAL LOTION			
00653209	TEVA-ECTOSONE	TEV	<b>FGNQSW</b>

0.1% TOPICAL LOTION

00750050 TEVA-ECTOSONE TEV FGNQSW

0.1% SCALP LOTION

00653217 TEVA-ECTOSONE TEV FGNQSW  
00716634 BETADERM TAR FGNQSW

**CLOBETASOL 17 PROPIONATE**

0.05% TOPICAL CREAM

01910272 TEVA-CLOBETASOL TEV FGNQSW  
02024187 MYLAN-CLOBETASOL MYL FGNQSW  
02213265 DERMOVATE TAR FGNQSW  
02245523 TARO-CLOBETASOL TAR FGNQSW

0.05% TOPICAL OINTMENT

01910280 TEVA-CLOBETASOL TEV FGNQSW  
02026767 MYLAN-CLOBETASOL MYL FGNQSW  
02213273 DERMOVATE TAR FGNQSW  
02245524 TARO-CLOBETASOL TAR FGNQSW

0.05% SCALP LOTION

01910299 TEVA-CLOBETASOL TEV FGNQSW  
02213281 DERMOVATE TAR FGNQSW  
02216213 MYLAN-CLOBETASOL MYL FGNQSW  
02245522 TARO-CLOBETASOL TAR FGNQSW

**CLOBETASONE BUTYRATE**

0.05% TOPICAL CREAM

02214415 SPECTRO ECZEMA CARE GSK FNQSW

**DESONIDE**

0.05% TOPICAL CREAM

02229315 PDP-DESONIDE PEN FGNQSW

0.05% TOPICAL OINTMENT

02229323 PDP-DESONIDE PEN FGNQSW

**DESOXIMETASONE**

0.05% TOPICAL EMOLLIENT CREAM

02221918 TOPICORT MILD VAL FNQSW

0.25% TOPICAL EMOLLIENT CREAM

02221896 TOPICORT VAL FNQSW

0.05% TOPICAL GEL

02221926 TOPICORT VAL FNQSW

0.25% TOPICAL OINTMENT			
02221934	TOPICORT	VAL	FNQSW
<b>FLUOCINONIDE</b>			
0.05% TOPICAL CREAM			
00716863	LYDERM	TAR	FGNQSW
02161923	LIDEX	VAL	FNQSW
0.05% TOPICAL GEL			
02161974	LIDEX	VAL	FNQSW
02236997	LYDERM	TAR	FGNQSW
0.05% TOPICAL OINTMENT			
02161966	LIDEX	VAL	FNQSW
02236996	LYDERM	TAR	FGNQSW
<b>HALOBETASOL PROPIONATE</b>			
0.01% LOTION			
02506262	BRYHALI	BAU	FNQSW
<b>HYDROCORTISONE</b>			
0.5% TOPICAL CREAM			
00716820	HYDERM	TAR	NW
1% TOPICAL CREAM			
00716839	HYDERM	TAR	FGNQSW
02412926	HYDROCORTISONE	SDZ	FGNQSW
80057178	JAMP-HC	JPC	FGNQSW
80057189	JAMP-HYDROCORTISONE	JPC	FGNQSW
1% OINTMENT			
00716693	CORTODERM	TAR	FGNQSW
<b>HYDROCORTISONE ACETATE/UREA</b>			
1%-10% TOPICAL CREAM			
00681989	DERMAFLEX HC	PAL	FNQSW
80061501	JAMP-HYDROCORTISONE ACET-UREA	JPC	FGNQSW
80073645	M-HC	MRA	FGNQSW
1%-10% TOPICAL LOTION			
00681997	DERMAFLEX HC	PAL	FNQSW
<b>HYDROCORTISONE &amp; PRAMOXINE &amp; ZINC</b>			
0.5% & 1% & 0.5% RECTAL OINTMENT			
02234466	PROCTODAN HC	ODN	FGNQSW

**HYDROCORTISONE & ZINC SULFATE**

0.5% & 0.5% RECTAL OINTMENT  
02128446 ANODAN-HC  
02387239 JAMPZINC-HC

ODN **FGNQSW**  
JPC **FGNQSW**

0.5% & 0.5% RECTAL SUPPOSITORY  
02236399 ANODAN-HC

ODN **FGNQSW**

**MOMETASONE FUROATE**

0.1% TOPICAL CREAM  
00851744 ELOCOM  
02367157 TARO-MOMETASONE

MSD **FNQSW**  
TAR **FGNQSW**

0.1% TOPICAL OINTMENT  
00851736 ELOCOM  
02248130 TEVA-MOMETASONE

MSD **FNQSW**  
TEV **FGNQSW**

0.1% LOTION  
00871095 ELOCOM

ORG **FNQSW**

**TRIAMCINOLONE ACETONIDE**

0.1% TOPICAL CREAM  
00716960 TRIADERM  
02194058 ARISTOCORT R

TAR **FGNQSW**  
VAL **FNQSW**

0.1% TOPICAL OINTMENT  
02194031 ARISTOCORT R

VAL **FNQSW**

0.1% ORAL TOPICAL OINTMENT  
01964054 ORACORT

TAR **FGNQSW**

**CLIOQUINOL & HYDROCORTISONE**

3% & 1% TOPICAL CREAM  
00074500 VIOFORM HYDROCORTISONE

PAL **FNQSW**

**HYDROCORTISONE & FRAMYCETIN & CINCHOCAINE HCL**

1% & 0.5% RECTAL OINTMENT  
02223252 PROCTOSEDYL  
02247322 PROCTOL

ALL **FNQSW**  
ODN **FGNQSW**

**TRIAMCINOLONE & NYSTATIN & NEOMYCIN & GRAMICIDIN**

2.5MG & 0.25MG & 100,000U & 1MG/G TOPICAL CREAM  
00717002 VIADERM K C

TAR **FGNQSW**

2.5MG & 0.25MG & 100,000U & 1MG/G TOPICAL OINTMENT



00717029

VIADERM K C

TAR FGNQSW

**84:08.00 ANTIPRURITICS AND TOPICAL ANESTHETICS**

**CALAMINE**

TOPICAL LOTION

00999829 CALAMINE LOTION

N

Note: The Drug Identification Number listed is for billing purposes only.

**LIDOCAINE HCL**

2% TOPICAL GEL

00001694 XYLOCAINE

ASN FNQSW

2% TOPICAL JELLY (SYRINGE)

00385484 XYLOCAINE

ASN NQ

02143879 LIDODAN

ODN NQ

5% OINTMENT

02083795 LIDODAN

ODN NW

**84:16.00 CELL STIMULANTS AND PROLIFERANTS**

**TRETINOIN**

0.05% TOPICAL CREAM

00443794 RETIN A

VAL FQW

0.025% TOPICAL GEL

00443816 RETIN A

VAL FQW

**84:24.00 EMOLLIENTS, DECMULCENTS, AND PROTECTANTS**

**DIMETHYLPOLYSILOXANE**

20% TOPICAL CREAM

02060841 BARRIERE

WES NW

**84:28.00 KERATOLYTIC AGENTS**

**UREA**

10% TOPICAL CREAM

80005397 URISEC 10

ODN NW

12% TOPICAL LOTION

00514896 URISEC

ODN NW

22% TOPICAL CREAM

00396125 URISEC 22

ODN NW

**84:32.00 KERATOPLASTIC AGENTS**

**COAL TAR**

1% TOPICAL SHAMPOO

02307146 T/GEL THERAPEUTIC

JJM NW

**84:92.00 MISCELLANEOUS SKIN AND MUCOUS MEMBRANE AGENTS**

**ACITRETIN**

10MG CAPSULE

02070847 SORIATANE  
02466074 TARO-ACITRETIN  
02468840 MINT-ACITRETIN

HLR FNQSW  
TAR FGNQSW  
MNT FGNQSW

25MG CAPSULE

02070863 SORIATANE  
02466082 TARO-ACITRETIN  
02468859 MINT-ACITRETIN

HLR FNQSW  
TAR FGNQSW  
MNT FGNQSW

**AZELAIC ACID**

15% TOPICAL GEL

02270811 FINACEA

LEO FNQSW

**BIMEKIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

160MG/ML PREFILLED SYRINGE

02525267 BIMZELX (SA)

UCB NMQW

160MG/ML AUTOINJECTOR

02525275 BIMZELX (SA)

UCB NMQW

**BRODALUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA  
210MG/1.5ML SYRINGE  
02473623 SILIQ (SA)

VAL **NMQW**

**CALCIPOTRIOL**

50MCG/G TOPICAL OINTMENT  
01976133 DOVONEX

LEO **FNQSW**

**DUPILUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA  
200MG/1.14ML PREFILLED PEN (SA)  
02524252 DUPIXENT (SA)

AVN **NMQW**

200MG/1.14ML SYRINGE  
02492504 DUPIXENT (SA)

AVN **NMQW**

300MG/2ML PREFILLED PEN  
02510049 DUPIXENT (SA)

AVN **NMQW**

300MG/2ML SYRINGE  
02470365 DUPIXENT (SA)

AVN **NMQW**

**FLUOROURACIL**

5% TOPICAL CREAM  
00330582 EFUDEX

VAL **FNQSW**

**GUSELKUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA  
100MG/ML AUTOINJECTOR  
02487314 TREMFYA (SA)

JAN **NMQW**

100MG/ML PREFILLED SYRINGE  
02469758 TREMFYA (SA)

JAN **NMQW**

**IMIQUIMOD**

5% TOPICAL CREAM  
02239505 ALDARA  
02482983 TARO-IMIQUIMOD

VAL **FNQSW**  
TAR **FGNQSW**

**ISOTRETINOIN**

COVERAGE IS LIMITED TO 30 DAYS AT A TIME

10MG CAPSULE  
00582344 ACCUTANE  
02257955 CLARUS

HLR **FQW**  
MYL **FGQW**

10MG CAPSULE 02396971	EPURIS	CIP	<b>FQW</b>
20MG CAPSULE 02396998	EPURIS	CIP	<b>FQW</b>
30MG CAPSULE 02397005	EPURIS	CIP	<b>FQW</b>
40MG CAPSULE 00582352 02257963	ACCUTANE CLARUS	HLR MYL	<b>FQW FGQW</b>
40MG CAPSULE 02397013	EPURIS	CIP	<b>FQW</b>

**IXEKIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

80MG/ML AUTOINJECTOR 02455102	TALTZ (SA)	LIL	<b>NMQW</b>
80MG/ML SYRINGE 02455110	TALTZ (SA)	LIL	<b>NMQW</b>

**RISANKIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

75MG/0.83ML PREFILLED SYRINGE 02487454	SKYRIZI (SA)	ABV	<b>NMQW</b>
150MG/ML PREFILLED PEN 02519291	SKYRIZI (SA)	ABV	<b>NMQW</b>
150MG/ML PREFILLED SYRINGE 02519283	SKYRIZI (SA)	ABV	<b>NMQW</b>
360MG/2.4ML PREFILLED CARTRIDGE 02532093	SKYRIZI (SA)	ABV	<b>NMQW</b>
600MG/10ML VIAL 02532107	SKYRIZI (SA)	ABV	<b>NMQW</b>

**SECUKINUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG/ML INJECTION

02438070	COSENTYX (SA)	NVR	<b>NMQW</b>
<b>TACROLIMUS</b>			
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
0.1% OINTMENT			
02244148	PROTOPIC (SA)	LEO	<b>FNQSW</b>
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
0.03% TOPICAL OINTMENT			
02244149	PROTOPIC (SA)	LEO	<b>FNQSW</b>
<b>TAZAROTENE</b>			
0.045% LOTION			
02517868	ARAZLO	BAU	<b>FNQSW</b>
<b>TAZAROTENE/HALOBETASOL PROPIONATE</b>			
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
0.01%/0.045% LOTION			
02499967	DUOBRII (SA)	BLO	<b>FNQSW</b>
<b>TILDRAKIZUMAB</b>			
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
100MG/ML PREFILLED SYRINGE			
02516098	ILUMYA (SA)	SUN	<b>NMQW</b>
<b>USTEKINUMAB</b>			
<a href="#">SEE APPENDIX A</a> FOR SA CRITERIA			
45MG/0.5ML SYRINGE			
02320673	STELARA (SA)	JAN	<b>NMQW</b>
02543036	JAMTEKI (SA)	JPC	<b>NMQW</b>
02544180	WEZLANA (SA)	AMG	<b>NMQW</b>
45MG/0.5ML VIAL			
02544202	WEZLANA (SA)	AMG	<b>NMQW</b>
90MG/ML SYRINGE			
02320681	STELARA (SA)	JAN	<b>NMQW</b>
02543044	JAMTEKI (SA)	JPC	<b>NMQW</b>
02544199	WEZLANA (SA)	AMG	<b>NMQW</b>
130MG/26ML VIAL			
02544210	WEZLANA I.V. (SA)	AMG	<b>NMQW</b>

## **86:12.00 GENITOURINARY SMOOTH MUSCLE RELAXANTS**

### **DARIFENACIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

7.5MG EXTENDED RELEASE TABLET

02273217	ENABLEX (SA)	MRS	<b>FNQSW</b>
02452510	APO-DARIFENACIN (SA)	APX	<b>FGNQSW</b>
02491869	JAMP-DARIFENACIN (SA)	JPC	<b>FGNQSW</b>

15MG EXTENDED RELEASE TABLET

02273225	ENABLEX (SA)	MRS	<b>FNQSW</b>
02452529	APO-DARIFENACIN (SA)	APX	<b>FGNQSW</b>
02491877	JAMP-DARIFENACIN (SA)	JPC	<b>FGNQSW</b>

### **FESOTERODINE FUMARATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

4MG EXTENDED RELEASE TABLET

02380021	TOVIAZ (SA)	PFI	<b>FNQSW</b>
02521768	SANDOZ-FESOTERODINE (SA)	SDZ	<b>FGNQSW</b>

8MG EXTENDED RELEASE TABLET

02380048	TOVIAZ (SA)	PFI	<b>FNQSW</b>
02521776	SANDOZ-FESOTERODINE (SA)	SDZ	<b>FGNQSW</b>

### **MIRABEGRON**

[SEE APPENDIX A](#) FOR SA CRITERIA

25MG EXTENDED RELEASE TABLET

02402874	MYRBETRIQ (SA)	AST	<b>FNQSW</b>
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50MG EXTENDED RELEASE TABLET

02402882	MYRBETRIQ (SA)	AST	<b>FNQSW</b>
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### **OXYBUTYNIN CHLORIDE**

1MG/ML SYRUP

02223376	PMS-OXYBUTYNIN	PMS	<b>FGNQSW</b>
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5MG TABLET

02163543	APO-OXYBUTYNIN	APX	<b>FGNQSW</b>
02230394	TEVA-OXYBUTYNIN	TEV	<b>FGNQSW</b>
02240550	PMS-OXYBUTYNIN	PMS	<b>FGNQSW</b>

02350238 OXYBUTYNIN SNS FGNQSW

**PROPIVERINE**

5MG TABLET

[SEE APPENDIX A](#) FOR SA CRITERIA  
02460289 MICTORYL PEDIATRIC

DUI FQW

**SOLIFENACIN**

5MG TABLET

02277263	VESICARE	AST	FNQSW
02397900	TEVA-SOLIFENACIN	TEV	FGNQSW
02399032	SANDOZ-SOLIFENACIN	SDZ	FGNQSW
02417723	PMS-SOLIFENACIN	PMS	FGNQSW
02423375	APO-SOLIFENACIN	APX	FGNQSW
02424339	JAMP-SOLIFENACIN	JPC	FGNQSW
02428911	JAMP-SOLIFENACIN	JPC	FGNQSW
02437988	TARO-SOLIFENACIN	RAN	FGNQSW
02439344	ACH-SOLIFENACIN	ACH	FGNQSW
02446375	AURO-SOLIFENACIN	ARO	FGNQSW
02458241	SOLIFENACIN	SNS	FGNQSW
02493039	PRZ-SOLIFENACIN	PRZ	FGNQSW
02529696	M-SOLIFENACIN SUCCINATE	MRA	FGNQSW

10MG TABLET

02277271	VESICARE	AST	FNQSW
02397919	TEVA-SOLIFENACIN	TEV	FGNQSW
02399040	SANDOZ-SOLIFENACIN	SDZ	FGNQSW
02417731	PMS-SOLIFENACIN	PMS	FGNQSW
02423383	APO-SOLIFENACIN	APX	FGNQSW
02424347	JAMP-SOLIFENACIN	JPC	FGNQSW
02428938	JAMP-SOLIFENACIN	JPC	FGNQSW
02437996	TARO-SOLIFENACIN	RAN	FGNQSW
02439352	ACH-SOLIFENACIN	ARO	FGNQSW
02446383	AURO-SOLIFENACIN	ARO	FGNQSW
02458268	SOLIFENACIN	SNS	FGNQSW
02493047	PRZ-SOLIFENACIN	PRZ	FGNQSW
02529718	M-SOLIFENACIN SUCCINATE	MRA	FGNQSW

**TOLTERODINE**

1MG TABLET

02239064	DETROL	UJC	FNQSW
02299593	TEVA-TOLTERODINE	TEV	FGNQSW
02423308	MINT-TOLTERODINE	MNT	FGNQSW
02496836	JAMP-TOLTERODINE	JPC	FGNQSW

2MG TABLET

02239065	DETROL	UJC	<b>FNQSW</b>
02299607	TEVA-TOLTERODINE	TEV	<b>FGNQSW</b>
02423316	MINT-TOLTERODINE	MNT	<b>FGNQSW</b>
02496844	JAMP-TOLTERODINE	JPC	<b>FGNQSW</b>

<b>2MG EXTENDED RELEASE CAPSULE</b>			
02244612	DETROL LA	UJC	<b>FNQSW</b>
02412195	TEVA-TOLTERODINE	TEV	<b>FGNQSW</b>
02413140	SANDOZ-TOLTERODINE	SDZ	<b>FGNQSW</b>

<b>4MG EXTENDED RELEASE CAPSULE</b>			
02244613	DETROL LA	UJC	<b>FNQSW</b>
02412209	TEVA-TOLTERODINE	TEV	<b>FGNQSW</b>
02413159	SANDOZ-TOLTERODINE	SDZ	<b>FGNQSW</b>

**TROSPIUM**

[SEE APPENDIX A](#) FOR SA CRITERIA

**20MG TABLET**

02275066	TROSEC (SA)	SNV	<b>FNQSW</b>
02488353	MAR-TROSPIUM (SA)	MAR	<b>FGNQSW</b>
02506661	JAMP-TROSPIUM (SA)	JPC	<b>FGNQSW</b>

**86:16.00 RESPIRATORY SMOOTH MUSCLE RELAXANTS**

**THEOPHYLLINE ANHYDROUS**

<b>100MG SUSTAINED RELEASE TABLET</b>			
00692689	AA-THEO-LA	AAA	<b>FGNQSW</b>

<b>200MG SUSTAINED RELEASE TABLET</b>			
00692697	AA-THEO-LA	AAA	<b>FGNQSW</b>

<b>300MG SUSTAINED RELEASE TABLET</b>			
00692700	AA-THEO-LA	AAA	<b>FGNQSW</b>

<b>400MG SUSTAINED RELEASE TABLET</b>			
02360101	THEO ER	AAA	<b>FGNQSW</b>

<b>600MG SUSTAINED RELEASE TABLET</b>			
02360128	THEO ER	AAA	<b>FGNQSW</b>

<b>5.33MG/ML ORAL SOLUTION</b>			
01966219	THEOLAIR	VAL	<b>FNQSW</b>



**88:08.00 VITAMIN B**

**CYANOCOBALAMIN**

1MG/ML INJECTION SOLUTION (10ML)

00521515	VITAMIN B12	SDZ	<b>NW</b>
01987003	CYANOCOBALAMIN	STE	<b>NW</b>
02413795	CYANOCOBALAMIN	MYL	<b>NW</b>

**FOLIC ACID**

1MG TABLET

00999899	FOLIC ACID		<b>OW</b>
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Note: The Drug Identification Number listed is for billing purposes only.

5MG TABLET

00426849	FOLIC ACID	AAA	<b>FGNQW</b>
02366061	JAMP-FOLIC ACID	JPC	<b>FGNQW</b>

**NIACIN**

100MG TABLET

00999879	NIACIN		<b>NW</b>
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Note: The Drug Identification Number listed is for billing purposes only.

500MG TABLET

00999889	NIACIN		<b>NW</b>
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Note: The Drug Identification Number listed is for billing purposes only.

**PYRIDOXINE**

25 MG Tablet

00268607	VITAMIN B6	VAL	<b>OX</b>
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**88:16.00 VITAMIN D**

**ALFACALCIDOL**

0.25MCG CAPSULE

02533316	SANDOZ-ALFACALCIDOL	SDZ	<b>FGNQSW</b>
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1MCG CAPSULE

00474525	ONE-ALPHA	XPI	<b>FNQSW</b>
02533324	SANDOZ-ALFACALCIDOL	SDZ	<b>FGNQSW</b>

**CALCITRIOL**

## 0.25MCG CAPSULE

00481823	ROCALTROL	HLR	<b>FNQSW</b>
02431637	CALCITRIOL-ODAN	ODN	<b>FGNQSW</b>
02485710	TARO-CALCITRIOL	TAR	<b>FGNQSW</b>
02495899	CALCITRIOL	STR	<b>FGNQSW</b>

## 0.5MCG CAPSULE

00481815	ROCALTROL	HLR	<b>FNQSW</b>
02431645	CALCITRIOL-ODAN	ODN	<b>FGNQSW</b>
02485729	TARO-CALCITRIOL	TAR	<b>FGNQSW</b>
02495902	CALCITRIOL	STR	<b>FGNQSW</b>

**VITAMIN D**

## 1000IU TABLET

00999869	VITAMIN D		<b>N</b>
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Note: The Drug Identification Number listed is for billing purposes only.

**88:20.00 VITAMIN E****VITAMIN E (D-ALPHA TOCOPHERYL ACETATE)**

## 200 UNIT CAPSULE

00999849	VITAMIN E		<b>CN</b>
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Note: The Drug Identification Number listed is for billing purposes only.

## 400 UNIT CAPSULE

00999859	VITAMIN E		<b>CN</b>
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Note: The Drug Identification Number listed is for billing purposes only.

**88:24.00 VITAMIN K ACTIVITY****PHYTONADIONE (VITAMIN K1)**

## 10MG/ML INJECTION SOLUTION (1ML)

00804312	VITAMIN K1	SDZ	<b>NQ</b>
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**92:00.00 MISCELLANEOUS THERAPEUTIC AGENTS****ABROCITINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG TABLET 02528363	CIBINQO (SA)	PFI	NMQW
100MG TABLET 02528371	CIBINQO (SA)	PFI	NMQW
200MG TABLET 02528398	CIBINQO (SA)	PFI	NMQW

**ALEMTUZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

12MG/1.2ML VIAL

02418320	LEMTRADA (SA)	GZY	NMQW
00904161	LEMTRADA (SA)*		NMQW
00904162	LEMTRADA (SA)*		NMQW
00904163	LEMTRADA (SA)*		NMQW
00904164	LEMTRADA (SA)*		NMQW
00904165	LEMTRADA (SA)*		NMQW
00904166	LEMTRADA (SA)*		NMQW

\*use when drug cost in excess of CPHA maximum

**AMIFAMPRIDINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET

02503034	RUZURGI (SA)	MDU	NMQW
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**AMIFAMPRIDINE PHOSPHATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET

02502984	FIRDAPSE (SA)	KYE	NMQW
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**ANIFROLUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG/ML VIAL

02522845	SAPHNELO (SA)	AZE	NMQW
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**BUROSUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG/ML VIAL

02483629	CRYSVITA (SA)	ULT	NMQW
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20MG/ML VIAL

02483637	CRYSVITA (SA)	ULT	NMQW
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30MG/ML VIAL

02483645	CRYSVITA (SA)	ULT	NMQW
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00904749            CRYSVITA (SA)\*  
\*use when drug cost in excess of CPHA maximum

**NMQW**

**CINACALCET**

[SEE APPENDIX A](#) FOR CRITERIA

**30MG TABLET**

02441624	TEVA-CINACALCET (SA)	TEV	<b>FGNQSW</b>
02452693	APO-CINACALCET (SA)	APX	<b>FGNQSW</b>
02478900	AURO-CINACALCET (SA)	ARO	<b>FGNQSW</b>
02480298	MAR-CINACALCET (SA)	MAR	<b>FGNQSW</b>
02481987	M-CINACALCET (SA)	MRA	<b>FGNQSW</b>
02500094	JAMP-CINACALCET (SA)	JPC	<b>FGNQSW</b>
02517604	PMS-CINACALCET (SA)	PMS	<b>FGNQSW</b>
02524880	CINACALCET (SA)	SNS	<b>FGNQSW</b>

**60MG TABLET**

02441632	TEVA-CINACALCET (SA)	TEV	<b>FGNQSW</b>
02452707	APO-CINACALCET (SA)	APX	<b>FGNQSW</b>
02478919	AURO-CINACALCET (SA)	ARO	<b>FGNQSW</b>
02480301	MAR-CINACALCET (SA)	MAR	<b>FGNQSW</b>
02481995	M-CINACALCET (SA)	MRA	<b>FGNQSW</b>
02500108	JAMP-CINACALCET (SA)	JPC	<b>FGNQSW</b>
02517612	PMS-CINACALCET (SA)	PMS	<b>FGNQSW</b>

**90MG TABLET**

02441640	TEVA-CINACALCET (SA)	TEV	<b>FGNQSW</b>
02452715	APO-CINACALCET (SA)	APX	<b>FGNQSW</b>
02478943	AURO-CINACALCET (SA)	ARO	<b>FGNQSW</b>
02480328	MAR-CINACALCET (SA)	MAR	<b>FGNQSW</b>
02482002	M-CINACALCET (SA)	MRA	<b>FGNQSW</b>
02500116	JAMP-CINACALCET (SA)	JPC	<b>FGNQSW</b>
02517620	PMS-CINACALCET (SA)	PMS	<b>FGNQSW</b>

**DIMETHYL FUMARATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

**120MG DELAYED RELEASE CAPSULE**

02404508	TECFIDERA (SA)	BIG	<b>NMQW</b>
02494809	GLN-DIMETHYL FUMARATE	GLN	<b>NMQW</b>
02495341	ACH-DIMETHYL FUMARATE	ACH	<b>NMQW</b>
02497026	PMS-DIMETHYL FUMARATE	PMS	<b>NMQW</b>
02502690	MAR-DIMETHYL FUMARATE	MAR	<b>NMQW</b>
02505762	APO-DIMETHYL FUMARATE	APX	<b>NMQW</b>
02513781	SANDOZ-DIMETHYL FUMARATE	SDZ	<b>NMQW</b>
02516047	JAMP-DIMETHYL FUMARATE	JPC	<b>NMQW</b>
02540746	AURO-DIMETHYL FUMARATE	ARO	<b>NMQW</b>

240MG DELAYED RELEASE CAPSULE			
02420201	TECFIDERA (SA)	BIG	NMQW
02494817	GLN-DIMETHYL FUMARATE	GLN	NMQW
02495368	ACH-DIMETHYL FUMARATE	ACH	NMQW
02497034	PMS-DIMETHYL FUMARATE	PMS	NMQW
02502704	MAR-DIMETHYL FUMARATE	MAR	NMQW
02505770	APO-DIMETHYL FUMARATE	APX	NMQW
02513803	SANDOZ-DIMETHYL FUMARATE	SDZ	NMQW
02516055	JAMP-DIMETHYL FUMARATE	JPC	NMQW
02540754	AURO-DIMETHYL FUMARATE	ARO	NMQW

**ETHINYL ESTRADIOL & CYPROTERONE**

0.035MG & 2MG TABLET			
02233542	DIANE-35	BAY	FQW
02290308	CYESTRA-35	PAL	FGQW
02309556	TEVA-CYPROTERONE/ETHINYL ESTRADIOL	TEV	FGQW

**FINGOLIMOD**

[SEE APPENDIX A](#) FOR CRITERIA

0.5MG CAPSULE			
02365480	GILENYA (SA)	NVR	NMQW
02469561	TEVA-FINGOLIMOD (SA)	TEV	NMQW
02469618	TARO-FINGOLIMOD (SA)	TAR	NMQW
02469715	MYLAN-FINGOLIMOD (SA)	MYL	NMQW
02469782	PMS-FINGOLIMOD (SA)	PMS	NMQW
02469936	APO-FINGOLIMOD (SA)	APX	NMQW
02474743	MAR-FINGOLIMOD (SA)	MAR	NMQW
02482606	SANDOZ-FINGOLIMOD (SA)	SDZ	NMQW
02487772	JAMP-FINGOLIMOD (SA)	JPC	NMQW

**GLATIRAMER ACETATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG PRE-FILLED SYRINGE			
02460661	GLATECT (SA)	PMS	NMQW

**⑤ GLUCAGON**

[SEE APPENDIX A](#) FOR SA CRITERIA

3MG NASAL SPRAY			
02492415	BAQSIMI (*)	LIL	DNQW

\*Regular benefit up to two devices per 12 months. Additional units require patient meets SA criteria

**⑤ GLUCAGON (RECOMBINANT DNA ORIGIN)**

[SEE APPENDIX A](#) FOR SA CRITERIA

INJECTION KIT

02243297	GLUCAGON KIT (*)	LIL	<b>DNQW</b>
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\*Regular benefit up to two devices per 12 months. Additional units require patient meets SA criteria

**⑤ GLUCAGON (HUMAN RECOMBINANT)**

[SEE APPENDIX A](#) FOR SA CRITERIA

INJECTION VIAL

02333619	GLUCAGEN VIAL (*)	PAL	<b>DNQW</b>
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INJECTION KIT

02333627	GLUCAGEN KIT (*)	PAL	<b>DNQW</b>
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\*Regular benefit up to two devices per 12 months. Additional units require patient meets SA criteria

**INTERFERON BETA-1A**

[SEE APPENDIX A](#) FOR SA CRITERIA

30MCG PREFILLED SYRINGE, 30MCG PEN WITH AUTO-INJECTOR

02269201	AVONEX PS (SA)	BIG	<b>NMQW</b>
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22MCG SYRINGE

02237319	REBIF (SA)	SRO	<b>NMQW</b>
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44MCG SYRINGE

02237320	REBIF (SA)	SRO	<b>NMQW</b>
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66MCG/1.5ML PRE-FILLED CARTRIDGE

02318253	REBIF MULTIDOSE (SA)	SRO	<b>NMQW</b>
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132MCG/1.5ML PRE-FILLED CARTRIDGE

02318261	REBIF MULTIDOSE (SA)	SRO	<b>NMQW</b>
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**INTERFERON BETA-1B**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.3MG INJECTION POWDER

02169649	BETASERON (SA)	BAY	<b>NMQW</b>
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**LANADELUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

300MG/2ML VIAL

02480948	TAKHZYRO (SA)	TAK	<b>NMQW</b>
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00904577	TAKHZYRO (SA)*		<b>NMQW</b>
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00904578	TAKHZYRO (SA)*		<b>NMQW</b>
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\*use when drug cost in excess of CPHA maximum

300MG/2ML PREFILLED SYRINGE			
02505614	TAKHZYRO (SA)	TAK	NMQW
00904638	TAKHZYRO (SA)*		NMQW
00904639	TAKHZYRO (SA)*		NMQW

\*use when drug cost in excess of CPHA maximum

**LANREOTIDE**

60MG/0.2ML PREFILLED SYRINGE			
02283395	SOMATULINE AUTOGEL	IPS	MNQW

90MG/0.3MG PREFILLED SYRINGE			
02283409	SOMATULINE AUTOGEL	IPS	MNQW

120MG/0.5ML PREFILLED SYRINGE			
02283417	SOMATULINE AUTOGEL	IPS	MNQW

**LEFLUNOMIDE**

10MG TABLET			
02241888	ARAVA	AVN	FNQSW
02256495	APO-LEFLUNOMIDE	APX	FGNQSW
02261251	TEVA-LEFLUNOMIDE	TEV	FGNQSW
02283964	SANDOZ-LEFLUOMIDE	SDZ	FGNQSW
02351668	LEFLUNOMIDE	SNS	FGNQSW
02543575	LEFLUNOMIDE	SIV	FGNQSW

20MG TABLET			
02241889	ARAVA	AVN	FNQSW
02256509	APO-LEFLUNOMIDE	APX	FGNQSW
02261278	TEVA-LEFLUNOMIDE	TEV	FGNQSW
02283972	SANDOZ -EFLUOMIDE	SDZ	FGNQSW
02351676	LEFLUNOMIDE	SNS	FGNQSW
02543583	LEFLUNOMIDE	SIV	FGNQSW

**LEUCOVORIN**

5MG TABLET			
02493357	RIV-LEUCOVORIN	RIV	FGNQSW
02496828	MINT- LEUCOVORIN	MNT	FGNQSW

**LEVOCARNITINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

330MG TABLET			
02144328	CARNITOR (SA)	LBI	FNQSW

100MG/ML ORAL SOLUTION			
02144336	CARNITOR (SA)	LBI	FNQSW
02492105	ODAN-LEVOCARNITINE (SA)	ODN	FGNQSW

**MONTELUKAST****4MG CHEWABLE TABLET**

02243602	SINGULAIR	MSD	<b>FQW</b>
02330385	SANDOZ-MONTELUKAST	SDZ	<b>FGQW</b>
02354977	PMS-MONTELUKAST	PMS	<b>FGQW</b>
02355507	TEVA-MONTELUKAST	TEV	<b>FGQW</b>
02377608	APO-MONTELUKAST	APX	<b>FGQW</b>
02382458	MONTELUKAST	SIV	<b>FGQW</b>
02399865	MAR-MONTELUKAST	MAR	<b>FGQW</b>
02408627	MINT-MONTELUKAST	MNT	<b>FGQW</b>
02422867	AURO-MONTELUKAST	ARO	<b>FGQW</b>
02442353	JAMP-MONTELUKAST	JPC	<b>FGQW</b>
02514877	JAMP-MONTELUKAST	JPC	<b>FGQW</b>
02522101	NAT-MONTELUKAST	NAT	<b>FGQW</b>

**5MG CHEWABLE TABLET**

02238216	SINGULAIR	MSD	<b>FQW</b>
02330393	SANDOZ-MONTELUKAST	SDZ	<b>FGQW</b>
02354985	PMS-MONTELUKAST	PMS	<b>FGQW</b>
02355515	TEVA-MONTELUKAST	TEV	<b>FGQW</b>
02377616	APO-MONTELUKAST	APX	<b>FGQW</b>
02379325	MONTELUKAST	SNS	<b>FGQW</b>
02382466	MONTELUKAST	SIV	<b>FGQW</b>
02399873	MAR-MONTELUKAST	MAR	<b>FGQW</b>
02408635	MINT-MONTELUKAST	MNT	<b>FGQW</b>
02422875	AURO-MONTELUKAST	ARO	<b>FGQW</b>
02442361	JAMP-MONTELUKAST	JPC	<b>FGQW</b>
02514885	JAMP-MONTELUKAST	JPC	<b>FGQW</b>
02522128	NAT-MONTELUKAST	NAT	<b>FGQW</b>

**10MG TABLET**

02238217	SINGULAIR	MSD	<b>FNQSW</b>
02328593	SANDOZ-MONTELUKAST	SDZ	<b>FGNQSW</b>
02355523	TEVA-MONTELUKAST	TEV	<b>FGNQSW</b>
02373947	PMS-MONTELUKAST	PMS	<b>FGNQSW</b>
02374609	APO-MONTELUKAST	APX	<b>FGNQSW</b>
02379236	MONTELUKAST SODIUM	ACH	<b>FGNQSW</b>
02379333	MONTELUKAST	SNS	<b>FGNQSW</b>
02382474	MONTELUKAST	SIV	<b>FGNQSW</b>
02389517	RAN-MONTELUKAST	RAN	<b>FGNQSW</b>
02391422	JAMP-MONTELUKAST	JPC	<b>FGNQSW</b>
02401274	AURO-MONTELUKAST	ARO	<b>FGNQSW</b>
02399997	MAR-MONTELUKAST	MAR	<b>FGNQSW</b>
02408643	MINT-MONTELUKAST	MNT	<b>FGNQSW</b>
02488183	M-MONTELUKAST	MRA	<b>FGNQSW</b>
02489821	NRA-MONTELUKAST	NRA	<b>FGNQSW</b>



02522136	NAT-MONTELUKAST	NAT	<b>FGNQSW</b>
4MG GRANULES IN PACKET			
02247997	SINGULAIR	MSD	<b>FQW</b>
02358611	SANDOZ-MONTELUKAST	SDZ	<b>FGQW</b>

**NATALIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA  
300MG & 15ML VIAL

02286386	TYSABRI (SA)	BIG	<b>NMQW</b>
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**NUSINERSEN**

[SEE APPENDIX A](#) FOR SA CRITERIA  
2.4MG/ML INTRATHECAL VIAL

02465663	SPINRAZA (SA)	BIG	<b>NMQW</b>
00904366	SPINRAZA (SA)*		<b>NMQW</b>
00904367	SPINRAZA (SA)*		<b>NMQW</b>
00904368	SPINRAZA (SA)*		<b>NMQW</b>
00904369	SPINRAZA (SA)*		<b>NMQW</b>
00904370	SPINRAZA (SA)*		<b>NMQW</b>
00904371	SPINRAZA (SA)*		<b>NMQW</b>
00904372	SPINRAZA (SA)*		<b>NMQW</b>
00904373	SPINRAZA (SA)*		<b>NMQW</b>
00904374	SPINRAZA (SA)*		<b>NMQW</b>
00904375	SPINRAZA (SA)*		<b>NMQW</b>
00904376	SPINRAZA (SA)*		<b>NMQW</b>

\*use when drug cost in excess of CPHA maximum

**OCTREOTIDE**

200MCG/ML INJECTION (5ML)

02049392	SANDOSTATIN	NVR	<b>FNQSW</b>
02248642	OCTREOTIDE OMEGA	OMG	<b>FGNQSW</b>

10MG PREFILLED SYRINGE

02239323	SANDOSTATIN LAR	NVR	<b>MNQW</b>
02503751	OCTREOTIDE	TEV	<b>MNQW</b>

20MG PREFILLED SYRINGE

02239324	SANDOSTATIN LAR	NVR	<b>MNQW</b>
02503778	OCTREOTIDE	TEV	<b>MNQW</b>

30MG PREFILLED SYRINGE

02239325	SANDOSTATIN LAR	NVR	<b>MNQW</b>
02503786	OCTREOTIDE	TEV	<b>MNQW</b>

**OCRELIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

300 MG/10 ML (VIAL)

02467224 OCREVUS (SA)

00904527 OCREVUS (SA)\*

\*use when drug cost in excess of CPHA maximum

HLR **NMQW**  
**NMQW**

**OFATUMUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG/0.4ML PREFILLED PEN

02511355 KESIMPTA (SA)

NVR **NMQW**

**ONABOTULINUMTOXINA**

[SEE APPENDIX A](#) FOR SA CRITERIA

200 UNITS/VIAL

02531585 BOTOX (SA)

ALL **FNQSW**

**OZANIMOD**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.23 MG (4)-0.46 MG (3) INITIATION PACK

02506009 ZEPOSIA (SA)

BMS **FNQSW**

0.92MG CAPSULE

02505991 ZEPOSIA (SA)

BMS **FNQSW**

**PAMIDRONATE DISODIUM**

[SEE APPENDIX A](#) FOR SA CRITERIA

30MG INJECTION

02244550 PAMIDRONATE DISODIUM (SA)

PFI **FNQSW**

60MG INJECTION

02244551 PAMIDRONATE DISODIUM (SA)

PFI **FNQSW**

90MG INJECTION

02244552 PAMIDRONATE DISODIUM (SA)

PFI **FNQSW**

**PEGINTERFERON BETA-1A**

[SEE APPENDIX A](#) FOR SA CRITERIA

63/94MCG/0.5ML

02444402 PLEGRIDY (SA)

BIG **NMQW**

125MCG/0.5ML

02444399 PLEGRIDY (SA)

BIG **NMQW**

**PENTOSAN POLYSULFATE SO4**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE

02029448 ELMIRON (SA) JAN FNQSW

**PHENYLALANINE-REDUCED FOODS**

NUTRITIONAL FORMULA

00030800 PHENEX-1 ROS P  
04444444 PHENEX-2 ROS P  
00368020 PHENYL-FREE MJS P

**RISDIPLAM**

[SEE APPENDIX A](#) FOR SA CRITERIA

Note: Risdiplam claims must be billed in mgs

0.75MG/ML POWDER FOR ORAL SOLUTION

02514931 EVRYSDI (SA) HLR NMQW  
00904768 EVRYSDI (SA)\* NMQW  
00904769 EVRYSDI (SA)\* NMQW  
00904770 EVRYSDI (SA)\* NMQW

\*use when drug cost in excess of CPHA maximum

**SATRALIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

120MG/ML PREFILLED SYRINGE

02499681 ENSPRYNG (SA) HLR NMQW

**SIPONIMOD**

[SEE APPENDIX A](#) FOR SA CRITERIA

0.25MG TABLET

02496429 MAYZENT (SA) NVR NMQW

2MG TABLET

02496437 MAYZENT (SA) NVR NMQW

**SODIUM CHLORIDE**

7% INHALATION LIQUID

80029414 HYPERSAL 7% KEG C

**SODIUM CROMOGLYCATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE

00500895 NALCROM (SA) AVN FQSW

**SORAFENIB TOSYLATE**

[SEE APPENDIX A](#) FOR SA CRITERIA

200MG TABLET

02284227 NEXAVAR (SA) BAY NMQW

**TERIFLUNOMIDE**[SEE APPENDIX A](#) FOR SA CRITERIA**14MG TABLET**

02416328	AUBAGIO (SA)	GZY	NMQW
02500310	NAT-TERIFLUNOMIDE (SA)	NAT	NMQW
02500434	PMS-TERIFLUNOMIDE (SA)	PMS	NMQW
02500469	MAR-TERIFLUNOMIDE (SA)	MAR	NMQW
02500639	APO-TERIFLUNOMIDE (SA)	APX	NMQW
02501090	TEVA-TERIFLUNOMIDE (SA)	TEV	NMQW
02502933	ACH-TERIFLUNOMIDE (SA)	ACH	NMQW
02504170	JAMP-TERIFLUNOMIDE (SA)	JPC	NMQW
02505843	SANDOZ-TERIFLUNOMIDE (SA)	SDZ	NMQW
02523833	M-TERIFLUNOMIDE (SA)	MRA	NMQW

**TETRABENAZINE****25MG TABLET**

02199270	NITOMAN	VAL	FNQSW
02402424	PMS-TETRABENAZINE	PMS	FGNQSW
02407590	APO-TETRABENAZINE	APX	FGNQSW
02410338	TETRABENAZINE	STE	FGNQSW

**TOCILIZUMAB**[SEE APPENDIX A](#) FOR SA CRITERIA**80MG/4ML IV VIAL**

02350092	ACTEMRA (SA)	HLR	NMQW
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**200MG/10ML IV VIAL**

02350106	ACTEMRA (SA)	HLR	NMQW
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**400MG/20ML IV VIAL**

02350114	ACTEMRA (SA)	HLR	NMQW
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**162MG/0.9ML SYRINGE**

02424770	ACTEMRA (SA)	HLR	NMQW
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**162MG/0.9ML PREFILLED AUTOINJECTOR**

02483327	ACTEMRA (SA)	HLR	NMQW
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**TRIENTINE**[SEE APPENDIX A](#) FOR SA CRITERIA**250MG CAPSULE**

02504855	MAR-TRIENTINE (SA)	MAR	NMQW
02515067	WAYMADE-TRIENTINE (SA)	WMD	NMQW

**TRIMEPRAZINE TARTRATE****2.5MG TABLET**

01926306 PANECTYL ERF FNQW

5MG TABLET  
01926292 PANECTYL ERF FNQW

**UPADACITINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA  
15MG EXTENDED RELEASE TABLET  
02495155 RINVOQ (SA)

ABV MNQW

30MG EXTENDED RELEASE TABLET  
02520893 RINVOQ (SA) ABV MNQW

**VEDOLIZUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA  
300MG VIAL

02436841 ENTYVIO (SA) TAK NMQW

108MG/0.68ML PREFILLED SYRINGE  
02497875 ENTYVIO (SA) TAK NMQW

108MG/0.68ML PREFILLED PEN  
02497867 ENTYVIO (SA) TAK NMQW

**ZOLEDRONIC ACID**

5MG/100ML INJECTION  
02269198 ACLASTA SDZ FNQSW  
02422433 ZOLEDRONIC ACID RCH FGNQSW

**92:00.08 ALFA REDUCTASE INHIBITORS**

**DUTASTERIDE**

0.5MG CAPSULE  
02247813 AVODART GSK FNQSW  
02393220 PMS-DUTASTERIDE PMS FGNQSW  
02404206 APO-DUTASTERIDE APX FGNQSW  
02408287 TEVA-DUTASTERIDE TEV FGNQSW  
02416298 MED-DUTASTERIDE GMP FGNQSW  
02424444 SANDOZ-DUTASTERIDE SDZ FGNQSW  
02428873 MINT-DUTASTERIDE MNT FGNQSW  
02429012 DUTASTERIDE SIV FGNQSW  
02443058 DUTASTERIDE SNS FGNQSW  
02469308 AURO-DUTASTERIDE ARO FGNQSW

02484870	JAMP-DUTASTERIDE	JPC	FGNQSW
<b>FINASTERIDE</b>			
5MG TABLET			
02010909	PROSCAR	MDS	FNQSW
02348500	TEVA-FINASTERIDE	TEV	FGNQSW
02322579	SANDOZ-FINASTERIDE	SDZ	FGNQSW
02310112	PMS-FINASTERIDE	PMS	FGNQSW
02355043	FINASTERIDE	ACH	FGNQSW
02357224	JAMP-FINASTERIDE	JPC	FGNQSW
02365383	APO-FINASTERIDE	APX	FGNQSW
02389878	MINT-FINASTERIDE	MNT	FGNQSW
02405814	AURO-FINASTERIDE	ARO	FGNQSW
02445077	FINASTERIDE	SNS	FGNQSW
02447541	FINASTERIDE	SIV	FGNQSW
02455013	RIVA-FINASTERIDE	RIV	FGNQSW
02522489	M-FINASTERIDE	MRA	FGNQSW

## **92:16.00 ANTIGOUT AGENTS**

### **ALLOPURINOL**

#### 100MG TABLET

00402818	APO-ALLOPURINOL	APX	FGNQSW
02396327	MAR-ALLOPURINOL	MAR	FGNQSW
02402769	APO-ALLOPURINOL	APX	FGNQSW

#### 200MG TABLET

00479799	APO-ALLOPURINOL	APX	FGNQSW
02396335	MAR-ALLOPURINOL	MAR	FGNQSW
02402777	APO-ALLOPURINOL	APX	FGNQSW

#### 300MG TABLET

00402796	APO-ALLOPURINOL	APX	FGNQSW
02396343	MAR-ALLOPURINOL	MAR	FGNQSW
02402785	APO-ALLOPURINOL	APX	FGNQSW

### **COLCHICINE**

#### 0.6MG TABLET

00287873	SANDOZ-COLCHICINE	SDZ	FGNQSW
00572349	COLCHICINE-ODAN	ODN	FGNQSW
02373823	JAMP-COLCHICINE	JPC	FGNQSW
02402181	PMS-COLCHICINE	PMS	FGNQSW

**FEBUXOSTAT**[SEE APPENDIX A](#) FOR SA CRITERIA**80MG TABLETS**

02466198	TEVA-FEBUXOSTAT	TEV	<b>FGNQSW</b>
02473607	MAR-FEBUXOSTAT	MAR	<b>FGNQSW</b>
02490870	JAMP-FEBUXOSTAT	JPC	<b>FGNQSW</b>
02533243	AURO-FEBUXOSTAT	ARO	<b>FGNQSW</b>
02539837	FEBUXOSTAT	SNS	<b>FGNQSW</b>

**92:24:00 BONE RESORPTION INHIBITORS****ALENDRONATE & CHOLECALCIFEROL****70MG/5600 UNIT TABLET**

02314940	FOSAVANCE	MSD	<b>FNQSW</b>
02454475	APO-ALENDRONATE/VITAMIN D3	APX	<b>FGNQSW</b>
02519836	JAMP-ALENDRONATE/VITAMIN D3	JPC	<b>FGNQSW</b>

**ALENDRONATE SODIUM****10MG TABLET**

02248728	APO-ALENDRONATE	APX	<b>FGNQSW</b>
02381486	ALENDRONATE SODIUM	ACH	<b>FGNQSW</b>
02384701	RAN-ALENDRONATE	RAN	<b>FGNQSW</b>
02388545	AURO-ALENDRONATE	ARO	<b>FGNQSW</b>

**70MG TABLET**

02245329	FOSAMAX	MSD	<b>FNQSW</b>
02248730	APO-ALENDRONATE	APX	<b>FGNQSW</b>
02261715	TEVA-ALENDRONATE	TEV	<b>FGNQSW</b>
02270889	RIVA-ALENDRONATE	RIV	<b>FGNQSW</b>
02284006	PMS-ALENDRONATE	PMS	<b>FGNQSW</b>
02288109	SANDOZ-ALENDRONATE	SDZ	<b>FGNQSW</b>
02299712	ALENDRONATE	SIV	<b>FGNQSW</b>
02352966	ALENDRONATE	SNS	<b>FGNQSW</b>
02381494	ALENDRONATE SODIUM	ACH	<b>FGNQSW</b>
02385031	JAMP-ALENDRONATE	JPC	<b>FGNQSW</b>
02388553	AURO-ALENDRONATE	ARO	<b>FGNQSW</b>
02394871	MINT-ALENDRONATE	MNT	<b>FGNQSW</b>
02485184	AG-ALENDRONATE	ANG	<b>FGNQSW</b>
02500175	JAMP-ALENDRONATE	JPC	<b>FGNQSW</b>
02529394	M-ALENDRONATE	MRA	<b>FGNQSW</b>

**DENOSUMAB**[SEE APPENDIX A](#) FOR SA CRITERIA

60MG/ML SC SYRINGE  
 02343541 PROLIA (SA) AMG **FNQSW**

**RALOXIFENE**

60MG TABLET  
 02279215 APO-RALOXIFENE APX **FGNQSW**  
 02358840 ACT-RALOXIFENE TEV **FGNQSW**  
 02540681 JAMP-RALOXIFENE JPC **FGNQSW**

**RISEDRONATE SODIUM**

5MG TABLET  
 02298376 TEVA-RISEDRONATE TEV **FGNQSW**

[SEE APPENDIX A](#) FOR SA CRITERIA

30MG TABLET  
 02298384 TEVA-RISEDRONATE (SA) TEV **FGNQSW**

35MG TABLET

02246896 ACTONEL ALL **FNQSW**  
 02298392 TEVA-RISEDRONATE TEV **FGNQSW**  
 02302209 PMS-RISEDRONATE PMS **FGNQSW**  
 02327295 SANDOZ-RISEDRONATE SDZ **FGNQSW**  
 02353687 APO-RISEDRONATE APX **FGNQSW**  
 02368552 JAMP-RISEDRONATE JPC **FGNQSW**  
 02370255 SANIS-RISEDRONATE SNS **FGNQSW**  
 02406306 AURO-RISEDRONATE ARO **FGNQSW**  
 02411407 RISEDRONATE SIV **FGNQSW**

**92:36.00 DISEASE-MODIFYING ANTIRHEUMATIC AGENTS**

**ABATACEPT**

[SEE APPENDIX A](#) FOR SA CRITERIA

250 MG VIAL  
 02282097 ORENCIA (SA) BMS **NMQW**

125MG/ML PREFILLED SC SYRINGE  
 02402475 ORENCIA (SA) BMS **NMQW**

**ADALIMUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

20MG/0.2 ML PREFILLED SYRINGE



02542315	HYRIMOZ (SA)	SDZ	<b>NMQW</b>
20MG/0.4 ML PREFILLED SYRINGE			
02459310	AMGEVITA (SA)	AMG	<b>NMQW</b>
02502380	HULIO (SA)	BGP	<b>NMQW</b>
02505258	HYRIMOZ (SA)	SDZ	<b>NMQW</b>
02511061	ABRILADA (SA)	PFI	<b>NMQW</b>
40MG/0.4 ML PREFILLED SYRINGE			
02523760	YUFLYMA (SA)	LIL	<b>NMQW</b>
02523949	SIMLANDI (SA)	JPC	<b>NMQW</b>
02533472	HADLIMA (SA)	MER	<b>NMQW</b>
02542323	HYRIMOZ (SA)	SDZ	<b>NMQW</b>
40MG/0.4 ML PEN INJECTOR			
02523779	YUFLYMA (SA)	LIL	<b>NMQW</b>
02523957	SIMLANDI (SA)	JPC	<b>NMQW</b>
02533480	HADLIMA PUSHTOUCH (SA)	MER	<b>NMQW</b>
02542331	HYRIMOZ (SA)	SDZ	<b>NMQW</b>
40MG/0.8 ML PREFILLED SYRINGE			
02459299	AMGEVITA (SA)	AMG	<b>NMQW</b>
02473097	HADLIMA (SA)	MER	<b>NMQW</b>
02492164	HYRIMOZ (SA)	SDZ	<b>NMQW</b>
02502399	HULIO (SA)	BGP	<b>NMQW</b>
02502682	IDACIO (SA)	FKB	<b>NMQW</b>
02511053	ABRILADA (SA)	PFI	<b>NMQW</b>
40MG/0.8 ML PREFILLED PEN			
02459302	AMGEVITA (SA)	AMG	<b>NMQW</b>
02473100	HADLIMA PUSHTOUCH (SA)	MER	<b>NMQW</b>
02492156	HYRIMOZ (SA)	SDZ	<b>NMQW</b>
02502402	HULIO (SA)	BGP	<b>NMQW</b>
02502674	IDACIO (SA)	FKB	<b>NMQW</b>
02511045	ABRILADA (SA)	PFI	<b>NMQW</b>

80MG/0.8 ML PREFILLED SYRINGE

02523965	SIMLANDI (SA)	JPC	NMQW
02535076	YUFLYMA (SA)	CLT	NMQW
02542358	HYRIMOZ (SA)	SDZ	NMQW
80MG/0.8 ML PEN INJECTOR			
02535084	YUFLYMA (SA)	CLT	NMQW
02542366	HYRIMOZ (SA)	SDZ	NMQW

### BOSENTAN

[SEE APPENDIX A](#) FOR CRITERIA

62.5 MG TABLET

02244981	TRACLEER (SA)	JAN	MSQ
02383012	PMS-BOSENTAN (SA)	PMS	GMSQ
02467984	NAT-BOSENTAN (SA)	NAT	GMSQ
02483130	TARO-BOSENTAN (SA)	TAR	GMSQ

125 MG TABLET

02244982	TRACLEER (SA)	JAN	MSQ
02383020	PMS-BOSENTAN (SA)	PMS	GMSQ
02467992	NAT-BOSENTAN (SA)	NAT	GMSQ
02483149	TARO-BOSENTAN (SA)	TAR	GMSQ

### CANAKINUMAB

[SEE APPENDIX A](#) FOR SA CRITERIA

02460351	ILARIS (SA)	NVR	NMQW
00904405	ILARIS (SA)*		NMQW

\*use when drug cost in excess of CPHA maximum

### CERTOLIZUMAB

[SEE APPENDIX A](#) FOR CRITERIA

200MG/ML SYRINGE KIT

02331675	CIMZIA (SA)	UCB	NMQW
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200MG/ML AUTO-INJECTOR KIT

02465574	CIMZIA (SA)	UCB	NMQW
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### ETANERCEPT

[SEE APPENDIX A](#) FOR SA CRITERIA

25MG/0.5ML PEN INJECTOR

02462877	ERELZI (SA)	SDZ	NMQW
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50MG/ML PEN INJECTOR

02455331	BRENZYS (SA)	MSD	NMQW
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02462850	ERELZI (SA)	SDZ	NMQW
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02530309	RYMTI (SA)	LUP	<b>NMQW</b>
50MG/ML PRE-FILLED SYRINGE			
02455323	BRENZYS (SA)	MSD	<b>NMQW</b>
02462869	ERELZI (SA)	SDZ	<b>NMQW</b>
02530309	RYMTI (SA)	LUP	<b>NMQW</b>

**GOLIMUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

50MG/0.5ML SYRINGE

02324776	SIMPONI (SA)	JAN	<b>NMQW</b>
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50MG/0.5ML AUTO-INJECTOR

02324784	SIMPONI (SA)	JAN	<b>NMQW</b>
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**ICATIBANT**

[SEE APPENDIX A](#) FOR SA CRITERIA

30MG/3ML SC SYRINGE

02425696	FIRAZYR (SA)	SHR	<b>NMQW</b>
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**INFLIXIMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG/VIAL INJECTION

02419475	INFLECTRA (SA)	HOS	<b>NMQW</b>
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02470373	RENFLEXIS (SA)	MSD	<b>NMQW</b>
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02496933	AVSOLA (SA)	AGA	<b>NMQW</b>
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**SARILUMAB**

[SEE APPENDIX A](#) FOR SA CRITERIA

150MG/1.14ML PEN

02472961	KEVZARA (SA)	AVN	<b>NMQW</b>
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200MG/1.14ML SYRINGE

02460548	KEVZARA (SA)	AVN	<b>NMQW</b>
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200MG/1.14ML PEN

02472988	KEVZARA (SA)	AVN	<b>NMQW</b>
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**TOFACITINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

5MG TABLET

02423898	XELJANZ (SA)	PFI	<b>NMQW</b>
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02511304	TARO-TOFACITINIB (SA)	TAR	<b>NMQW</b>
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02522799	PMS-TOFACITINIB (SA)	PMS	<b>NMQW</b>
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02522896	JAMP-TOFACITINIB (SA)	JPC	NMQW
02530007	AURO-TOFACITINIB (SA)	ARO	NMQW
10MG TABLET			
02480786	XELJANZ (SA)	PFI	NMQW
02511312	TARO-TOFACITINIB (SA)	TAR	NMQW
02530015	AURO-TOFACITINIB (SA)	ARO	NMQW
11MG TABLET			
02470608	XELJANZ XR (SA)	PFI	NMQW

## **92:44.00 IMMUNOSUPPRESSIVE AGENTS**

### **AZATHIOPRINE**

50MG TABLET			
00004596	IMURAN	ASN	FNQSW
02236819	TEVA-AZATHIOPRINE	TEV	FGNQSW
02242907	APO-AZATHIOPRINE	APX	FGNQSW

### **BARICITINIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

2MG TABLET			
02480018	OLUMIANT (SA)	LIL	NMQW

### **CLADRIBINE**

[SEE APPENDIX A](#) FOR SA CRITERIA

10MG TABLET			
02470179	MAVENCLAD (SA)	EMD	NMQW
00904524	MAVENCLAD (SA)*		NMQW
00904525	MAVENCLAD (SA)*		NMQW
00904526	MAVENCLAD (SA)*		NMQW

\*use when drug cost in excess of CPHA maximum

### **CYCLOSPORINE**

10MG CAPSULE			
02237671	NEORAL	NVR	FNQSTW
25MG CAPSULE			
02150689	NEORAL	NVR	FNQSTW
02247073	SANDOZ-CYCLOSPORINE	SDZ	FGNQSTW
02495805	CYCLOSPORINE	STR	FGNQSTW
50MG CAPSULE			
02150662	NEORAL	NVR	FNQSTW

02247074	SANDOZ-CYCLOSPORINE	SDZ	<b>FGNQSTW</b>
02495821	CYCLOSPORINE	STR	<b>FGNQSTW</b>

100MG CAPSULE

02150670	NEORAL	NVR	<b>FNQSTW</b>
02242821	SANDOZ-CYCLOSPORINE	SDZ	<b>FGNQSTW</b>
02495813	CYCLOSPORINE	STR	<b>FGNQSTW</b>

100MG/ML ORAL SOLUTION

02150697	NEORAL	NVR	<b>FNQSTW</b>
02244324	APO-CYCLOSPORINE	APX	<b>FGNQSTW</b>

**MYCOPHENOLATE MOFETIL**

250MG CAPSULE

02192748	CELLCEPT	HLR	T
02320630	SANDOZ-MYCOPHENOLATE	SDZ	T
02352559	APO-MYCOPHENOLATE	APX	T
02364883	TEVA-MYCOPHENOLATE	TEV	T
02383780	MYCOPHENOLATE MOFETIL	ACH	T
02386399	JAMP-MYCOPHENOLATE	JPC	T
02457369	MYCOPHENOLATE MOFETIL	SNS	T

500MG TABLET

02237484	CELLCEPT	HLR	T
02313855	SANDOZ-MYCOPHENOLATE	SDZ	T
02352567	APO-MYCOPHENOLATE	APX	T
02348675	TEVA-MYCOPHENOLATE	TEV	T
02378574	MYCOPHENOLATE MOFETIL	ACH	T
02380382	JAMP-MYCOPHENOLATE	JPC	T
02457377	MYCOPHENOLATE MOFETIL	SNS	T

**MYCOPHENOLATE SODIUM**

180MG ENTERIC-COATED TABLET

02264560	MYFORTIC	NVR	T
02372738	APO-MYCOPHENOLIC ACID	APX	T
02511673	MAR-MYCOPHENOLIC ACID	MAR	T

360MG ENTERIC-COATED TABLET

02264579	MYFORTIC	NVR	T
02372746	APO-MYCOPHENOLIC ACID	APX	T
02511681	MAR-MYCOPHENOLIC ACID	MAR	T

**NINTEDANIB**

[SEE APPENDIX A](#) FOR SA CRITERIA

100MG CAPSULE

02443066	OFEV (SA)	BOE	<b>NMQW</b>
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150MG CAPSULE			
02443074	OFEV (SA)	BOE	NMQW

**PIRFENIDONE**

[SEE APPENDIX A](#) FOR SA CRITERIA

267MG CAPSULE			
02393751	ESBRIET (SA)	HLR	NMQW
02488833	SANDOZ-PIRFENIDONE (SA)	SDZ	NMQW
02509938	JAMP-PIRFENIDONE (SA)	JPC	NMQW

267MG TABLET

02464489	ESBRIET (SA)	HLR	NMQW
02488507	SANDOZ-PIRFENIDONE (SA)	SDZ	NMQW
02514702	JAMP-PIRFENIDONE (SA)	JPC	NMQW
02531526	PMS-PIRFENIDONE (SA)	PMS	NMQW
02537753	AURO-PIRFENIDONE (SA)	ARO	NMQW

801MG TABLET

02464500	ESBRIET (SA)	HLR	NMQW
02488515	SANDOZ-PIRFENIDONE (SA)	SDZ	NMQW
02514710	JAMP-PIRFENIDONE (SA)	JPC	NMQW
02531534	PMS-PIRFENIDONE (SA)	PMS	NMQW
02537761	AURO-PIRFENIDONE (SA)	ARO	NMQW

**SIROLIMUS**

1MG/ML ORAL SOLUTION			
02243237	RAPAMUNE	PFI	T

1MG TABLET			
02247111	RAPAMUNE	PFI	T

**TACROLIMUS**

0.5MG CAPSULE			
02243144	PROGRAF	AST	T
02416816	SANDOZ-TACROLIMUS	SDZ	T
02454068	ACH-TACROLIMUS	ACH	T

1MG CAPSULE			
02175991	PROGRAF	AST	T
02416824	SANDOZ-TACROLIMUS	SDZ	T
02456095	ACH-TACROLIMUS	ACH	T

5MG CAPSULE			
02175983	PROGRAF	AST	T
02416832	SANDOZ-TACROLIMUS	SDZ	T

02456109	ACH-TACROLIMUS	ACH	T
0.5MG EXTENDED RELEASE CAPSULE			
02296462	ADVAGRAF	AST	T
1MG EXTENDED RELEASE CAPSULE			
02296470	ADVAGRAF	AST	T
3MG EXTENDED RELEASE CAPSULE			
02331667	ADVAGRAF	AST	T
5MG EXTENDED RELEASE CAPSULE			
02296489	ADVAGRAF	AST	T
0.75MG EXTENDED RELEASE TABLET			
02485877	ENVARUSUS PA	END	T
1MG EXTENDED RELEASE TABLET			
02485885	ENVARUSUS PA	END	T
4MG EXTENDED RELEASE TABLET			
02485893	ENVARUSUS PA	END	T

## **92:92 OTHER MISCELLANEOUS AGENTS**

### **SAPROPTERIN**

[SEE APPENDIX A](#) FOR SA CRITERIA

100 MG POWDER FOR ORAL SOLUTION

02534533 REDDY-SAPROPTERIN (SA)

96599937 SAPROPTERIN (SA)\*

RCH **MNQW**  
**MNQW**

500 MG POWDER FOR ORAL SOLUTION

02535610 REDDY-SAPROPTERIN (SA)

96599936 SAPROPTERIN (SA)\*

RCH **MNQW**  
**MNQW**

\*use when drug cost in excess of CPHA maximum

## **PROFESSIONAL SERVICES**

### **MEDICATION REVIEW**

93899926	BASIC MEDICATION REVIEW	<b>DSW</b>
93899924	BASIC MEDICATION REVIEW FOLLOW-UP	<b>DSW</b>
93899925	DIABETES MEDICATION REVIEW	<b>DW</b>

93899923

DIABETES MEDICATION REVIEW FOLLOW-UP

**DW**

**OTHER SERVICES**

93899914

COMPLIANCE PACKAGING

**DFSW**

93899916

THERAPEUTIC SUBSTITUTION

**FGNSW**

93899917

REFUSAL TO FILL

**FLNSW**

93899918

PRESCRIPTION ADAPTATION

**DFGMNSVWZ**



## APPENDIX A Special Authorization Criteria

### NOTES REGARDING SPECIAL AUTHORIZATION (SA) COVERAGE

The following prescribers are permitted to submit and sign special authorization requests:

- medical practitioners and nurse practitioners
- hospital and community pharmacists for medications affiliated with a Common Ailment assessment (as per *Pharmacist and Pharmacy Technician Regulations*)
- Hospital pharmacists for medications prescribed under a practice directive approved by a regulatory licensing body in approved settings.
  
- Special Authorizations are reviewed by drug program staff.
  
- Not all medications currently approved for sale in Canada will be considered for Special Authorization coverage.
  
- Special Authorization coverage will not be considered for any medications approved for sale in Canada since January 2000 that have not been reviewed, and approved, for coverage by either the Canadian Expert Drug Advisory Committee (CEDAC), the Pan-Canadian Oncology Drug Review (P-CODR) or the Atlantic Expert Advisory Committee (AEAC).
  
- Special Authorization coverage will normally only be approved for the treatment of indications and in dosages listed in the official product monograph approved by Health Canada and published in the most recent edition of the Compendium of Pharmaceuticals and Specialities (CPS).
  
- Special Authorization coverage will potentially be considered for any drug not listed as an open benefit under the:
  - Family Health Benefit Drug Program
  - Financial Assistance Program
  - Nursing Home / Institutional Program
  - Seniors Drug Program
  
- Special Authorization coverage will be limited to selected drugs with specific criteria under the:
  - HIV Program
  - Diabetes Drug Program
  - Generic Drug Program
  - High Cost Drug Program
  - Home Oxygen Program
  - Substance Use Harm Reduction Drug Program
  - Transplant Drugs Program
  
- Special Authorization coverage will not be considered under the:
  - Community Mental Health Program
  - Cystic Fibrosis Program
  - Eprex Program
  - Growth Hormone Program
  - Hepatitis Program

- Phenylketonuria Program
  - Smoking Cessation Program
  - Sexually Transmitted Diseases Program
  - Tuberculosis Program
- Prescribers may apply for Special Authorization coverage by mailing or faxing a completed Special Authorization to:
    - Special Authorizations
    - PEI Pharmacare
    - P.O. Box 2000
    - Charlottetown, PEI, C1A 7N8
    - Fax: 1-902-368-4905
- Information that must be completed on, or included with the Special Authorization includes:
    - Patient's name, personal health number (PHN), date of birth, mailing address, and telephone number;
    - Name, dose, and dosage regimen of the medication requested;
    - Anticipated length of therapy of the medication requested;
    - Specific diagnosis or indication being treated using the medication requested;
    - Reason(s) for the request;
    - Other comments, including copies of culture and sensitivity reports for antibiotic requests, copies of relevant test results and relevant advice received from consultants or specialists; and
    - Prescriber's name, address, and signature. **No request will be considered without a valid prescriber's signature.**
- Special Authorizations with insufficient information to properly assess the request will be returned to the prescriber.
  - Please allow up to three weeks for the processing of Special Authorizations.
  - Copies of the Special Authorization Forms are available by contacting the PEI Pharmacare office at 1-877-577-3737 or online at <http://www.princeedwardisland.ca/pharmacareforms>.
  - For some drugs a patient application is required in addition to the Special Authorization form. The patient application form is available by contacting the PEI Pharmacare office at 1-877-577-3737 or online at <http://www.princeedwardisland.ca/pharmacareforms>.
  - Patients and prescribers are notified by letter if coverage has been approved. Patients should take a copy of the approval letter to their pharmacy to initiate coverage.
  - The duration of approval of Special Authorization coverage may range from a one time only fill to coverage with no end date. This will be based upon the medication requested and the condition being treated.

- Medications approved through the Special Authorization process are limited to a maximum 30 (thirty) day supply per fill unless otherwise noted in drug criteria.
- If additional information is required **or** if the request is denied, a letter is sent to the patient and prescriber notifying them of the need for additional information **or** reason for the denial. Payment of the medication is the responsibility of the patient in these cases.
- If the request is approved, patients may be reimbursed for one fill of the medication received during the assessment period, after which all of the requested information has been received. **No reimbursement will be provided for medication received by the patient prior to receipt of the Special Authorization by the Drug Programs Office.**
- If it is anticipated that a patient will continue to require the product beyond the last day of approval, the prescriber is required **to request an extension of coverage at least four weeks before its expiration.** Coverage will not be continued automatically.

## CRITERIA FOR COVERAGE OF SPECIFIC MEDICATIONS

The following are criteria for Special Authorization coverage of specific medications. Coverage may be granted for other products in certain instances.

### **Abatacept, vial, 250mg; syringe, 125mg/mL; syringe, 125mg/mL (Orencia-BMS)**

Maximum IV adult dose is 500mg for patients < 60kg, 750mg for patients 60 to 100kg, 1000mg for patients > 100kg, given at 0, 2, 4, 8 weeks and every 4 weeks thereafter. Pediatric patients 6-17 years of age and < 75kg, coverage is for IV dose 10mg/kg based on weight at administration (pediatric patients > 75kg to be treated at adult dose) given at 0, 2, 4, 8 weeks and every 4 weeks thereafter.

For adult Orencia-naïve patients, a single loading dose of up to 1000mg, then 125mg sc injection should be give within a day, and once weekly thereafter.

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks;

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Abemaciclib, tablet, 50mg, 100mg, 150mg (Verzenio-LIL)**

In combination with endocrine therapy (ET) for the adjuvant treatment of adult patients with hormone receptor (HR) positive, human epidermal growth factor receptor 2 (HER2) negative, node-positive early breast cancer at high risk of disease recurrence and a Ki-67 score of at least 20%.

Clinical Notes:

- Patients should have a good performance status.
- Treatment should continue until disease progression, unacceptable toxicity, or completion of 2 years of adjuvant therapy. ET may be continued after abemaciclib is completed.
- Patients are not eligible if they have inflammatory breast cancer, or prior treatment with a cyclin-dependent kinases 4 and 6 (CDK4/6) inhibitor.

- Retreatment with a CDK4/6 inhibitor may be reasonable in the metastatic setting if disease recurrence occurs greater than or equal to 6 months after completion of adjuvant abemaciclib.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Abiraterone, tablet, 250mg, 500mg (Zytiga-JAN and generics)**

1. In combination with prednisone for the treatment of metastatic prostate cancer (castration-resistant prostate cancer) in patients who are asymptomatic or mildly symptomatic after failure of androgen deprivation therapy or have received prior chemotherapy containing docetaxel after failure of androgen deprivation therapy.
2. In combination with androgen deprivation therapy (ADT) for the treatment of patients with metastatic castration sensitive prostate cancer who have had no prior ADT, or are within 6 months of beginning ADT, in the metastatic setting.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Abilify Maintena** – see Aripiprazole

**Abrilada** – see Adalimumab

**Abrocitinib, tablet, 50mg, 100mg, 200mg (Cibinqo-PFI)**

For the treatment of moderate to severe atopic dermatitis (AD) in patients 12 years of age and older who meet all the following criteria:

- Patients must have had an adequate trial (with a documented refractory disease, including the relief of pruritis), or were intolerant (with documented intolerance), or are ineligible for each of the following therapies:
  - Maximally tolerated medical topical therapies for AD combined with phototherapy (where available), and
  - Maximally tolerated medical topical therapies for AD combined with at least 1 of the 4 systemic immunomodulators (methotrexate, cyclosporine, mycophenolate mofetil, or azathioprine)

- Baseline Physician Global Assessment score of 3 or greater and Eczema Area and Severity Index (EASI) of 7.1 or greater.

Renewal Criteria:

- Requests for renewal must provide proof of beneficial clinical effect defined as a 75% or greater improvement from baseline in the EASI score (EASI-75) six months after treatment initiation.
- Proof of maintenance of EASI-75 response from baseline must be provided for subsequent authorizations.

Clinical Note:

- Not to be used in combination with phototherapy or any immunomodulatory agents (including biologics or other janus kinase inhibitor treatment) for moderate to severe AD. Treatment should continue until disease progression or unacceptable toxicity.

Claim Notes:

- The patient must be under the care of a dermatologist, allergist, clinical immunologist, or pediatrician who has expertise in the management of moderate to severe AD.
- Approvals will be for a maximum of 200 mg once daily.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Acalabrutinib, capsule, 100 mg ; tablet, 100mg (Calquence-AZE)**

1. As monotherapy for adult patients with previously untreated chronic lymphocytic leukemia (CLL) / small lymphocytic lymphoma (SLL) for whom fludarabine-based treatment is inappropriate due to high-risk cytogenetic markers (i.e., del17p, TP53 mutation, or unmutated IGHV).
2. As monotherapy for adult patients with relapsed or refractory CLL / SLL who have received at least one prior therapy.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.

2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Claim Notes:**

Requests will not be considered for patients who experience disease progression on a Bruton's tyrosine kinase (BTK) inhibitor or idelalisib.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Acamprosate, delayed release, tablet, 333mg (Campral-MYL)**

Note: For Substance Use Harm Reduction Drug Program, no Special Authorization is required.

For the treatment of alcohol use disorder.

**Actemra** – see Tocilizumab

**Actonel 30mg** - see Risedronate

**Adalimumab, 20mg/0.2mL prefilled syringe (Hyrimoz-SDZ); 20mg/0.4mL prefilled syringe (Abrilada-PFI, Amgevita-AMG, Hulio-BGP, Hyrimoz-SDZ); 40mg/0.4mL prefilled syringe, 40mg/0.4mL pen injector (Hadlima-MER, Hyrimoz-SDZ, Simlandi-JPC, Yuflyma-LIL) 40mg/0.8mL prefilled syringe (Abrilada-PFI, Amgevita-AMG, Hadlima-MER, Hulio-BGP, Hyrimoz-SDZ, Idacio-FKB), 40mg/0.8mL prefilled pen (Abrilada-PFI, Amgevita-AMG, Hadlima-MER, Hulio-BGP, Hyrimoz-SDZ, Idacio-FKB), 80mg/0.8mL prefilled syringe (Hyrimoz-SDZ, Simlandi-JPC, Yuflyma-CLT), 80mg/0.8mL pen injector (Hyrimoz-SDZ, Yuflyma-CLT)**

For Adalimumab naïve patients, approved requests will be for a biosimilar product.

**Ankylosing Spondylitis**

For the treatment of patients with moderate to severe ankylosing spondylitis (Bath AS Disease Activity Index (BASDAI) score  $\geq 4$  on 10 point scale) who:

- a) have axial symptoms\* and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation or in whom NSAIDs are contraindicated OR

- b) have peripheral symptoms and who have failed to respond to, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.

\*Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease, do not require a trial of NSAIDs alone.

Claim Notes:

- Approvals will be for a maximum of 40 mg every two weeks.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial Approval: 6 months
- Renewal Approval: maximum of 12 months. Requests for renewal must include information showing the beneficial effects of the treatment, specifically:
  - a) a decrease of at least two points on the BASDAI scale, compared with pretreatment score OR
  - b) patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as Health Assessment Questionnaire (HAQ) or ability to return to work).

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Ankylosing Spondylitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Crohn's Disease**

For the treatment of patients with moderate to severe Crohn's disease who have active disease and are refractory, intolerant or have contraindications to:

- Prednisone 40mg (or equivalent) daily for ≥ 2 weeks, AND
- Azathioprine ≥ 2 mg/kg/day for ≥ 3 months, OR
- Mercaptopurine ≥ 1 mg/kg/day for ≥ 3 months, OR
- Methotrexate (SC or IM) ≥ 15 mg/week for ≥ 3 months

Clinical Notes:

- Refractory is defined as lack or loss of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.



- Consideration will be given for the approval of a biologic DMARD (disease modifying antirheumatic drug) without a trial of a traditional DMARD for patients who have an aggressive/severe disease course (e.g. extensive disease, a modified Harvey Bradshaw Index score > 16) and are refractory, intolerant or have contraindications to systemic corticosteroids.

Claim notes:

- Initial 12 week approval for an induction dose of 160mg followed by 80mg two weeks later, then 40mg every two weeks thereafter. Renewal of coverage will require reassessment of the patient and submission of a new Crohn's Disease Special Authorization form. Continued coverage may be approved at a dose not exceeding 40mg every 2 weeks.
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a gastroenterologist using the Crohn's Disease Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Hidradenitis Suppurative**

For the treatment of patients with active moderate to severe hidradenitis suppurativa (HS) who have not responded to conventional therapy and who meet all of the following criteria:

- A total abscess and nodule count of 3 or greater
- Lesions in at least two distinct anatomic areas, one of which must be Hurley Stage II or III
- An inadequate response to a 90-day trial of oral antibiotics

Initial renewal criteria:

- Requests for renewal should provide objective evidence of a treatment response, defined as at least a 50% reduction in abscess and inflammatory nodule count with no increase in abscess or draining fistula count relative to baseline at week 12.

Subsequent renewal criteria:

- Requests for renewal should provide objective evidence of the preservation of treatment effect (i.e. the current abscess and inflammatory nodule count and

draining fistula count should be compared to the count prior to initiating treatment with adalimumab).

Claim Notes:

- Approvals will be for a maximum of 160mg followed by 80mg two weeks later, then 40mg every week beginning four weeks after the initial dose.
- Must be prescribed by a dermatologist or physician with experience in the treatment of HS.

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Plaque Psoriasis**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20$ mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Concurrent use of biologics not approved.
- Approvals will be for a maximum dose of 80 mg administered once, followed by 40 mg after 1 week of initial dose, then 40mg every other week thereafter up to 16 weeks. If response criteria is met at 16 weeks, approval will be continued at a dose of 40 mg every two weeks up to one year.
- Initial approval: 16 weeks.
- Renewal approval: 1 year

**The request must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Polyarticular Juvenile Idiopathic Arthritis (pJIA)**

For the treatment of pJIA for patients aged 4-17 years with moderately or severe pJIA who have had an inadequate response to one or more disease-modifying anti-rheumatic drugs (DMARDs). Treatment must be initiated by a rheumatologist who is familiar with the use of DMARDs and/or biologic DMARDs in children.

Claim Notes:

- Approvals will be for a maximum dose of 40mg every two weeks
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and
  - Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and

- Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum dose of 40mg every two weeks
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Rheumatoid Arthritis**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks

AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks;

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.

- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Approvals will be for a maximum dose of 40mg every two weeks
- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ulcerative Colitis**

For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score > 4, and a rectal bleeding subscore ≥ 2 and are:

- Refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks AND prednisone ≥ 40mg daily for two weeks or IV equivalent for one week) OR
- Corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.

- Initial Approval: 8 week approval is for an induction dose of 160 mg followed by 80 mg two weeks later, then 40 mg every two weeks thereafter. Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease in the partial Mayo score  $\geq 2$  from baseline, and
  - a decrease in the rectal bleeding subscore  $\geq 1$ .
- Renewal Approval: 1 year. Maximum approved dose is 40 mg every two weeks.
- Combined use of more than one biologic DMARD will not be reimbursed.

Clinical Notes:

- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.
- Patients with severe disease (partial Mayo > 6) do not require a trial of 5-ASA

**The request for coverage must be made by a gastroenterologist using the Ulcerative Colitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Uveitis**

For the treatment of patients with non-infectious uveitis who are refractory, intolerant or have contraindications to conventional therapy.

Claim Notes:

- Must be prescribed by, or in consultation with an ophthalmologist.
- Approvals will be for a maximum of 80 mg followed by 40mg in one week, then 40 mg every two weeks thereafter
- Combined use of more than one biologic DMARD will not be reimbursed.

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Adempas** – see Riociguat

**Advair** - see Salmeterol & Fluticasone

**Advair Diskus** - see Salmeterol & Fluticasone

**Afatinib, tablet, 20mg, 30mg, 40mg (Giotrif-BOE)**

For the first line treatment of patients with EGFR mutation positive advanced or metastatic adenocarcinoma of the lung and with an ECOG performance status of 0 or 1.

**NOTE**

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

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Aflibercept, vial, 2mg/0.5mL (Eylea-BAY)**

**1. Neovascular Age-Related Macular Degeneration:**

Criteria For Initial Coverage:

For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD) where all of the following apply:

- a) Best Corrected Visual Acuity (BCVA) is between 6/12 and 6/96 **AND**
- b) The lesion size is less than or equal to 12 disc areas in greatest linear dimension **AND**
- c) There is evidence of recent (<3 months) presumed disease progression (blood vessel growth, as indicated by fluorescein angiography, optical coherence tomography (OCT), or recent visual acuity changes. The interval between doses should not be shorter than one month. Administration is to be done by a qualified ophthalmologist experienced in intravitreal injections.

Criteria For Continued Coverage:

Treatment with aflibercept should be continued only in people who maintain adequate response to therapy.

Aflibercept should be discontinued if any of the following occur:

- a) Reduction in BCVA in the treated eye to less than 15 letters (absolute) on 2 consecutive visits in the treated eye, attributed to AMD in the absence of other pathology Or

- b) 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 OR
- c) There is evidence of deterioration of the lesion morphology despite optimum treatment over 3 consecutive visits.

Coverage will not be approved for patients:

- a) Receiving concurrent treatment with verteporfin.
- b) With permanent retinal damage as defined by the Royal College of Ophthalmology guidelines.

Coverage is limited to a maximum of one vial per eye in any 30-day period. The request for coverage must be made by an ophthalmologist.

Approval Period: 1 year

## **2. Diabetic macular edema (DME)**

Initial coverage:

For the treatment of visual impairment due to diabetic macular edema (DME) in patients who meet all of the following criteria:

- clinically significant centre-involving macular edema for whom laser photocoagulation is also indicated
- hemoglobin A1c test in the past 6 months with a value of less than or equal to 11%
- best corrected visual acuity of 20/32 to 20/400
- central retinal thickness greater than or equal to 250 micrometers

Renewal Criteria:

- confirm that a hemoglobin A1c test in the past 6 months had a value of less than or equal to 11%
- date of last visit and results of best corrected visual acuity at that visit
- date of last OCT and central retinal thickness on that examination
- if aflibercept is being administered monthly, please provide details on the rationale

Clinical Notes:

1. Treatment should be given monthly until maximum visual acuity is achieved (i.e. stable visual acuity for three consecutive months). Thereafter, visual acuity should be monitored monthly.
2. Treatment should be resumed when monitoring indicates a loss of visual acuity due to DME and continued until stable visual acuity is reached again for three consecutive months.
3. Treatment should be discontinued if there is no improvement of retinal thickness or visual acuity after five consecutive treatments.



4. Injection will be by a qualified ophthalmologist with experience in administering intravitreal injections.

Approval Period: 1 year

### **3. Retinal vein occlusion (RVO)**

For the treatment of visual impairment due to macular edema secondary to central retinal vein occlusion (CRVO) or branch retinal vein occlusion (BRVO).

Clinical Notes:

1. Treatment should be given monthly until maximum visual acuity is achieved (i.e. stable visual acuity for three consecutive months). Thereafter, visual acuity should be monitored monthly.
2. Treatment should be resumed 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 months.
3. Treatment should be discontinued if there is no improvement after 6 months of initial treatment.
4. Injection will be by a qualified ophthalmologist with experience in administering intravitreal injections.

Approval Period: 1 year

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Agalsidase alfa** – see Replagal

**Agalsidase beta** – see Fabrazyme

**Agrylin** - see Anagrelide

**Ajovy** – see Fremanezumab

**Akynzeo** – see Netupitant & Palonosetron

**Alecensaro** – see Alectinib

### **Alectinib, capsule, 150mg (Alecensaro-HLR)**

For the treatment of patients with anaplastic lymphoma kinase (ALK)-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer

when used:

- as first-line therapy, or
- following disease progression on, or intolerance to, crizotinib.

Renewal Criteria:

- Confirmation that the patient is responding to treatment.

Claim Notes:

- Requests for alectinib will not be considered for patients who experience disease progression on any ALK inhibitor other than crizotinib.
- No further ALK inhibitor will be reimbursed following disease progression on alectinib.
- Initial approval period: 1 year.
- Renewal approval period: 1 year

\* Claims that exceed the maximum claim amount of \$9999.99 must be divided as separate transactions, using DIN first and PDIN\* second.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Alertec** - see Modafinil

**Alirocumab, prefilled pen, 75mg/mL, 150mg/mL (Praluent-SAV)**

For the treatment of heterozygous familial hypercholesterolemia (HeFH) in adult patients who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following criteria are met:

- Definite or probable diagnosis of HeFH using the Simon Broome or Dutch Lipid Network criteria or genetic testing; and
- Patient is unable to reach LDL-C target (less than 2.0 mmol/L or at least a 50% reduction in LDL-C from untreated baseline) despite confirmed adherence to at least 3 months of continuous treatment with:  
high-dose statin (e.g. atorvastatin 80 mg, rosuvastatin 40 mg) in combination with ezetimibe; or—ezetimibe alone, if high dose statin is not possible due to rhabdomyolysis, contraindication or intolerance.

**Initial renewal criteria:**

- A reduction in LDL-C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L.

**Subsequent renewal criteria:**

- The patient continues to maintain a reduction in LDL-C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L.

**Clinical Notes:**

1. LDL-C levels must be provided.
2. Intolerance to high dose statin will be considered if patient has developed documented myopathy or abnormal biomarkers (i.e. creatinine kinase greater than 5 times the upper limit of normal) after trial of at least two statins and—for each statin, dose reduction was attempted rather than statin discontinuation, and intolerance was reversible upon statin discontinuation, but reoccurred with statin re-challenge where clinically appropriate; and
  - at least one statin was initiated at the lowest daily starting dose; and
  - other known causes of intolerance have been ruled out.
3. For patients who cannot take ezetimibe due to an intolerance or contraindication, details must be provided.

**Claim Notes:**

- Approvals will be for a maximum of 300mg every 4 weeks.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**Alunbrig** – see Brigatinib

**Ambrisentan, 5mg, 10mg (Volibris-GSK and generics)**

For treatment of patients with pulmonary arterial hypertension (PAH), of at least World Health Organization (WHO) functional class III, which is associated with either idiopathic or connective tissue disease and who have failed to respond to or who have contraindications to, or who are not a candidate for sildenafil.

**Clinical Notes:**

1. Diagnosis of PAH should be confirmed by cardiac catheterization
2. Ambrisentan will not be approved when used concurrently with other endothelin receptor antagonists, epoprostenol, treprostinil or sildenafil.
3. Coverage must be requested and medications prescribed by a Pulmonary Arterial Hypertension (PAH) specialist.

**Claim Note:**

The maximum dose of ambrisentan that will be reimbursed is 10mg daily

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at**

<http://www.princeedwardisland.ca/pharmacareforms>

**Amerge** - see Naratriptan HCl

**Amgevita** – see Adalimumab

**Amifampridine, tablet, 10mg (Ruzurgi-MDU)**

For the treatment of Lambert-Eaton myasthenic syndrome (LEMS) in patients 6 years of age or older.

Initial Renewal Criteria:

- An improvement of at least 30% on the Triple Timed Up and Go (3TUG) test compared to baseline measurement.

Subsequent Renewal Criteria:

- The patient continues to maintain an improvement of at least 30% on the 3TUG test compared to baseline measurement.

Clinical Note:

1. The 3TUG test score must be provided with initial and renewal requests.

Claim Notes:

1. Must be prescribed by a neurologist.
2. Approvals will be up to a maximum daily dose of 40 mg for patients weighing less than 45 kg and 100 mg for patients weighing 45 kg or more.
3. Initial approval period: 3 months. Renewal approval period: 1 year.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Amifampridine phosphate, tablet, 10mg (Firdapse-KYE)**

For the treatment of Lambert-Eaton myasthenic syndrome (LEMS) in patients 18 years of age and older.

Initial Renewal Criteria:

- An improvement of at least 30% on the Triple Timed Up and Go (3TUG) test compared to baseline measurement.

Subsequent Renewal Criteria:

- The patient continues to maintain an improvement of at least 30% on the 3TUG test compared to baseline measurement.

Clinical Note:

1. The 3TUG test score must be provided with initial and renewal requests.

Claim Notes:

1. Must be prescribed by a neurologist.
2. Approvals will be up to a maximum daily dose of 80mg.
3. Initial Approval: 3 months. Renewal approval period: 1 year.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Amlodipine, oral solution, 1mg/mL (Generic)**

- For patients who require administration through a feeding tube.
- For patients 19 years of age and younger, who cannot use a tablet or capsule.
- Pediatric patients 12 and under will not require written Special Authorization.

**Anagrelide, capsule, 0.5mg (Agrylin-SHR and generics)**

For the treatment of essential thrombocythemia (ET) in patients who have:

- a) Failed Hydroxyurea therapy (does not provide sufficient platelet reduction) or
- b) Have intolerable side effects to Hydroxyurea therapy.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Anifrolumab, vial, 150mg/mL (Saphnelo-AZE)**

For the treatment of adult patients with moderate to severe autoantibody positive, systemic lupus erythematosus (SLE) who meet all of the following criteria:

- Systemic lupus erythematosus disease activity index 2000 (SLEDAI-2K) score of 6 or greater.
- Refractory to oral corticosteroids (OCS) at a dose of at least 10 mg per day of prednisone or its equivalent, in addition to standard of care.

Renewal criteria:

- OCS dose has decreased to less than or equal to 7.5 mg per day of prednisone or its equivalent OR OCS dose decreased by at least 50% from baseline; and
- Reduction in disease activity as measured by:
  - Reduction in the SLEDAI-2K index score to 5 or less; or
  - British Isles lupus assessment group (BILAG)-2004 index score improvement in involved organ systems and no new worsening in other organ systems.

Subsequent renewal criteria:

- Initial response achieved after the first twelve months of treatment with anifrolumab has been maintained.

Clinical notes:

- Standard of care is defined as using an immunosuppressive drug (e.g., rituximab, hydroxychloroquine, mycophenolic acid, or azathioprine) with or without NSAIDs.
- A baseline SLEDAI-2K must be provided. If BILAG-2004 is used for assessment on renewal, then a baseline BILAG-2004 assessment of organ systems must also be provided. The same scale should be used on all subsequent renewals.
- Improvement in organ systems is defined as a reduction of all severe BILAG-2004 A) or moderately severe BILAG-2004 B) to lower rating levels.
- Worsening in organ systems is defined as at least one new BILAG-2004 A item or at least two new BILAG-2004 B items.

Exclusion criteria:

- Severe or unstable neuropsychiatric SLE.
- Active severe SLE nephritis.

Claim notes:

- Patient should be under the care of a physician with expertise in the diagnosis and management of SLE.
- Combined use with other biologic drugs will not be reimbursed.
- Approvals will be for a maximum of 300 mg every four weeks.
- Approval period: 1 year.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**AndroGel-** see Testosterone

**Anoro Ellipta –** see Umeclidinium Bromide & Vilanterol Trifenatate

**Apalutamide, tablet, 60mg, 240mg (Erleada-LIL)**

1. In combination with androgen deprivation therapy (ADT) for the treatment of patients with castration-resistant prostate cancer (CRPC) who have no detectable distant metastasis (M0) by either CT, MRI or technetium-99m bone scan and who are at high risk of developing metastases<sup>1</sup>.  
Patients should have a good performance status and no risk factors for seizures. Treatment should continue until unacceptable toxicity or radiographic disease progression.

Clinical Notes:

- Castration-resistance must be demonstrated during continuous ADT and is defined as 3 PSA rises at least one week apart, with the last PSA > 2 ng/mL.
- Castrate levels of testosterone must be maintained.
- Patients with N1 disease, pelvic lymph nodes < 2cm in short axis located below the common iliac vessels are eligible for apalutamide.
- Apalutamide will not be funded for patients who experience disease progression on enzalutamide.
- Patients receiving apalutamide for the treatment of non-metastatic CRPC will be eligible for funding of abiraterone at the time of disease progression to metastatic CRPC. Enzalutamide is not funded for patients who experience disease progression to metastatic CRPC while on apalutamide.
- Either abiraterone or enzalutamide may be used to treat metastatic CRPC in patients who discontinued apalutamide in the non-metastatic setting due to intolerance without disease progression.

<sup>1</sup> High risk of developing metastases is defined as a prostate-specific antigen (PSA) doubling time of ≤ 10 months during continuous ADT

2. In combination with androgen deprivation therapy (ADT) for the treatment of patients with metastatic castration-sensitive prostate cancer (mCSPC). Patients must have had either no prior ADT, or are within six months of beginning ADT in the metastatic setting.

Clinical Notes:

- Patients should have a good performance status and no risk factors for seizures.
- Treatment should continue until unacceptable toxicity or disease progression.

Claim Notes:

- Patients receiving apalutamide for the treatment of metastatic CSPC will be eligible for funding of abiraterone at the time of disease progression to metastatic CRPC.
- Enzalutamide is not funded for patients who experience disease progression to metastatic CRPC while on apalutamide.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Aprepitant, capsule, 80mg, 125mg, 80mg & 125mg package (Emend, Emend Tri-Pack)**

In combination with dexamethasone for the prevention of acute and delayed nausea and vomiting in patients receiving:

- highly emetogenic chemotherapy or
- moderately emetogenic chemotherapy who have had inadequate symptom control using a 5-HT<sub>3</sub> antagonist and dexamethasone in a previous cycle.

Clinical Notes:

- Highly emetogenic chemotherapy (HEC) includes but it not limited to : cisplatin regimens, anthracycline and cyclophosphamide combination regimens, and regimens containing carmustine, mechlorethamine, streptozocin, dacarbazine, and cyclophosphamide > 1500mg/m<sup>2</sup>
- Patients who receive carboplatin-based regimens with AUC ≥ 4 are also eligible to receive aprepitant in combination a 5-HT3 antagonist and dexamethasone for primary prevention of acute and delayed nausea and vomiting

**Aptiom** – see Eslicarbazepine Acetate

**Aptivus** – see Tipranavir

**Aranesp** - see Darbepoetin Alfa

**Aripiprazole, injection, 300mg, 400mg (Abilify Maintena-OTS)**

Note: For Community Mental Health Drug Program, no Special Authorization is required.

For the treatment of schizophrenia in patients with documented compliance issues with an oral antipsychotic OR who are currently receiving a conventional depot antipsychotic and experiencing significant side effects (EPS or TD) or lack of efficacy.

NOTE: Must be requested and prescribed by a psychiatrist. Only doses up to 400mg monthly will be approved.

*In accordance with the manufacturer's product monograph:*

For patients who have never taken aripiprazole, establish tolerability with oral aripiprazole prior to initiating treatment with Abilify Maintena.

**Asciminib, tablet, 20mg, 40mg (Scemblix-NVR)**

For the treatment of adult patients with Philadelphia chromosome-positive chronic myeloid leukemia (Ph+ CML) in the chronic phase who meet the following criteria:

- Treatment failure on or intolerance to a minimum of two prior tyrosine kinase inhibitor (TKI) therapies.
- No evidence of a T315I or V299L mutation.

Clinical Notes:

1. Patients should have a good performance status.
2. Not for use in the acute phase or blast phase.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**



**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Asenapine, sublingual tablet, 5mg, 10mg (Saphris-MSD)**

For the acute treatment of manic or miXPI episodes associated with bipolar I disorder as either:

- Monotherapy, after a trial of lithium or divalproex sodium has failed, and trials of less expensive atypical antipsychotic agents have failed due to intolerance or lack of response.
- Co-therapy with lithium or divalproex sodium, after trials of less expensive atypical antipsychotic agents have failed due to intolerance or lack of response.

**Atectura Breezhaler** – see Indacaterol & Mometasone

**Atogepant, tablet, 10mg, 30mg 60mg (Qulipta-ABV)**

For the prevention of migraine in patients with a confirmed diagnosis of episodic migraine who have experienced an inadequate response, intolerance, or contraindication to at least two classes of oral prophylactic migraine medications.

Renewal Criteria:

- A reduction of at least 50% in the average number of migraine days per month at the time of initial renewal compared with baseline.
- At subsequent renewals, the patient continues to maintain the reduction of at least 50% in average number of migraine days per month.

Clinical Notes:

- The average number of headache and migraine days per month must be provided on initial and renewal requests.
- According to the International Headache Society criteria, episodic migraine is defined as:
  - migraine headaches on at least 4 days per month and less than 15 headache days per month for more than 3 months.
- Atogepant should not be reimbursed for use in combination with other CGRP inhibitors for the prevention of migraine in adult patients with episodic migraine.

Claim Notes:

- Initial approval: 6 months
- Renewal approval: 1 year

**Aubagio** – see Multiple Sclerosis Agents

**Avonex** - see Multiple Sclerosis Agents

**Avsola** – see Infliximab

**Axitinib, tablet, 1 mg, 5 mg (Inlyta-PFI)**

For the treatment of patients with advanced or metastatic renal cell carcinoma when used as:

- As first-line therapy in combination with pembrolizumab; or
- Second-line therapy following disease progression on a vascular endothelial growth factor receptor tyrosine kinase inhibitor (i.e., sunitinib or pazopanib); or
- Third-line therapy following disease progression on first-line nivolumab and ipilimumab combination therapy and a second-line vascular endothelial growth factor receptor tyrosine kinase inhibitor.
- Patients must have a good performance status.
- Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Sequential use of axitinib and everolimus is not permitted except in the case of intolerability or contraindication.
- Sequential use of axitinib (as a single agent) and cabozantinib is not permitted for patients following progression on first-line axitinib + pembrolizumab.
- For patients treated with nivolumab + ipilimumab first-line and VEGFR TK1 second line, either cabozantinib or axitinib may be used as third-line therapy.
- Both clear cell and non-clear cell histology are eligible for treatment.
- Approval period: 1 year

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Azacitidine, tablet, 200mg, 300mg (Onureg-CEL)**

As maintenance therapy for adult patients with acute myeloid leukemia (AML) who meet all of the following criteria:

- Intermediate or poor risk cytogenetics
- Complete remission (CR) or complete remission with incomplete blood count recovery (CRi) following induction therapy, with or without consolidation treatment
- Not eligible for hematopoietic stem cell transplantation (HSCT)

Clinical Notes:

1. Newly diagnosed includes patients with AML de novo or secondary to prior myelodysplastic syndrome (MDS) or chronic myelomonocytic leukemia (CMML).
2. Last dose of chemotherapy should be within 4 months of starting azacitidine maintenance.
3. Treatment should be discontinued upon disease relapse (i.e., appearance of greater than 5% blasts in the bone marrow or peripheral blood), unacceptable toxicity, or if patient becomes eligible for allogeneic bone marrow or stem cell transplant during the treatment period.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Azithromycin, tablet, 250mg, 600mg; oral suspension, 20mg/mL, 40mg/mL (Zithromax-PFI and generics)**

Note: For HIV, Cystic Fibrosis, Sexually Transmitted Diseases, and Tuberculosis Programs, no Special Authorization is required.

- a) For the treatment of infections requiring a macrolide antibiotic when the patient has a documented intolerance to clarithromycin
- b) For the completion of hospital initiated treatment with azithromycin (maximum 5 days)
- c) For the treatment and prevention of non-tuberculosis mycobacterial
- d) For the treatment of infections requiring a macrolide antibiotic when the patient is taking medications that would significantly interact with erythromycin/clarithromycin

**Aztreonam, inhalation vial, 75mg/mL (Cayston-GIL)**

For the treatment of chronic pulmonary *Pseudomonas aeruginosa* infections, when used as a cyclic treatment, in patients with moderate to severe cystic fibrosis and deteriorating clinical condition despite treatment with inhaled tobramycin.

**Clinical Note:**

Cyclic treatment measured in 28-day cycles is defined as 28 days of treatment, followed by 28 days without treatment.

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Baqsimi** – see Glucagon

**Baraclude** – see Entecavir

### **Baricitinib, tablet, 2mg (Olumiant-LIL)**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

#### Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

#### Claim Notes:

- Approvals will be for a maximum adult dose of 2 mg daily.
- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

### **Benralizumab, syringe, autoinjector, 30mg/mL (Fasenra-AZN)**

As add-on maintenance treatment for adult patients with severe eosinophilic asthma, if the following criteria are met:

Initiation Criteria:

- Patient must have a documented diagnosis of asthma.
- Patient is inadequately controlled with high-dose inhaled corticosteroids, defined as greater or equal to 500mcg of fluticasone propionate or equivalent daily, and one or more additional asthma controller(s) (e.g., long-acting beta agonists).
- Patient has one of the following:
  - blood eosinophil count of  $\geq 300$  cells/ $\mu$ L within the past 12 months AND has experienced two or more clinically significant asthma exacerbations in the past 12 months, or
  - blood eosinophil count of  $\geq 150$  cells/ $\mu$ L AND is receiving maintenance treatment with oral corticosteroids (OCS).

Renewal Criteria:

- The effects of treatment should be assessed every 12 months to determine whether reimbursement should continue
- Reimbursement of treatment should be discontinued if:
  - the 12 months asthma control questionnaire score has not improved from baseline, when baseline represents the initiation of treatment, or
  - the asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
  - the number of clinically significant exacerbations has increased within the previous 12 months, or
  - in patients on maintenance treatment with OCS, there has been no decrease in the OCS dose in the first 12 months of treatment, or
  - in patients on maintenance treatment with OCS, the reduction in the dose of OCS achieved after the first 12 months of treatment is not maintained subsequently.

Clinical Notes:

- Benralizumab should not be used in combination with other biologics used to treat asthma.
- A baseline assessment of asthma symptom control using a validated asthma control questionnaire must be completed prior to initiation of benralizumab treatment.
- Patients should be managed by a physician with expertise in treating asthma.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Benzydamine HCl, oral rinse, 0.15% (Generic)**

For oncology patients only.

**Betahistine HCL, tablet, 16mg, 24mg (Serc-BGP and generics)**

For the symptomatic treatment of recurrent episodes of vertigo associated with Meniere's disease.

**Betaseron** - see Multiple Sclerosis Agents

**Beovu** – see Brovacumab

**Bimekizumab, prefilled syringe, 160mg/mL; autoinjector, 160mg/mL (Bimzelx-UCB)**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score ; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Concurrent use of biologics not approved.
- Approvals will be for 320mg given every 4 weeks for 16 weeks then 320 mg every 8 weeks thereafter
- Initial approval: 12 weeks.

- Renewal approval: 1 year.

**The request must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Bimzelx** – see Bimekizumab

**Biphentin** - see Methylphenidate

**Binimetinib, tablet, 15mg (Mektovi-PFI)**

For the treatment of patients with BRAF V600 mutation-positive locally advanced unresectable or metastatic melanoma when used in combination with encorafenib.

**Renewal Criteria:**

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

**Clinical Notes:**

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Claim Notes:**

- Binimetinib will not be reimbursed in patients who have progressed on BRAF targeted therapy
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Bisacodyl, suppository (water based), 10mg (Magic Bullet)**

For the treatment of bowel incontinence where alternative therapies have failed.

For use as part of a bowel program for neurogenic bowel dysfunction in patients with spinal cord injuries.

**Bosentan, tablet, 62.5mg, 125mg (Tracleer-ACT and generics)**

For treatment of pulmonary arterial hypertension (PAH) in patients with World Health Organization (WHO) functional class III or IV.

Clinical Notes:

- Idiopathic pulmonary arterial hypertension (IPAH) in patients who do not demonstrate vasoreactivity on testing or who demonstrate vasoreactivity on testing but fail a trial of, or are intolerant to, calcium channel blockers.
- Pulmonary arterial hypertension associated with connective tissue disease or congenital heart disease or human immunodeficiency virus (HIV) who do not respond adequately to conventional therapy.

Coverage must be requested and medications prescribed by a Pulmonary Arterial Hypertension (PAH) specialist.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Bosulif** – see Bosutinib

**Bosutinib, tablet, 100mg, 500mg (Bosulif-PFI)**

For treatment of adult patients with chronic, accelerated, or blast phase Philadelphia chromosome-positive (Ph+) chronic myelogenous leukemia (CML) with resistance or intolerance to prior TKI therapy.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Braftovi** – see Encorafenib

**Breztri Aerosphere** – see Budesonide Glycopyrronium Formoterol

**Brigatinib, tablet, 30mg, 90mg, 180mg; initiation kit, 90mg (7) & 180mg (21) (Alunbrig-TAK)**



For the treatment of adult patients with anaplastic lymphoma kinase (ALK)-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer who have not been previously treated with an ALK inhibitor.

**Renewal Criteria:**

- Written confirmation that the patient is responding to treatment.

**Clinical Note:**

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

**Claim Notes:**

- No further ALK inhibitor will be reimbursed following disease progression on brigatinib.
- Initial approval period: 1 year.
- Renewal approval period: 1 year

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Brilinta** – see Ticagrelor

**Brivaracetam, tablet, 10mg, 25mg 50mg, 75mg, 100mg (Brivlera-UCB)**

For the treatment of partial onset seizures in patients with epilepsy who are not satisfactorily controlled with conventional therapy if the following clinical criteria and conditions are met:

1. Patients are currently receiving two or more antiepileptic drugs (AEDs).
2. Patients are not receiving concurrent therapy with levetiractam.
3. Patients are those for whom less costly AEDs are ineffective or not clinically appropriate.

**Brivlera** – see Brivaracetam

**Breo Ellipta** – see Fluticasone Furoate/Vilanterol

**Brenzys** - see Etanercept

**Brolucizumab, prefilled syringe, 6 mg/0.05 mL (Beovu-NVR)**

**1. Neovascular Age-Related Macular Degeneration**

Initial Coverage:

For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD).

Discontinuation Criteria:

- Reduction in Best Corrected Visual Acuity (BCVA) in the treated eye of 15 letters or more on 2 consecutive visits, attributed to AMD in the absence of other pathology, or
- Reduction in BCVA in the treated eye of 30 letters or more compared to either baseline and/or best recorded level, or
- There is evidence of deterioration of the lesion morphology despite optimum treatment over 3 consecutive visits.

Clinical Note:

- BCVA must be provided with initial request and with subsequent renewal requests.

Claim Notes:

- Approvals will be for a maximum of 1 prefilled syringe per eye every 4 weeks for 12 weeks, followed by 1 prefilled syringe per eye every 8 weeks thereafter.
- Approval Period: 1 year.

**2. Diabetic Macular Edema (DME)**

Initial coverage:

For the treatment of visual impairment due to diabetic macular edema (DME) in patients who meet all of the following criteria:

- clinically significant centre-involving macular edema for whom laser photocoagulation is also indicated
- hemoglobin A1c test in the past 6 months with a value of less than or equal to 11%
- best corrected visual acuity of 20/32 to 20/400
- central retinal thickness greater than or equal to 250 micrometers

Renewal Criteria:

- confirm that a hemoglobin A1C test in the past 6 months had a value of less than or equal to 11%
- date of last visit and results of best corrected visual acuity at that visit
- date of last OCT and central retinal thickness on that examination

Clinical Notes:

- Treatment should be given monthly until maximum visual acuity is achieved (i.e. stable visual acuity for three consecutive months). Thereafter, visual acuity should be monitored monthly.

- Treatment should be resumed when monitoring indicates a loss of visual acuity due to DME and continued until stable visual acuity is reached again for three consecutive months.
- Treatment should be discontinued if there is no improvement of retinal thickness or visual acuity after three consecutive treatments.
- Injection will be by a qualified ophthalmologist with experience in administering intravitreal injections.

Approval Period: 1 year

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

### **Brodalumab, syringe, 210mg/1.5mL (Siliq-VAL)**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score ; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Concurrent use of biologics not approved.

- Approvals will be for a maximum adult dose of 210mg administered at 0, 1 and 2 weeks followed by 210mg every 2 weeks. If response criteria is met at 16 weeks, approval will be continued at a dose of 210mg every two weeks up to one year.
- Initial approval period: 16 weeks
- Renewal approval period: 1 year

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Brukinsa** – see Zanubrutinib

**Budesonide, inhalation solution, 0.125mg/mL, 0.25mg/mL, 0.5mg/mL (Pulmicort Nebuamp-AZE and generics)**

Note: For Nursing Home Program, no Special Authorization is required.

- For use in patients on the Nursing Home Program.
- For use in children under 6 years of age. The pharmacy must call the drug programs office to have coverage set up initially. Coverage will be in place until the child's sixth birthday.
- Other uses will be considered on a case by case basis where there are extreme circumstances.

**Burosumab, vial, 10mg/mL, 20mg/ml, 30mg/mL (Crysvita-ULT)**

For the treatment of patients with X-linked hypophosphatemia (XLH) who meet the following criteria:

- Initiated in a pediatric patient who is at least one year of age and in whom epiphyseal closure has not yet occurred
- Fasting hypophosphatemia
- Normal renal function (defined as a serum creatinine below the age-adjusted upper limit of normal)
- Radiographic evidence of rickets with a rickets severity score (RSS) of two or greater
- Confirmed phosphate-regulating endopeptidase homolog, X-linked (PHEX) gene variant in either the patient or in a directly related family member with appropriate X-linked inheritance

Discontinuation Criteria:

In pediatric patients under 18 years of age in whom epiphyseal closure has not yet occurred and who met the above criteria, treatment should be discontinued if

- there is no demonstrated improvement in the 12-month RSS total score from baseline RSS total score; or
- the patient's RSS total score achieved after the first 12 months of therapy has not been maintained subsequently.

In adolescent patients who are 13 to 17 years of age in whom epiphyseal closure has occurred and who met the above criteria and initiated treatment as a pediatric patient, treatment should be discontinued if any of the following occur:

- Hyperparathyroidism; or
- Nephrocalcinosis; or
- Evidence of fracture or pseudo-fracture based on radiographic assessment.

In adult patients who met the above criteria and initiated treatment as a pediatric patient, treatment should be discontinued if any of the following occur:

- Hyperparathyroidism; or
- Nephrocalcinosis; or
- Evidence of fracture or pseudo-fracture based on radiographic assessment.

Claim Notes:

- Requests will not be considered for treatment-naïve adults
- Must be prescribed by a physician working in a multidisciplinary team of health care providers who are experienced in the diagnosis and management of XLH
- Approvals for children (1-17 years of age) will be up to a maximum of 90mg every 2 weeks
- Approvals for adults (18 years of age and older) will be up to a maximum of 90mg every 4 weeks.
- Approval period: 1 year.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Byooviz** – see Ranibizumab

**Cabenuva** – see Cabotegravir & Rilpivirine

**Cabometyx** – see Cabozantinib

**Cabotegravir, tablet, 30mg (Vocabria-VII)**

For the treatment of adult patients with HIV-1 infection who are virologically stable and suppressed (HIV-1 RNA less than 50 copies per mL).

## **Cabotegravir & Rilpivirine, vial, 400mg/600mg, 600mg/900mg (Cabenuva-VII)**

For the treatment of adult patients with HIV-1 infection who are virologically stable and suppressed (HIV-1 RNA less than 50 copies per mL).

## **Cabozantinib, tablet, 20mg, 40mg, 60mg (Cabometyx-IPS)**

### **Advanced or Metastatic Renal Cell Carcinoma (RCC)**

For the treatment of patients with advanced or metastatic renal cell carcinoma who have received at least one prior vascular endothelial growth factor receptor (VEGFR) tyrosine kinase inhibitor (TKI) therapy when used as:

- second-line therapy following disease progression on sunitinib, pazopanib or pembrolizumab in combination with axitinib; or
- third-line therapy following disease progression on immunotherapy and VEGFR TKI (i.e., sunitinib or pazopanib), used in any sequence.

#### Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of clinically meaningful disease progression.

#### Clinical Note:

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

#### Claim Notes:

- Requests for cabozantinib will not be considered for patients who experience disease progression on everolimus or axitinib monotherapy.
- Initial approval period: 1 year.
- Renewal approval period: 1 year

### **Differentiated Thyroid Carcinoma (DTC)**

For the treatment of adult patients with locally advanced or metastatic differentiated thyroid carcinoma (DTC) who have progressed on at least one prior line of vascular endothelial growth factor receptor (VEGFR)-targeted tyrosine kinase inhibitor (TKI) therapy.

#### Clinical Notes:

- Patients should have a good performance status.
- Patients should be refractory to radioactive iodine therapy (RAI-R) or not eligible for radioactive iodine therapy.
- Treatment should continue until disease progression or unacceptable toxicity.
- Patients will be eligible for funding if intolerant to the prior line of VEGFR-targeted TKI therapy.

- Cabozantinib may be used in the third line setting for RET fusion positive patients after progression on or intolerance to selpercatinib.

### **Unresectable Hepatocellular Carcinoma (HCC)**

For the second-line treatment of adult patients with unresectable hepatocellular carcinoma who meet all of the following criteria:

- Disease progression on sorafenib or lenvatinib
- Child-Pugh class status of A
- ECOG performance status of 0 or 1

Clinical Note:

- Treatment should continue until the patient no longer experiences clinical benefit or experiences unacceptable toxicity.

Claim Notes:

- Requests for cabozantinib will not be considered for patients who experience disease progression on regorafenib or atezolizumab in combination with bevacizumab.
- Approval period: 6 months

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Calquence** – see Acalabrutinib

**Campral** – see Acamprosate

### **Canagliflozin, tablet, 100mg, 300mg (Invokana-JAN)**

For the treatment of type 2 diabetes as a third drug added on to metformin and a sulfonylurea for patients with inadequate glycemic control on optimal doses of metformin and a sulfonylurea, and for whom insulin is not an option.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Canakinumab, vial, 150mg/mL (Ilaris-NVR)**

For the treatment of active systemic juvenile idiopathic arthritis, in patients 2 years of

age or older, who have an inadequate response or intolerance to systemic corticosteroids (with or without methotrexate) and tocilizumab.

**Clinical Note:**

- Intolerance is defined as a serious adverse effect as described in the product monograph. The nature of the intolerance(s) must be clearly documented.

**Claim Notes:**

- Must be prescribed by, or in consultation with, a rheumatologist, who is familiar with the use of biologic DMARDs in children.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Approvals will be for 4 mg/kg for patients > 9 kg, to a maximum of 300mg, administered every four weeks.
- Initial approval period: 16 weeks.
- Renewal approval period: 1 year. Confirmation of continued response is required

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Caprelsa** – see Vandetanib

**Carbamazepine, suspension, 100mg/5mL (Tegretol-NVR and generics)**

For use in patients for indications as defined in the CPS, and who cannot use carbamazepine chewable, regular and controlled release tablets.

**Carbidopa & Levodopa & Entacapone, tablet, 12.5mg/50mg/200mg, 25mg/100mg/200mg, 37.5mg/150mg/200mg, 18.75mg/75mg/200mg, 31.25mg/125mg/200mg (Stalevo-NVR)**

For the treatment of Parkinson's disease in patients who are not well controlled and are experiencing significant "wearing off" symptoms despite optimal therapy with levodopa/carbidopa and are currently stabilized on levodopa/carbidopa and entacapone separately.

**Carnitor** – see Levocarnitine

**Caripul** – see Epoprostenol

**Cayston** – see Aztreonam

**Ceritinib, capsule, 150mg (Zykadia-NVR)**

As monotherapy treatment for patients with anaplastic lymphoma kinase (ALK)-positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer who experience disease progression on, or intolerance to, crizotinib.



Renewal Criteria:

- Confirmation that the patient is responding to treatment.

Clinical Note:

- Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Certolizumab, syringe kit, 400mg/2mL; auto-injector kit, 400mg/2mL (Cimzia-UCB)**

**Ankylosing Spondylitis**

For the treatment of patients with moderate to severe ankylosing spondylitis (Bath AS Disease Activity Index (BASDAI) score  $\geq 4$  on 10 point scale) who:

- a) have axial symptoms\* and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation or in whom NSAIDs are contraindicated OR
- b) have peripheral symptoms and who have failed to respond to, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.

\*Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease, do not require a trial of NSAIDs alone.

Claim Notes:

- Approvals will be for a maximum adult dose of 400 mg (given as two SC injections of 200 mg) given at 0, 2, 4 weeks then 200 mg every two weeks thereafter.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial Approval: 6 months
- Renewal Approval: maximum of 12 months. Requests for renewal must include information showing the beneficial effects of the treatment, specifically:
  - a) a decrease of at least two points on the BASDAI scale, compared with pretreatment score OR
  - b) patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as Health Assessment Questionnaire (HAQ) or ability to return to work).

The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Ankylosing Spondylitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>

Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>

### **Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and
  - Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and
  - Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

#### Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

#### Claim notes:

- Approvals will be for a maximum adult dose of 400 mg (given as two SC injections of 200 mg) given at 0, 2, 4 weeks then 200 mg every two weeks thereafter.
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>

Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at

<http://www.princeedwardisland.ca/pharmacareforms>

### **Rheumatoid Arthritis**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

#### Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

#### Claim Notes:

- Approvals will be for a maximum adult dose of 400 mg (given as two SC injections of 200 mg) given at 0, 2, 4 weeks then 200 mg every two weeks thereafter.
- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at**

**<http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at**

**<http://www.princeedwardisland.ca/pharmacareforms>**

**Cesamet** - see Nabilone

**Chlorhexidine, oral rinse, 0.12% (Peridex-MDA, Perichlor-PMS)**

For the treatment of periodontal disease in long term care residents who need assistance in mouth care upon request or recommendation from a dentist. **A copy of the recommendation from the dentist may be required.**

**Cholinesterase Inhibitors (ChEI)**

**Galantamine, extended-release capsule, 8mg, 16mg, 24mg (Generics)**  
**Rivastigmine, capsule, 1.5mg, 3mg, 4.5mg, 6mg (Exelon-NVR and generics)**

For the treatment of patients with mild to moderate dementia who have had an intolerance to donepezil and who meet the following criteria:

- Mini-Mental State Exam (MMSE) score of 10 to 30

Clinical Notes:

1. Requests must contain an updated MMSE score completed within 6 months of the request.
2. The nature of the intolerance must be described.

Claim Note:

Approval period: 1 year

**Chronic Obstructive Pulmonary Disease Medications**

**Acclidinium Bromide, aerosol powder for inhalation, 400mcg/dose (Tudorza Genuair-ALM)**

**Acclidinium Bromide & Formoterol Fumarate Dyhydrate, aerosol powder, 400mcg/12mcg actuation (Duaklir Genuair-AZE)**

**Budesonide & Glycopyrronium & Formoterol, metered dose inhaler, 160mcg-7.2mcg-5mcg (Breztri Aerosphere-AZE)**

**Fluticasone & Umeclidinium & Vilanterol, dry powder for inhalation, 100mcg-62.5mcg-25mcg/dose (Trelegy Ellipta-GSK)**

**Fluticasone Furoate/Vilanterol, blister with inhalation device, 100mcg-25mcg/dose (Breo Ellipta-GSK)**

**Formoterol Fumerate, powder for inhalation (capsule), 12mcg/dose (Foradil-NVR); powder for inhalation (inhaler), 6mcg/dose, 12mcg/dose (Oxeze Turbuhaler-AZE)**

**Formoterol & Budesonide, powder for inhalation, 6mcg & 100mcg per dose, 6mcg & 200mcg per dose (Symbicort Turbuhaler-AZE)**

**Glycopyrronium Bromide, capsule for inhalation, 50mcg (Seebri Breezhaler-NVR)**

**Indacaterol, capsule, inhalation powder, 75mcg (Onbrez-NVR)**

**Indacaterol & Glycopyrronium powder for inhalation (capsule), 110mcg-50mcg (Ultibro Breezhaler – NVR)**

**Salmeterol Xinafoate, aerosol powder disk, 50µg/dose (Serevent Diskus-GSK)**  
**Salmeterol & Fluticasone, aerosol inhalation, 25mcg & 125mcg per dose, 25mcg & 250mcg per dose (Advair-GSK); inhaled powder disk, 50mcg & 100mcg per dose, 50mcg & 250mcg per dose, 50mcg & 500mcg per dose (Advair Diskus- GSK)**  
**Tiotropium, capsule for inhalation, 18mcg/dose (Spiriva-BOE); mist inhaler, 2.5mcg/dose (Spiriva Respimat-BOE)**  
**Tiotropium & Olodaterol mist inhaler, 2.5mcg-2.5mcg, (Inspiolto Respimat - BOE)**  
**Umeclidinium Bromide, blister with inhalation device, 62.5mcg (Incruse Ellipta-GSK)**  
**Umeclidinium Bromide & Vilanterol Trifenatate, blister, 62.5mcg/25mcg (Anoro Ellipta-GSK)**

<b>Table 1 (of 4)</b>	
Formoterol fumarate dehydrate (Oxeze Turbuhaler)	Acclidinium (Tudorza Genuair)
Formoterol fumarate (Foradil)	Glycopyrronium Bromide (Seebri)
Indacaterol maleate (Onbrez)	Tiotropium (Spiriva) 18mcg; (Spiriva Respimat) 2.5mcg
Salmeterol (Serevent)	Umeclidinium Bromide (Incruse Ellipta)

**For any one agent listed in Table 1:**

For the treatment of chronic obstructive pulmonary disease (COPD) as defined by spirometry<sup>i</sup> in patients

AND

- Experiencing persistent symptoms, as defined by Medical Research Council (MRC) score of at least 3<sup>ii</sup> or a COPD Assessment test (CAT) score ≥ 10<sup>iii</sup> and a post-bronchodilator FEV<sub>1</sub> <80% predicted
- OR
- Experiencing 2 or more moderate exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids OR at least 1 acute severe exacerbation of COPD (AECOPD) requiring hospitalization.

**NOTE:** Coverage for both a LABA and a LAAC as separate inhalers will not be considered. See below for combination LABA/LAAC coverage criteria.

<b>Table 2 (of 4)</b>
Acclidinium Bromide & Formoterol Fumarate Dihydrate (Duaklir Genuair)
Indacaterol/Glycopyrronium (Ultibro Breezhaler)
Tiotropium/Olodaterol (Inspiolto Respimat)

Umeclidinium Bromide & Vilanterol Trifenatate (Anoro Ellipta)

**For any one agent listed in Table 2:**

- For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry<sup>i</sup>, in patients with inadequate control<sup>iv</sup> with either a long-acting beta-2 agonist (LABA) or long-acting anticholinergic (LAAC).

Coverage for both a LABA and a LAAC as separate inhalers will not be considered. LABA/LAAC inhalers are not intended to be used in combination with an inhaled corticosteroid (ICS) unless criteria for triple therapy<sup>v</sup> is fulfilled.

<b>Table 3 (of 4)</b>
Budesonide/formoterol (Symbicort)
Fluticasone/umeclidinium/vilanterol (Trelegy Ellipta)
Fluticasone/vilanterol (Breo Ellipta)
Salmeterol /fluticasone (Advair)

**For any one agent listed in Table 3:**

- For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry<sup>i</sup>  
AND
- When the LABA/ICS is part of triple therapy<sup>v</sup> in patients with COPD  
OR
- In patients with asthma/COPD (ACO) overlap, based on patient history and lung function studies indicating an ACO diagnosis

<b>Table 4 (of 4)</b>
Budesonide/glycopyrronium/formoterol (Breztri Aerosphere)

**For agent listed in Table 4:**

- For the treatment of chronic obstructive pulmonary disease (COPD), as defined by spirometry<sup>i</sup>, in patients who experience inadequate control<sup>iv</sup> while being treated with a long-acting beta-2 agonist/long-acting muscarinic antagonist (LABA/LAMA).

Patients should not be started on a LABA, LAMA and an inhaled corticosteroid (triple inhaled therapy) as initial therapy.

**Clinical Notes:**

- (i) COPD is defined by spirometry as a post bronchodilator FEV<sub>1</sub>/FVC ratio of < 0.70. Spirometry reports from any point in time will be accepted.
- (ii) MRC Grade 3 is described as: walks slower than people of the same age on the level because of shortness of breath from COPD or has to stop for breath when walking at own pace on the level because of COPD.
- (iii) The COPD assessment test (CAT) is an 8-item tool for measuring health status impairment with scores from 0-40. It is available online at <http://www.catestonline.org/images/pdfs/CATest.pdf>
- (iv) Inadequate control is defined as persistent symptoms after at least 1 month of long- acting beta-agonist (LABA) or long-acting anticholinergic therapy (LAAC); and an MRC score of at least 3 or a CAT score ≥ 10.
- (v) Triple therapy criteria:  
Combination therapy with LABA/LAAC/ICS will be considered for patients who experience inadequate control (persistent symptoms or experiencing 2 or more exacerbations of COPD in the previous year requiring treatment with antibiotics and/or systemic corticosteroids or at least 1 exacerbation requiring hospitalization) while being treated with a LABA/LAAC combination for at least two months.
- (vi) Prescriptions written by PEI respirologists do not require written Special Authorization.

**Cibinqo** – see Abrocitinib

**Ciloxan** - see Ciprofloxacin, ophthalmic solution

**Cimzia** – see Certolizumab

**Cinacalcet, tablet, 30mg, 60mg 90mg (Generics)**

For the treatment of dialysis patients with severe hyperparathyroidism (PTH > 88 pmol/L measured twice in 3 months at least 6 weeks apart) who have maximized phosphate binder therapy and vitamin D therapy.

Patients must have one of the following:

- corrected serum calcium > 2.54mmol/L; serum phosphate > 1.8mmol/L; **or**
- presence of symptoms related to hyperparathyroidism (i.e. bone pain)

**Ciprodex** - see Ciprofloxacin & Dexamethasone

**Ciprofloxacin, ophthalmic solution, 0.3%; ophthalmic ointment, 0.3% (Ciloxan-ALC and generics)**

For the treatment of ophthalmic infections caused by susceptible bacteria and not responding to alternative agents.

**Ciprofloxacin HCl, tablet, 250mg, 500mg, 750mg; (Generics) oral suspension, 100mg/mL (Cipro-BAY)**

Note: For Cystic Fibrosis, Nursing Home and Tuberculosis Programs, no Special Authorization is required.

- For treatment of complicated urinary tract infections (UTI), early pyelonephritis, or bacterial prostatitis.
- For treatment of severe (malignant) otitis externa
- For empiric treatment of acute exacerbations of chronic obstructive pulmonary disease (AECOPD) in patients at risk of Pseudomonas infection (e.g. previously isolated Pseudomonas, end stage lung disease, concomitant bronchiectasis, frequent or recent broad spectrum antibiotic use).
- For empiric treatment of outpatient febrile neutropenia.
- For the prevention of endophthalmitis in patients who have had cataract surgery with unplanned vitrectomy.
- For treatment of lung infections in patients with cystic fibrosis.
- Pseudomonas aeruginosa susceptible disease (or if previous Pseudomonas susceptible disease).
- For the oral treatment of multi-resistant, aerobic, gram-negative infections traditionally requiring parenteral therapy for which other oral agents are not effective or available.
- For the treatment of patients intolerant or allergic (hypersensitivity reaction) to all other effective oral agents.
- For the empiric treatment of peritonitis in patients currently receiving peritoneal dialysis.

**Ciprofloxacin & Dexamethasone, otic suspension, 0.3% / 0.1% (Ciprodex-ALC and generics)**

- For the treatment of patients with acute otitis media with otorrhea through tympanostomy tubes; or with known or suspected tympanic membrane perforation with otorrhea.
- For the treatment of patients with acute otitis externa in the presence of tympanostomy tubes or known perforation of the tympanic membrane.



### **Cladribine, tablet, 10mg (Mavenclad-EMD)**

For the treatment of adult patients with relapsing-remitting multiple sclerosis (RRMS) who meet all the following criteria:

- Confirmed diagnosis based on McDonald criteria.
- Has experienced one or more disabling relapses or new MRI activity in the past year.
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5).
- Refractory or intolerant to at least one disease modifying therapy (e.g., interferon, glatiramer, dimethyl fumarate, teriflunomide, ocrelizumab).

#### **Clinical Notes:**

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.
- A relapse is defined as the appearance of new or worsening neurological symptoms in the absence of fever or infection, lasting at least 24 hours yet preceded by stability for at least one month and accompanied by new objective neurological findings observed through evaluation by a neurologist.

#### **Claim Notes:**

- Must be prescribed by a neurologist with experience in the treatment of multiple sclerosis.
- Approvals will be for 1.75mg/kg to a maximum of 200mg per treatment year.
- Approval period: 2 years

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Clozapine, tablet, 25mg, 50mg, 100mg, 200mg (Clozaril-NVR and generics)**

Clozapine is only available upon registration of the patient, prescriber, and pharmacy with a Clozapine-Support and Assistance Network.

Clozapine is only to be dispensed to patients upon receipt of 7 day, 14 day or 28 day hematological test results by the pharmacy.

For the treatment of patients with schizophrenia refractory to other treatments upon written request or recommendation of a psychiatrist. **A copy of the recommendation must accompany the Special Authorization.**

**Clozaril** - see Clozapine

### **Cobimetinib – tablet, 20mg (Cotellic-HLR)**

In combination with vemurafenib, for the treatment of patients with previously untreated

BRAF V600 mutation-positive unresectable stage III or stage IV melanoma who have a good performance status. Treatment should continue until unacceptable toxicity or disease progression. If brain metastases are present, patients should be asymptomatic or have stable symptoms.

Approvals are for a maximum daily dose of 60mg during 21 consecutive days per 28 day cycle.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Codeine, controlled release tablet, 50mg, 100mg, 150mg, 200mg (Codeine Contin-PFR)**

For the treatment of documented mild to moderate chronic pain that is not well controlled by short-acting codeine products or where patients are well controlled on acetaminophen or ASA combinations but the codeine dose is limited by the amount of acetaminophen or ASA. **The maximum dose of Codeine Contin that will be reimbursed is 200mg every 12 hours.**

**Codeine Contin** - see Codeine

**Cosentyx** – see Secukinumab

**Cotellic** – see Cobimetinib

**Cresemba** – see Isavuconazole

**Crizotinib, capsule, 200mg, 250mg (Xalkori-PFI)**

1. For the treatment of patients with anaplastic lymphoma kinase (ALK)-positive locally advanced non-small cell lung cancer (NSCLC) with an ECOG performance status  $\leq 2$  when used as:
  - a) first line therapy or
  - b) second line therapy following chemotherapy
2. For the first-line treatment of patients with ROS-1 positive non-small cell lung cancer (NSCLC).

Clinical Notes:

- Eligible patients should be previously untreated and have a good performance status.
- Treatment may continue until disease progression or unacceptable toxicity.

- Patients with ROS-1 positive NSCLC who are currently receiving first-line chemotherapy or have been previously treated with chemotherapy or immunotherapy will be eligible for treatment with crizotinib.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Crysvita** – see Burosumab

**Cyclobenzaprine, tablet, 10mg (Generics)**

As an adjunct to rest and physical therapy for relief of muscle spasm associated with acute, painful musculoskeletal conditions not responding or experiencing severe adverse reactions to alternative therapy. **A maximum of three weeks (21 days) of therapy will be considered.**

**Cyclosporine, ophthalmic emulsion, 0.1% (Verkazia-SNN)**

For the treatment of pediatric patients between the age of 4 and 18 years of age with severe vernal keratoconjunctivitis (VKC) who meet the following criteria:

- Grade 3 (severe) or 4 (very severe) on the Bonini scale, *OR*
- Grade 4 (marked) or 5 (severe) on the modified Oxford scale.

Discontinuation Criteria:

- Treatment should be discontinued if no improvement in signs and symptoms of VKC is observed, *OR*
- Treatment should be discontinued if signs and symptoms of VKC have resolved.

Clinical Note:

- Documentation of the severity of signs and symptoms of VKC at treatment initiation and renewal must be provided.

Claim Notes:

- The patient must be under the care of a physician experienced in the diagnosis and treatment of VKC.
- Initial approval period: 6 months.
- Renewal approval period: 1 year

**Dabigatran, capsule, 110mg, 150mg (Pradaxa-BOE and generic)**

For the prevention of stroke and systemic embolism in at-risk patients with non-valvular atrial fibrillation for whom:

- a) Anticoagulation is inadequate following at least a two month trial of warfarin; or
- b) Warfarin is contraindicated or not possible due to inability to regularly monitor through International Normalized Ratio (INR) testing (i.e. no access to INR testing services at a laboratory, clinic, pharmacy, and at home).

The following patient groups are **excluded** from coverage for dabigatran for atrial fibrillation:

- a) Patients with impaired renal function (creatinine clearance or estimated glomerular filtration rate < 30mL/min)
- b) Patients 75 years of age or older without documented stable renal function
- c) Patients with hemodynamically significant rheumatic valvular heart disease, especially mitral stenosis
- d) Patients with prosthetic heart valves

Notes:

1. At-risk patients with atrial fibrillation are defined as those with a CHADS<sub>2</sub> score of  $\geq 1$ .
2. Inadequate anticoagulation is defined as INR testing results that are outside the desired INR range for at least 35% of the tests during the monitoring period (i.e. adequate anticoagulation is defined as INR test results that are within the desired INR range for at least 65% of the tests during the monitoring period).
3. Since renal impairment can increase bleeding risk, renal function should be regularly monitored. Other factors that increase bleeding risk should also be assessed and monitored (see dabigatran product monograph).
4. Documented stable renal function is defined as creatinine clearance or estimated glomerular filtration rate that maintained for at least three months (i.e. 30-49 mL/min for 110 mg twice daily dosing or  $\geq 50$  mL/min for 150 mg twice daily dosing).
5. There is currently no data to support that dabigatran provides adequate anticoagulation in patients with rheumatic valvular disease or those with prosthetic heart valves, so dabigatran is not recommended in these populations.
6. Patients starting dabigatran should have ready access to appropriate medical services to manage a major bleeding event.

## **Dabrafenib, capsule, 50mg, 75mg (Tafinlar-NVR)**

### **Adjuvant Melanoma**

In combination with trametinib for the adjuvant treatment of patients with cutaneous melanoma who meet all of the following criteria:

- Stage IIIA (limited to lymph node metastases of greater than 1 mm) to stage IIID disease (AJCC 8th edition)
- BRAF V600-mutation positive
- Completely resected disease including in-transit metastases

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should continue until disease recurrence, unacceptable toxicity, or up to a maximum of 12 months.

Claim Notes:

1. Requests will be considered for patients with regional lymph nodes with micrometastases after sentinel lymph node biopsy.
2. Requests will not be considered for patients who received adjuvant immunotherapy for greater than three months. Patients may switch to BRAF targeted therapy within the first three months of initiating immunotherapy to complete a total of 12 months of adjuvant treatment.
3. Approval period: up to 12 months

**Metastatic Melanoma**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used alone or in combination with trametinib.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Dalteparin** - see Low Molecular Weight Heparins

**Darbepoetin Alfa, pre-filled syringe 25mcg/mL, 40mcg/mL, 100mcg/mL, 200mcg/mL (Aranesp-AMG)**

For the treatment of severe anemia related to chronic renal failure in patients with:

- a) Normocytic normochromic anemia, requiring transfusions in patients who have evidence of iron overload (Ferritin > 1000 ng/mL), **OR**
- b) Anemia requiring blood transfusions in patients having symptomatic angina and/or heart failure, **OR**
- c) Anemia requiring transfusion in patients with difficulties in blood grouping and febrile reactions due to antibodies, **OR**

- d) Anemia requiring transfusions in patients who have high levels of panel reactive anti HLA antibodies, **OR**
- e) Severe normocytic normochromic anemia (Hb < 100 g/L) whose only symptom is fatigue and have never received transfusions.

**The request for coverage must be made by or in consultation with a nephrologist, internal medicine specialist, or oncologist. A copy of the consultation must accompany the request.**

**The request for coverage must be made using the Erythropoietin Program Approval Form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Darifenacin, extended release tablet, 7.5mg, 15mg (Enablex-NVR and generic)**

For the treatment of over-active bladder (not stress incontinence) in patients who cannot tolerate or have an insufficient response to an adequate trial (e.g. 3 months) of immediate release oxybutynin, solifenacin, or tolterodine.

**Darolutamide, tablet, 300mg (Nubeqa-BAY)**

**Non-Metastatic Castration-Resistant Prostate Cancer (nmCRPC)**

In combination with androgen deprivation therapy (ADT) for the treatment of patients with non-metastatic castration-resistant prostate cancer (nmCRPC) who are at high risk of developing metastases<sup>1</sup>.

- Patients should have a good performance status. Treatment should continue until unacceptable toxicity or radiographic disease progression.

**Clinical Notes:**

- Castration-resistance must be demonstrated during continuous ADT and is defined as 3 PSA rises at least one week apart, with the last PSA > 2 ng/mL.
- Patients should have no detectable distant metastases by either CT, MRI or technetium-99m bone scan.
- Castrate levels of testosterone must be maintained.
- Patients with N1 disease, pelvic lymph nodes < 2cm in short axis located below the aortic bifurcation are eligible for darolutamide.
- Darolutamide will not be funded for patients who experience disease progression on apalutamide or enzalutamide.
- Patients receiving darolutamide for the treatment of non-metastatic CRPC will be eligible for funding of abiraterone at the time of disease progression to metastatic CRPC. Enzalutamide is not funded for patients who experience disease progression to metastatic CRPC while on darolutamide.

- Either abiraterone or enzalutamide may be used to treat metastatic CRPC in patients who discontinued darolutamide in the non-metastatic setting due to intolerance without disease progression.

<sup>1</sup>High risk of developing metastases is defined as a prostate-specific antigen (PSA) doubling time of  $\leq 10$  months during continuous ADT.

### **Metastatic Castration-Sensitive Prostate Cancer (mCSPC)**

In combination with docetaxel and androgen deprivation therapy (ADT) for the treatment of patients with metastatic castration-sensitive prostate cancer (mCSPC). Patients must have had either no prior ADT or are within six months of beginning ADT in the metastatic setting.

Renewal Criteria:

- Confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

- Patients should have a good performance status.
- Treatment should continue until unacceptable toxicity or disease progression.

Claim Notes:

- Requests will not be considered for patients who are within 1 year of completing adjuvant ADT in the nonmetastatic setting.
- Patients who experience disease progression on apalutamide or enzalutamide are not eligible.
- Approval period: 1 year.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Dasatinib, tablet, 20mg, 50mg, 70mg, 80mg, 100mg, 140mg (Sprycel-BMS and generics)**

For use as a single agent for the treatment of adults with chronic, accelerated or blast phase chronic myelogenous leukemia (CML) and Philadelphia chromosome acute lymphoblastic leukemia (Ph+ ALL) with resistance or intolerance to prior therapy including Imatinib.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a**

patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

**DDAVP** - see Desmopressin

**Decitabine & Cedazuridine, tablet, 35mg & 100mg (Inqovi-TAI)**

For the treatment of patients with myelodysplastic syndromes (MDS), including previously treated and untreated, who meet all of the following criteria:

- De novo or secondary MDS including all French-American-British subtypes (i.e., refractory anemia, refractory anemia with ringed sideroblasts, refractory anemia with excess blasts, refractory anemia with excess blasts in transformation, and chronic myelomonocytic leukemia)
- Intermediate-1, intermediate-2, or high-risk MDS, according to the International Prognostic Scoring System
- Have not experienced disease progression on a hypomethylating agent

Clinical Notes:

1. Patients should have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Deferasirox, tablet, 90mg, 180mg, 360mg; dispersible tablet, 125mg, 250mg, 500mg (Exjade – NVR and generics)**

For the treatment of patients who require iron chelation.

**Denosumab, pre-filled syringe, 60mg/mL (Prolia – AMG)**

For the treatment of osteoporosis in patients who have:

- A high fracture risk, and
- a contraindication, severe gastrointestinal intolerance, or are refractory to bisphosphonates.

Clinical Notes:

- Refractory is defined as a fragility fracture or evidence of a decline in bone mineral density below pre-treatment baseline levels, despite adherence for one year to other available osteoporosis therapies.
- High fracture risk is defined as:



- Moderate 10-year fracture risk (10% to 20%) as defined by the Canadian Association of Radiologists and Osteoporosis Canada (CAROC) tool or the World Health Organization's Fracture Risk Assessment (FRAX) tool with a prior fragility fracture; or
- High 10-year fracture risk ( $\geq 20\%$ ) as defined by the CAROC or FRAX tool.

**Desmopressin, oral disintegrating tablet, 60mcg, 120mcg, 240mcg (DDAVP Melt-FEI); tablet, 0.1mg, 0.2mg (DDAVP-FEI and generics)**

1. For the treatment of diabetes insipidus in patients unable to tolerate the intranasal solution or when the intranasal solution is ineffective.
2. For the treatment of enuresis in children over 5 years and under 16 years of age.

**Desmopressin, intranasal solution (spray pump), 10mcg/dose (Generic)**

For the treatment of diabetes insipidus. The maximum recommended daily dosage is 40µg.

**Diacomit** – see Stiripentol

**Dienogest, tablet, 2mg (Visanne-BAY and generic)**

For the management of pelvic pain associated with endometriosis in patients for whom one or more less costly options are either ineffective or cannot be used.

**Dificid** – see Fidaxomicin

**Dojolvi** – see Triheptanoin

**Dornase Alfa, inhalation solution, 1mg/mL (Pulmozyme-HLR)**

For cystic fibrosis patients with a FEV1 < 70% predicted with clinically significant decline in FEV1 not responsive to usual treatment.

**Duaklir Genuair** – see Acclidinium Bromide & Formoterol Fumarate Dihydrate

**Duobrii** – see Tazarotene/Halobetasol propionate

**Duodopa** – see Levodopa/Carbidopa

**Dupilumab, syringe, prefilled pen, 200mg/1.14mL; syringe, prefilled pen, 300mg/2mL (Dupixent-AVN)**

**Atopic Dermatitis**

For the treatment of moderate to severe atopic dermatitis (AD) in patients 12 years of age and older who meet all of the following criteria:

Patients must have had an adequate trial (with a documented refractory disease), or were intolerant (with documented intolerance), or are ineligible for each of the following therapies:

- maximally tolerated medical topical therapies for AD combined with phototherapy (where available), and;
- maximally tolerated medical topical therapies for AD combined with at least 1 of the 4 systemic immunomodulators (methotrexate, cyclosporine, mycophenolate mofetil, or azathioprine).

AND

Baseline Physician Global Assessment score of 3 or greater and Eczema Area and Severity Index (EASI) of 7.1 or greater.

Renewal criteria:

- Requests for renewal must provide proof of beneficial clinical effect defined as a 75% or greater improvement from baseline in the Eczema Area and Severity Index (EASI-75) score six months after treatment initiation.
- Proof of maintenance of EASI-75 response from baseline must be provided for subsequent authorizations.

Clinical Note:

- Not to be used in combination with phototherapy or any immunomodulatory drugs (including biologics) or a Janus kinase inhibitor treatment for moderate-to-severe AD.

Claim Notes:

- The patient must be under the care of a dermatologist, allergist, clinical immunologist, or pediatrician who has expertise in the management of moderate to severe AD.
- Approvals will be for a maximum of 600 mg at week 0, then 300 mg every two weeks thereafter.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**Asthma**

1. For the adjunctive treatment of severe asthma with a type 2 or eosinophilic phenotype in patients aged 6 to 11 years of age who are inadequately controlled with medium-to high-dose inhaled corticosteroids (ICS) and one or more additional asthma controller(s) (e.g., long-acting beta-agonist) or high-dose ICS alone and meet the following criteria:

- blood eosinophil count  $\geq 0.15 \times 10^9/L$  within the past 12 months; and
- uncontrolled asthma with at least one clinically significant asthma exacerbation in the past 12 months.

Initial Discontinuation Criteria:

- Baseline asthma control questionnaire score has not improved at 12 months since initiation of treatment, or
- The number of clinically significant asthma exacerbations has increased within the previous 12 months

Subsequent Discontinuation Criteria:

- Asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
- The number of clinically significant asthma exacerbations has increased within the previous 12 months.

Clinical Notes:

1. A baseline and annual assessment of asthma symptom control using a validated asthma control questionnaire must be provided.
2. Medium dose ICS is defined as between 200 mcg and 400 mcg of fluticasone propionate or equivalent daily dose and high-dose ICS is defined as greater than 400 mcg of fluticasone propionate or equivalent daily dose.
3. A significant clinical exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least 3 days or the patient visited an emergency department or was hospitalized.

Claim Notes:

- Must be prescribed by a pediatric respirologist or allergist experienced in the treatment of severe asthma.
  - Combined use of dupilumab with other biologics used to treat asthma will not be reimbursed.
  - Approvals will be for a maximum of 200 mg every two weeks or 300 mg every four weeks.
  - Approval period: 1 year.
2. For the adjunctive treatment of severe asthma with a type 2 or eosinophilic phenotype or oral corticosteroid (OCS) dependent severe asthma in patients 12 years of age and older who are inadequately controlled with high-dose inhaled corticosteroids (ICS) and one or more additional asthma controller(s) (e.g., long-acting beta-agonist) and meets one of the following criteria:
    - blood eosinophil count  $\geq 0.15 \times 10^9/L$  within the past 12 months, or
    - have OCS dependent asthma.

Initial Discontinuation Criteria:

- Baseline asthma control questionnaire score has not improved at 12 months since initiation of treatment, or
- No decrease in the daily maintenance OCS dose in the first 12 months of treatment, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months.

Subsequent Discontinuation Criteria:

- Asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
- Reduction in the daily maintenance OCS dose achieved after the first 12 months of treatment is not maintained subsequently, or
- Number of clinically significant asthma exacerbations has increased within the previous 12 months.

Clinical Notes:

- A baseline and annual assessment of asthma symptom control using a validated asthma control questionnaire must be provided.
- A baseline and annual number of clinically significant asthma exacerbations must be provided.
- High-dose ICS is defined as greater than or equal to 500 mcg of fluticasone propionate or equivalent daily dose.
- A significant clinical exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least 3 days or the patient visited an emergency department or was hospitalized.

Claim Notes:

- Must be prescribed by a respirologist, clinical immunologist, allergist or internist experienced in the treatment of severe asthma.
- Combined use of dupilumab with other biologics used to treat asthma will not be reimbursed.
- Approvals will be for a maximum of 600 mg at week 0, then 300 mg every two weeks thereafter.
- Approval period: 1 year.

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Dupixent** – see Dupilumab

**Edaravone, solution for injection, 0.3mg/mL; oral solution, 105mg/5mL (Radicava-BMT)**

For the treatment of patients with probable or definite amyotrophic lateral sclerosis (ALS) who meet all of the following criteria:

Initiation criteria:

- Scores of at least two points on each item of the ALS Functional Rating Scale-Revised (ALSFRRS-R).
- Forced vital capacity is greater than or equal to 80% of predicted.
- ALS symptoms for two years or less.
- Not currently requiring permanent non-invasive or invasive ventilation.

Discontinuation Criteria:

- Patient becomes non-ambulatory (ALSFRRS-R score  $\leq$  1 for item 8) AND is unable to cut food and feed themselves without assistance, irrespective of whether a gastrostomy is in place (ALSFRRS-R score  $<$  1 for item 5a or 5b); or
- Patient requires permanent non-invasive or invasive ventilation.

Clinical Note :

- Patient must be under the care of a specialist with experience in the diagnosis and management of ALS.
- Approval period: 6 months

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Elexacaftor & Tezacaftor & Ivacaftor, tablet, 50mg & 25mg & 37.5mg (day) & Ivacaftor, tablet, 75mg (night); 100mg & 50mg & 75mg (day) & Ivacaftor, tablet, 150mg (night) (Trikafta-VER)**

For the treatment of cystic fibrosis (CF) in patients 6 years of age and older who meet all of the following criteria:

- Confirmed diagnoses of CF with at least one F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR gene); AND
- Patient has been optimized on best supportive care for their CF prior to starting Trikafta; AND
- Prescribed by a clinical specialist affiliated with a Canadian cystic fibrosis centre.

The following measurements must be completed prior to initiating treatment with Trikafta:

- Baseline spirometry measurements of FEV1 in liters and percent predicted (within the last 30 days); AND
- Number of days treated with oral and/or intravenous (IV) antibiotics for pulmonary exacerbations in the previous 6 months OR number of pulmonary exacerbations requiring oral and/or IV antibiotics in the previous 6 months; AND
- Number of CF-related hospitalizations in the previous 6 months; AND

- Weight, height, and body mass index (BMI); AND
- A score from an age-appropriate cystic fibrosis questionnaire as follows:
  - Cystic Fibrosis Questionnaire Child (CFQ-C) and Cystic Fibrosis Questionnaire-Parent (CFQ-P), if the Patient is 6 to 13 years of age, inclusive; or
  - Cystic Fibrosis Questionnaire Revised (CFQ-R teen/adult) Respiratory Domain score, if the Patient is 14 years of age or older.

**Exclusion Criteria:**

- Patient has undergone lung transplantation; OR
- Patient is using Trikafta as combination therapy with another cystic fibrosis transmembrane conductance regulator (CFTR) modulator.

Initial approval duration: 6 months

**Initial renewal criteria:**

Renewal of funding will be considered in patients demonstrating at least ONE of the following improvements after 6 months of treatment with Trikafta;

1. Improvement or percent predicted FEV1 by 5% or more above the baseline measurement; OR
2. A decrease in the total number of days for which the patient received treatment with oral and/or IV antibiotics for pulmonary exacerbations compared with the 6-month period prior to initiating treatment OR a decrease in the total number of pulmonary exacerbations requiring oral and/or IV antibiotics compared with the 6-month period prior to initiating treatment; OR
3. Decreased number of CF-related hospitalizations in the 6 months after initiation of Trikafta treatment compared with the 6-month period prior to initiating Trikafta; OR
4. No decline in BMI at 6 months compared with the baseline BMI assessment; OR
5. Improvement by 4 points or more in the CFQ-R Respiratory Domain scale compared to baseline scores.

**Subsequent renewal criteria:**

For patients who have met the initiation criteria and initial renewal criteria.

- Ongoing renewal of funding will be provided for those who are continuing to benefit from therapy with Trikafta and who do not meet any of the exclusion criteria.
- At the time of renewal application, please include the patient's most recent ppFEV1 and a clinical update to confirm the treatment benefits or response experienced by the patient.

Approval Duration of renewals: 1 year

Approved doses:

- 6 to < 12 years of age (weight < 30kg): 2 tablets (each containing elexacaftor/tezacaftor/ ivacaftor 50mg/ 25mg/ 37.5mg) taken in the morning & one tablet (ivacaftor 75mg) taken in the evening approximately 12 hours apart.

- 6 to < 12 years of age (weight ≥ 30kg): 2 tablets (each containing elexacaftor/ tezacaftor/ ivacaftor 100mg/ 50mg/ 75mg) taken in the morning & one tablet (ivacaftor 150mg) taken in the evening approximately 12 hours apart.
- 12 years of age and older: 2 tablets (each containing elexacaftor/ tezacaftor/ ivacaftor 100mg/ 50mg/ 75mg) taken in the morning & one tablet (ivacaftor 150mg) taken in the evening approximately 12 hours apart.

**Elexacaftor & Tezacaftor & Ivacaftor, tablet, 80mg & 40mg & 60mg (day) & Ivacaftor, tablet, 59.5mg (night); 100mg & 50mg & 75mg (day) & Ivacaftor, tablet, 75mg (night) (Trikafta-VER)**

For the treatment of cystic fibrosis (CF) in patients aged 2 to 5 years who have at least one F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene.

Initiation Criteria:

1. Confirmed diagnosis of CF with at least one F508del mutation in the CFTR gene
2. Aged 2 to 5 years
3. Prescribed by a specialist affiliated with a Canadian cystic fibrosis centre
4. The following measurements must be completed prior to initiating treatment:
  - Number of days treated with oral and IV antibiotics for pulmonary exacerbations in the previous 6 months OR number of pulmonary exacerbations requiring oral and/or IV antibiotics in the previous 6 months;
  - Weight, height, and BMI

Renewal Criteria:

1. For renewal after initial authorization, the physician must provide evidence of continuing benefit from treatment with ELZ-TEZ-IVA for subsequent renewal of reimbursement. Patients on therapy should be monitored for response (e.g., no decrease in BMI z-score) using clinical judgment and/or standard procedures.
2. Assessment for clinical response should occur every 12 months

Exclusion Criteria:

- Patient has undergone lung transplantation.
- Patient is using Trikafta as combination therapy with another cystic fibrosis transmembrane conductance regulator (CFTR) modulator.

**Elmiron** - see Pentosan Polysulfate Sodium

**Emend** - see Aprepitant

**Emerade** – see Epinephrine Bitartrate

**Emgality** – see Galcanezumab

**Empagliflozin, tablet, 10mg, 25mg (Jardiance-BOE)**

For the treatment of type 2 diabetes as a third drug added on to metformin and a sulfonylurea for patients with inadequate glycemic control on optimal doses of metformin and a sulfonylurea, and for whom insulin is not an option

**OR**

As an adjunct to diet, exercise, and standard care therapy to reduce the incidence of cardiovascular (CV) death in patients with type 2 diabetes mellitus and established cardiovascular disease, if the following criteria are met:

- Patients have inadequate glycemic control despite an adequate trial of metformin

Clinical Notes:

Established cardiovascular disease is defined as one of the following (details must be provided):

- History of myocardial infarction (MI).
- Multi-vessel coronary artery disease in two or more major coronary arteries (irrespective of revascularization status).
- Single-vessel coronary artery disease with significant stenosis and either a positive non-invasive stress test or discharged from hospital with a documented diagnosis of unstable angina within 12 months prior to selection.
- Last episode of unstable angina  $\geq 2$  months prior with confirmed evidence of coronary multi-vessel or single-vessel disease.
- History of ischemic or hemorrhagic stroke.
- Occlusive peripheral artery disease.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Empagliflozin & Metformin, tablet, 5mg & 500mg, 5mg & 850mg, 5mg & 1000mg, 12.5mg & 500mg, 12.5mg & 850mg, 12.5mg & 1000mg (Synjardy-BOE)**

For patients with type 2 diabetes mellitus who are already stabilized on therapy with metformin and empagliflozin, to replace the individual components of metformin and empagliflozin in these patients.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Enablex** – see Darifenacin

**Encorafenib, capsule, 75mg (Braftovi-PFI)**

**Metastatic Colorectal Cancer**



In combination with panitumumab for the treatment of patients with metastatic colorectal cancer who meet all of the following criteria:

- Presence of BRAF V600E mutation
- Disease progression following at least one prior therapy in the metastatic setting
- No previous treatment with an EGFR inhibitor

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Encorafenib will not be reimbursed in patients who have progressed on BRAF targeted therapy.

### **Metastatic Melanoma**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used in combination with binimetinib.

Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Encorafenib will not be reimbursed in patients who have progressed on BRAF targeted therapy.
- Requests will be considered for patients who received adjuvant BRAF targeted therapy if disease progression occurred at least 6 months following completion of therapy.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a**

patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

**Energair Breezhaler** – see Indacaterol & Glycopyrronium & Mometasone

**Enfuvirtide, injection kit, 90mg/mL (Fuzeon-HLR)**

For patients:

- a) Who have a CD4 count greater than 100 cells/mm<sup>3</sup>; **AND**
- b) Who have a viral load less than 100,000 copies/mL; **AND**
- c) Who have previously received less than 11 antiretroviral agents; **AND**
- d) Where therapy with Enfuvirtide is planned in combination with at least one other antiretroviral drug to which sensitivity has been demonstrated on resistance testing.

**Requests for Enfuvirtide (Fuzeon-HLR) must be made using the Enfuvirtide Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Enoxaparin** - see Low Molecular Weight Heparins

**Enspryng** – see Satralizumab

**Entecavir, tablet, 0.5mg (Baraclude – BMS and generics)**

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

**Entrectinib, capsule, 100mg, 200 mg (Rozlytrek-HLR)**

**ROS-1 Positive Non-Small Cell Lung Cancer or Metastatic Non-Small Cell Lung Cancer**

For the first-line treatment of patients with ROS-1 positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer (NSCLC).

Clinical Notes:

- Patients should have a good performance status.
- Treatment should continue until disease progression or unacceptable toxicity.

**Unresectable Locally Advanced or Metastatic Extracranial Solid Tumors with a NTRK Gene Fusion**

- For the treatment of adult patients with unresectable locally advanced or metastatic extracranial solid tumors with NTRK gene fusion without a known acquired resistance mutation. Eligible patients are not candidates for surgery and/or radiation due to risk of substantial morbidity and have no satisfactory treatment options.

Clinical Notes:

- Patients should have a good performance status.
- Treatment should be discontinued upon disease progression or unacceptable toxicity.
- CNS metastases are stable if present.
- Patients with prior progression on an NTRK inhibitor are not eligible.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Entresto** – see Sacubitril & Valsartan

**Entuzity** – see Insulin Regular

**Entyvio** - see Vedolizumab

**Enzalutamide, capsule, 40mg (Xtandi-AST)**

1. For treatment of patients with metastatic castration resistant prostate cancer who:
  - Are asymptomatic or mildly symptomatic after failure of androgen deprivation therapy with an ECOG performance status  $\leq 1$  and have not received prior chemotherapy and would be an alternative to abiraterone for patients and not sequential therapy in this asymptomatic or mildly symptomatic patient population **OR**
  - Have progressed on docetaxel-based chemotherapy with an ECOG performance status  $\leq 2$  and no risk factors for seizures and would be an alternative to abiraterone for patients and not sequential therapy in this symptomatic post docetaxel chemotherapy setting

Notes:

- Enzalutamide will not be reimbursed in combination with abiraterone.
- Use of enzalutamide in the past docetaxel setting is not permitted if previously used in the prechemotherapy setting

2. In combination with androgen deprivation therapy (ADT) for the treatment of patients with metastatic castration-sensitive prostate cancer (mCSPC). Patients must have had either no prior ADT or are within six months of beginning ADT in the metastatic setting.

Clinical Notes:

- Patients should have a good performance status and no risk factors for seizures.
- Treatment should continue until unacceptable toxicity or disease progression.

3. In combination with androgen deprivation therapy (ADT) for the treatment of patients with non-metastatic castration-resistant prostate cancer (nmCRPC) who are at high risk of developing metastases<sup>1</sup>.
- Patients should have a good performance status and no risk factors for seizures. Treatment should continue until unacceptable toxicity or radiographic disease progression.

Clinical Notes:

- Castration-resistance must be demonstrated during continuous ADT and is defined as 3 PSA rises at least one week apart, with the last PSA > 2 ng/mL.
- Castrate levels of testosterone must be maintained.
- Patients with N1 disease, pelvic lymph nodes < 2cm in short axis located below the common iliac vessels are eligible for enzalutamide.
- Enzalutamide will not be funded for patients who experience disease progression on apalutamide.
- Patients receiving enzalutamide for the treatment of non-metastatic CRPC will be eligible for funding of abiraterone at the time of disease progression to metastatic CRPC.

<sup>1</sup>High risk of developing metastases is defined as a prostate-specific antigen (PSA) doubling time of ≤ 10 months during continuous ADT

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Epinephrine, auto-injector, 0.15mg per dose, 0.3mg per dose (EpiPen-ALX)**

For the emergency treatment of anaphylactic reactions, when out of reach of immediate medical attention.

Note:

- Regular benefit, but with a quantity limit of two injections per 12 month period (one unit per dispense). Additional units require an exception status request.

**Epinephrine, pre-filled pen, 0.15mg per dose, 0.3mg per dose, 0.5mg per dose (Emerade-BAU)**

For the emergency treatment of anaphylactic reactions, when out of reach of immediate medical attention.

Note:

- Regular benefit, but with a total quantity limit of two injections per 12 month period. Additional units require an exception status request.

**EpiPen** - see Epinephrine

**EpiPen Jr.** - see Epinephrine

**Eplerenone, tablet, 25mg, 50mg (Generic)**

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 a complement to standard therapy.

Clinical Note:

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

**Epoetin Alfa, pre-filled syringe, 10,000IU/mL (Eprex-JAN)**

For the treatment of severe anemia related to chronic renal failure in patients with:

- a) Normocytic normochromic anemia, requiring transfusions in patients who have evidence of iron overload (Ferritin > 1000 ng/mL), **OR**
- b) Anemia requiring blood transfusions in patients having symptomatic angina and/or heart failure, **OR**
- c) Anemia requiring transfusion in patients with difficulties in blood grouping and febrile reactions due to antibodies, **OR**
- d) Anemia requiring transfusions in patients who have high levels of panel reactive anti HLA antibodies, **OR**
- e) Severe normocytic normochromic anemia (Hb < 100 g/L) whose only symptom is fatigue and have never received transfusions.

**The request for coverage must be made by or in consultation with a nephrologist, internal medicine specialist, or oncologist. A copy of the consultation must accompany the request.**

**The request for coverage must be made using the Erythropoietin Program Approval Form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Epoprostenol, vials, 0.5mg, 1.5mg (Caripul-ACT, Flolan-GSK)**

For the treatment of World Health Organization (WHO) class III or IV idiopathic pulmonary arterial hypertension in patients who do not demonstrate vasoreactivity on testing or who demonstrate vasoreactivity on testing but fail a trial of, or are intolerant to, calcium channel blockers.

For the treatment of WHO class III or IV pulmonary arterial hypertension associated with scleroderma in patients who do not respond adequately to conventional therapy.

Note: Coverage will be limited to medication and associated diluent costs only. No coverage will be provided for equipment or medical supplies (e.g. pumps, IV tubing, IV catheters, etc.).

Coverage must be requested and medications prescribed by a Pulmonary Arterial Hypertension (PAH) specialist.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Eprex** - see Epoetin Alfa

**Eptinezumab, vial, 100mg/1.0mL (Vyepiti-LUD)**

For the prevention of migraine in patients with a confirmed diagnosis of episodic or chronic migraine who have experienced an inadequate response, intolerance, or contraindication to at least two classes of oral prophylactic migraine medications.

Renewal Criteria:

- A reduction of at least 50% in the average number of migraine days per month at the time of initial renewal compared with baseline.
- At subsequent renewals, the patient continues to maintain the reduction of at least 50% in average number of migraine days per month.

Clinical Notes:

- The average number of headache and migraine days per month must be provided on initial and renewal requests.
- According to the International Headache Society criteria, episodic or chronic migraine are defined as:
  - Episodic migraine: migraine headaches on at least 4 days per month and less than 15 headache days per month for more than 3 months.
  - Chronic migraine: headaches for at least 15 days per month for more than 3 months of which at least eight days per month are with migraine.

Claim Notes:

- Initial approval period: 6 months
- Renewal approval period: 1 year

**Erelzi** – see Etanercept

**Erivedge** – see Vismodegib

**Erleada** – see Apalutamide

**Erlotinib, tablet, 25mg, 100mg, 150mg (Tarceva-HLR and generics)**

For use as monotherapy for the treatment of patients with locally advanced or metastatic NSCLC after failure of at least one prior chemotherapy regimen and whose EGFR expression status is positive or unknown.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage under the High Cost Drug Program must apply to this program using the application that is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Esbriet** – see Pirfenidone

**Eslicarbazepine Acetate, tablet, 200mg, 400mg, 600mg, 800mg (Aptiom-SNV)**

For the treatment of partial-onset seizures in patients with epilepsy who are currently receiving two or more antiepileptic drugs (AEDs) and for whom less costly AEDs are ineffective or not clinically appropriate.

**Etanercept, pre-filled syringe, 50mg/mL (Brenzys-MSD; Erelzi-SDZ; Rymti-LUP); pen injector, 25mg/0.5mL (Erelzi-SDZ); 50mg/mL (Brenzys-MSD; Erelzi-SDZ; Rymti-LUP)**

For etanercept naïve patients, approved requests will be for a biosimilar product

**Ankylosing Spondylitis**

For the treatment of patients with moderate to severe ankylosing spondylitis (Bath AS Disease Activity Index (BASDAI) score  $\geq 4$  on 10 point scale) who:

- a) have axial symptoms\* and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation or in whom NSAIDs are contraindicated OR
- b) have peripheral symptoms and who have failed to respond to, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.

\*Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease, do not require a trial of NSAIDs alone.

Claim Notes:

- Approvals will be for a maximum adult dose of 50 mg weekly or 25 mg twice weekly.
- Combined use of more than one biologic DMARD will not be reimbursed.

- Initial Approval: 6 months
- Renewal Approval: maximum of 12 months. Requests for renewal must include information showing the beneficial effects of the treatment, specifically:
  - a) a decrease of at least two points on the BASDAI scale, compared with pretreatment score OR
  - b) patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as Health Assessment Questionnaire (HAQ) or ability to return to work).

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Ankylosing Spondylitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Plaque Psoriasis**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score ; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.



- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 50 mg twice weekly for 12 weeks. If response criteria is met at 12 weeks, approval will be continued at a dose of 50 mg weekly.
- Initial approval: 12 weeks. Renewal approval: 1 year. Confirmation of continued response is required
- Concurrent use of biologics not approved.

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Polyarticular Juvenile Idiopathic Arthritis (pJIA)**

For the treatment of pJIA for patients aged 4-17 years with moderately or severe pJIA who have had an inadequate response to one or more disease-modifying anti-rheumatic drugs (DMARDs). Treatment must be initiated by a rheumatologist who is familiar with the use of DMARDs and/or biologic DMARDs in children.

Claim Notes:

- Approvals will be for a dose of 0.8 mg/kg weekly to a maximum of 50 mg weekly.
- Initial Approval: 6 months.
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and
  - Methotrexate (oral or parenteral) at a dose of  $\geq 20$ mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and

- Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 50 mg per week or 25 mg twice weekly.
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Rheumatoid Arthritis**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks  
AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.

- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Approvals will be for a maximum adult dose of 50 mg weekly or 25 mg twice weekly.
- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Everolimus, tablet, 2.5mg, 5mg, 10mg (Generics)**

**Metastatic Renal Cell Carcinoma (RCC)**

For the treatment of patients with advanced or metastatic renal cell carcinoma following disease progression on tyrosine kinase inhibitor therapy.

Clinical Notes:

- Patients must have a good performance status.
- Treatment should be discontinued upon disease progression or unacceptable toxicity.
- Requests for everolimus will not be considered for patients who experience disease progression on axitinib, cabozantinib or nivolumab monotherapy.

Claim Notes:

- Initial approval period: 6 months
- Renewal approval period: 1 year

### **Hormone Receptor Positive, HER2 Negative-Advanced Breast Cancer**

In combination with exemestane for postmenopausal patients (ECOG PS ≤2) with documented hormone receptor positive, HER2 negative-advanced breast cancer after recurrence or progression following a non-steroidal aromatase inhibitor (NSAI).

#### Clinical Notes:

- Patients must have a good performance status.
- Treatment should be discontinued upon disease progression or unacceptable toxicity.

#### Claim Notes:

- Requests for everolimus will not be considered for patients who experience disease progression on CDK4/6 inhibitor therapy.
- Approval period: 1 year

### **Metastatic Pancreatic Neuroendocrine Tumors (pNET)**

For the treatment of patients with progressive, unresectable, well or moderately differentiated, locally advanced or metastatic pancreatic neuroendocrine tumors (pNET) with good performance status (ECOG 0-2), until disease progression.

#### Claim Notes:

- Patients whose disease progresses on sunitinib are not eligible for funded treatment with everolimus for pNET
- Approval period: 1 year

### **Neuroendocrine Tumors of Gastrointestinal Or Lung Origin**

As a single agent treatment for patients with unresectable, locally advanced or metastatic; well-differentiated nonfunctional neuroendocrine tumors (NETs) of gastrointestinal or lung origin (GIL) in adults with documented radiological disease progression within six months and with a good performance status.

- Treatment should continue until confirmed disease progression or unacceptable toxicity.

#### Claim Notes:

- Approval period: 1 year.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a**

patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

### **Evolocumab, prefilled autoinjector, 140mg/mL (Repatha-AMG)**

For the treatment of heterozygous familial hypercholesterolemia (HeFH) in adult patients who require additional lowering of low-density lipoprotein cholesterol (LDL-C) if the following criteria are met:

- Definite or probable diagnosis of HeFH using the Simon Broome or Dutch Lipid Network criteria or genetic testing; and
- Patient is unable to reach LDL-C target (less than 2.0 mmol/L or at least a 50% reduction in LDL-C from untreated baseline) despite confirmed adherence to at least 3 months of continuous treatment with:
  - high-dose statin (e.g. atorvastatin 80 mg, rosuvastatin 40 mg) in combination with ezetimibe; or
  - ezetimibe alone, if high dose statin is not possible due to rhabdomyolysis, contraindication or intolerance

Initial renewal criteria:

- A reduction in LDL-C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L.

Subsequent renewal criteria:

- The patient continues to maintain a reduction in LDL-C of at least 40% from baseline or has reached a target LDL-C less than 2.0 mmol/L

Clinical Notes:

1. LDL-C levels must be provided.
2. Intolerance to high dose statin will be considered if patient has developed documented myopathy or abnormal biomarkers (i.e. creatinine kinase greater than 5 times the upper limit of normal) after trial of at least two statins and
  - for each statin, dose reduction was attempted rather than statin discontinuation, and intolerance was reversible upon statin discontinuation, but reoccurred with statin re-challenge where clinically appropriate; and
  - at least one statin was initiated at the lowest daily starting dose; and
  - other known causes of intolerance have been ruled out.
3. For patients who cannot take ezetimibe due to an intolerance or contraindication, details must be provided.

Claim Notes:

- Approvals will be for a maximum of 140mg every 2 weeks or 420mg monthly.
- Initial approval period: 6 months.
- Renewal approval period: 1 year

**Evrysdi** – see Risdiplam

**Exelon** - see Cholinesterase Inhibitors (ChEI)

**Eylea** – see Aflibercept

**Fabrazyme – vial, 5mg, 25mg (AVN)**

Coverage may be available for Fabrazyme for the treatment of Fabry Disease through the High Cost Drug Plan and Catastrophic Drug Plan, for eligible patients who meet the criteria set out in the Canadian Fabry Disease Treatment Guidelines.

The treatment guidelines are supported by the Canadian Fabry Disease Initiative (CFDI), and may be amended by the CFDI from time to time.

Please contact the PEI Pharmacare Program office at 1-877-577-3737 for more information regarding coverage availability and the Special Authorization application process for this product.

**Faricimab, vial, 6mg/0.05mL (Vabysmo-HLR)**

**Neovascular Age-Related Macular Degeneration**

Criteria For Initial Coverage:

For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD) where all of the following apply:

- a) Best Corrected Visual Acuity (BCVA) is between 6/12 and 6/96 **AND**
- b) The lesion size is less than or equal to 12 disc areas in greatest linear dimension **AND**
- c) There is evidence of recent (<3 months) presumed disease progression (blood vessel growth, as indicated by fluorescein angiography, optical coherence tomography (OCT), or recent visual acuity changes.

The interval between doses should not be shorter than one month. Administration is to be done by a qualified ophthalmologist experienced in intravitreal injections.

Criteria For Continued Coverage:

Treatment with faricimab should be continued only in people who maintain adequate response to therapy.

Faricimab should be discontinued if any of the following occur:

- a) Reduction in BCVA in the treated eye to less than 15 letters (absolute) on 2 consecutive visits in the treated eye, attributed to AMD in the absence of other pathology **OR**
- b) 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 **OR**

- c) There is evidence of deterioration of the lesion morphology despite optimum treatment over 3 consecutive visits.

Coverage will not be approved for patients:

- a) Receiving concurrent treatment with verteporfin.  
b) With permanent retinal damage as defined by the Royal College of Ophthalmology guidelines. Coverage is limited to a maximum of one vial per eye in any 30-day period. The request for coverage must be made by an ophthalmologist.

Approval Period: 1 year

### **Diabetic macular edema (DME)**

Criteria For Initial Coverage:

For the treatment of visual impairment due to diabetic macular edema (DME) in patients who meet all of the following criteria:

- clinically significant centre-involving macular edema for whom laser photocoagulation is also indicated
- hemoglobin A1c test in the past 6 months with a value of less than or equal to 11%
- best corrected visual acuity of 20/32 to 20/400
- central retinal thickness greater than or equal to 250 micrometers

Criteria For Continued Coverage:

- confirm that a hemoglobin A1c test in the past 6 months had a value of less than or equal to 11%
- date of last visit and results of best corrected visual acuity at that visit
- date of last OCT and central retinal thickness on that examination
- if faricimab is being administered monthly, please provide details on the rationale.

Clinical Notes:

- Treatment should be given monthly until maximum visual acuity is achieved (i.e. stable visual acuity for three consecutive months). Thereafter, visual acuity should be monitored monthly.
- Treatment should be resumed when monitoring indicates a loss of visual acuity due to DME and continued until stable visual acuity is reached again for three consecutive months.
- Treatment should be discontinued if there is no improvement of retinal thickness or visual acuity after three consecutive treatments.
- Injection will be by a qualified ophthalmologist with experience in administering intravitreal injections.
- Approval Period: 1 year

**Patients requesting coverage through the High Cost Drug Program must submit a**

patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

**Fasenra** – see Benralizumab

**Febratinib, capsule, 100mg (Inrebic-CEL)**

For the treatment of splenomegaly and/or disease-related symptoms in adult patients with intermediate-2 or high-risk primary myelofibrosis, post-polycythemia vera myelofibrosis, or post-essential thrombocythemia myelofibrosis, who have a contraindication or intolerance to ruxolitinib.

Clinical Notes:

1. Patients should have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Febuxostat, tablet, 80mg (Generics)**

For the treatment of symptomatic gout in patients who have documented hypersensitivity to allopurinol.

**Note: Intolerance or lack of response to allopurinol will not be covered by these criteria.**

**Fentanyl, transdermal patch, 12mcg/hr, 25mcg/hr, 37mcg/hr, 50mcg/hr, 75mcg/hr, 100mcg/hr (Generics)**

For the treatment of severe chronic pain that is not well controlled by short and long-acting Morphine and Hydromorphone products.

Maximum reimbursable coverage is for dosing every 72 hours.

**Fesoterodine Fumarate, extended release tablet, 4mg,8mg (Toviaz-PFI)**

For the treatment of over active bladder (not stress incontinence) in patients who cannot tolerate or have an insufficient response to an adequate trial (e.g. 3 months) of immediate release oxybutynin, solifenacin, or tolterodine.

**Fidaxomicin, tablet, 200mg (Dificid-MER)**

For the treatment of patients with Clostridium Difficile Infection (CDI) where the patient



has:

- a second or subsequent recurrence following treatment with oral vancomycin; OR
- treatment failure with oral vancomycin for the current CDI episode; OR
- an intolerance or contraindication to oral vancomycin.

Re-treatment criteria:

- Re-treatment with fidaxomicin will only be considered for an early relapse occurring within 8 weeks of the start of the most recent fidaxomicin course.

Clinical Notes:

- Treatment failure is defined as 14 days of vancomycin therapy without acceptable clinical improvement.
- Intolerance is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Should be prescribed by, or in consultation with, an infectious disease specialist/medical microbiologist (preferred) or an internist (if infectious disease or medical microbiology consult is not available).
- Requests will be approved for 200mg twice a day for 10 days.

**Filgrastim, prefilled syringe, 300mcg/0.5mL, 480mcg/0.8mL (Grastofil-APX, Nivestym-PFI, Nypozi-TAV), vial, 300mcg/mL, 480mcg/1.6mL (Nivestym-PFI)**

### **Chemotherapy Support**

For the prevention of febrile neutropenia in patients receiving myelosuppressive chemotherapy with curative intent who:

- are at high risk of febrile neutropenia due to chemotherapy regimen, co-morbidities or pre-existing severe neutropenia; or
- have had an episode of febrile neutropenia, neutropenic sepsis or profound neutropenia in a previous cycle of chemotherapy; or
- have had a dose reduction, or treatment delay greater than one week due to neutropenia.

Clinical Note:

- Patients with non-curative cancer receiving chemotherapy with palliative intent are not eligible for coverage of filgrastim for prevention of febrile neutropenia.

### **High Dose Chemotherapy with Stem Cell Support:**

For use in mobilizing stem cells in preparation for stem cell collection.

**Must be requested and prescribed by a specialist in hematology or medical oncology.**

Claim Notes:

- All requests for coverage of filgrastim will be approved for the biosimilar versions only.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Fingolimod** - See Multiple Sclerosis Agents

**Firazyr** – see Icatibant

**Firdapse** – see Amifampridine Phosphate

**Flolan** - see Epoprostenol

**Fludara** - see Fludarabine

**Fludarabine, tablet, 10mg (Fludara-BAY)**

For the treatment of chronic lymphocytic leukemia (CLL) in patients with an ECOG performance status of 0 to 2 when the patient has failed to respond to, or relapsed during/ after previous therapy with an alkylating agent and intravenous administration is not desirable.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Fluoxetine, oral solution, 20mg/5mL (Generics)**

For use in patients for whom oral capsules are not an option. Pediatric patients 12 and under will not require written Special Authorization.

**Fluticasone Furoate/Vilanterol, Inhaler, 100/25mcg/dose, 200/25mcg/dose (Breo Ellipta-GSK)**

- a. For the treatment of asthma in patients who are not well controlled on a regular and adequate course of inhaled steroid therapy prior to the request for combination therapy.

**NOTE**

Patients using this product must also have access to a short acting beta-2 agonist bronchodilator for the relief of acute symptoms.

- b. For the treatment of COPD, see Chronic Obstructive Pulmonary Disease.

**Foquest** – see Methylphenidate

**Foradil** - see Formoterol

**Formoterol Fumerate, powder for inhalation (capsule), 12mcg/dose (Foradil-NVR); powder for inhalation (inhaler), 6mcg/dose, 12mcg/dose (Oxeze Turbuhaler-AZE)**

- a) For the treatment of asthma when used in patients on concurrent steroid therapy.
- b) For the treatment of COPD, see Chronic Obstructive Pulmonary Disease.

Note: Patients using these products must also have access to a short-acting beta-2 agonist bronchodilator for the relief of acute symptoms.

**Formoterol & Budesonide, powder for inhalation, 6mcg & 100mcg per dose, 6mcg & 200mcg per dose (Symbicort Turbuhaler-AZE)**

- a) For the treatment of asthma in patients who are not well controlled on a regular and adequate course of inhaled steroid therapy prior to the request for combination therapy.
- b) For the treatment of COPD, see Chronic Obstructive Pulmonary Disease.

Note: Patients using this product must also have access to a short-acting beta-2 agonist bronchodilator for the relief of acute symptoms.

**Fosfomycin, sachet, 3g (Monurol-PAL & generic)**

For the treatment of uncomplicated urinary tract infections in adult female patients where:

- The infecting organism is resistant to other oral agents, or
- Other less costly treatments are not tolerated

Note: Regular benefit, but with a quantity limit of 3 doses per 12 month period (one sachet per dispense). Additional sachets require an exception status request.

**Fragmin** - see Low Molecular Weight Heparins

**Fremanezumab, prefilled syringe, autoinjector, 225mg/1.5 mL (Ajovy-TEV)**

For the prevention of migraine in patients with a confirmed diagnosis of episodic or chronic migraine who have experienced an inadequate response, intolerance, or contraindication to at least two classes of oral prophylactic migraine medications.

Renewal Criteria:

- A reduction of at least 50% in the average number of migraine days per month at the time of initial renewal compared with baseline.
- At subsequent renewals, the patient continues to maintain the reduction of at least 50% in

average number of migraine days per month.

Clinical Notes:

- The average number of headache and migraine days per month must be provided on initial and renewal requests.
- According to the International Headache Society criteria, episodic or chronic migraine are defined as:
  - Episodic migraine: migraine headaches on at least 4 days per month and less than 15 headache days per month for more than 3 months.
  - Chronic migraine: headaches for at least 15 days per month for more than 3 months of which at least eight days per month are with migraine.

Claim Notes:

- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**Fulphila** – see Pegfilgrastim

**Fulvestrant, syringe, 250mg/5mL (Generics)**

See palbociclib criteria  
See ribociclib criteria

**Fuzeon** - see Enfuvirtide

**Fycompa** – see Perampanel

**Galantamine** - see Cholinesterase Inhibitors (ChEI)

**Galcanezumab, 120mg/mL, prefilled pen, prefilled syringe (Emgality-LIL)**

For the prevention of migraine in patients with a confirmed diagnosis of episodic or chronic migraine who have experienced an inadequate response, intolerance, or contraindication to at least two classes of oral prophylactic migraine medications.

Renewal Criteria:

- A reduction of at least 50% in the average number of migraine days per month at the time of initial renewal compared with baseline.
- At subsequent renewals, the patient continues to maintain the reduction of at least 50% in average number of migraine days per month.

Clinical Notes:

- The average number of headache and migraine days per month must be provided on initial and renewal requests.
- According to the International Headache Society criteria, episodic or chronic migraine are defined as:

- Episodic migraine: migraine headaches on at least 4 days per month and less than 15 headache days per month for more than 3 months.
- Chronic migraine: headaches for at least 15 days per month for more than 3 months of which at least eight days per month are with migraine.

Claim Notes:

- Combined use with other calcitonin gene-related peptide (CGRP) antagonists will not be reimbursed.
- Initial approval: 6 months
- Renewal approval: 1 year

**Gatifloxacin, ophthalmic drops, 0.3% (Zymar-ALL and generic)**

For the treatment/prevention of bacterial conjunctivitis associated with eye surgery.

**Glatect** - see Multiple Sclerosis Agents

**Glatiramer Acetate** - see Multiple Sclerosis Agents

**Gleevec** - see Imatinib

**Gilenya** – see Multiple Sclerosis Agents

**Gilteritinib, tablet, 40 mg (Xospata-AST)**

As monotherapy for the treatment of adult patients with relapsed or refractory FMS-like tyrosine kinase 3 (FLT3)-mutated acute myeloid leukemia who meet all of the following criteria:

- Confirmed positive for FLT3 mutation at the time of relapse or determination of refractory disease
- Presence of FLT3-ITD, FLT3-TKD/D835 or FLT3-TKD/I836 mutation

Renewal Criteria:

- Written confirmation that the patient is responding to treatment.

Clinical Notes:

- Patients must have a good performance status.
- Treatment should continue as long as clinical benefit is observed or until unacceptable toxicity occurs.

Claim notes:

- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions using the DIN first, and then PDINs.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Giotrif** – see Afatinib

**Glucagen** – see Glucagon (Human Recombinant)

**Glucagon, nasal spray, 3mg (Baqsimi-LIL)**

Regular benefit up to two devices per 12 months. Additional units require patient meets criteria as below:

For the treatment of severe hypoglycemia (SH) reactions in patients with diabetes mellitus receiving insulin and at high risk for SH, when impaired consciousness precludes oral carbohydrate administration.

- Request should detail the clinical need for greater than 2 glucagon devices per 12 months, including the number of devices anticipated
- Request must be from a medical practitioner or nurse practitioner
- Special authorization requests for additional doses will be considered for up to one dose per month
- SA is valid for 12 months
- Coverage is limited to one unit at a time

**Glucagon (Human Recombinant), vial, 1mg; kit, 1mg (Glucagen - PAL)**  
**Note: IM administration only.**

Regular benefit up to two devices per 12 months. Additional units require patient meets criteria as below:

For the treatment of severe hypoglycemia (SH) reactions in patients with diabetes mellitus receiving insulin and at high risk for SH, when impaired consciousness precludes oral carbohydrate administration.

- Request should detail the clinical need for greater than 2 glucagon devices per 12 months, including the number of devices anticipated
- Request must be from a medical practitioner or nurse practitioner
- Special authorization requests for additional doses will be considered for up to one dose per month
- SA is valid for 12 months
- Coverage is limited to one unit at a time

**Glucagon** – see Glucagon (Recombinant DNA Origin)

**Glucagon (Recombinant DNA Origin), vial,1mg (Glucagon – LIL)**

Regular benefit up to two devices per 12 months. Additional units require patient meets criteria as below:

For the treatment of severe hypoglycemia (SH) reactions in patients with diabetes mellitus receiving insulin and at high risk for SH, when impaired consciousness precludes oral carbohydrate administration.

- Request should detail the clinical need for greater than 2 glucagon devices per 12 months, including the number of devices anticipated
- Request must be from a medical practitioner or nurse practitioner
- Special authorization requests for additional doses will be considered for up to one dose per month.
- SA is valid for 12 months
- Coverage is limited to one unit at a time

**Golimumab, Syringe, 50mg/0.5mL; auto-injector, 50mg/0.5mL (Simponi-JAN)**

**Ankylosing Spondylitis**

For the treatment of patients with moderate to severe ankylosing spondylitis (Bath AS Disease Activity Index (BASDAI) score  $\geq 4$  on 10 point scale) who:

- a) have axial symptoms\* and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation or in whom NSAIDs are contraindicated OR
- b) have peripheral symptoms and who have failed to respond to, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.

\*Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease, do not require a trial of NSAIDs alone.

**Claim Notes:**

- Approvals will be for a maximum adult dose of 50 mcg once monthly.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial Approval: 6 months
- Renewal Approval: maximum of 12 months. Requests for renewal must include information showing the beneficial effects of the treatment, specifically:
  - a) a decrease of at least two points on the BASDAI scale, compared with pretreatment score OR
  - b) patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as Health Assessment Questionnaire (HAQ) or ability to return to work).

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Ankylosing Spondylitis Special Authorization**

form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>

Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

### **Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and
  - Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and
  - Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

#### Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

#### Claim notes:

- Approvals will be for a maximum adult dose of 50 mcg monthly.
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>

Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

### **Rheumatoid Arthritis**



For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks  
AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Approvals will be for a maximum adult dose of 50 mcg once monthly.
- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Granisetron, tablet, 1 mg (Generic)**

For the treatment of emesis in cancer patients receiving highly emetogenic chemotherapy (i.e. containing Cisplatin); receiving moderately emetogenic chemotherapy (i.e. containing Cyclophosphamide, Doxorubicin, Epirubicin, or Melphalan); or receiving radiation therapy

and who have:

- a) Experienced adverse effects to Metoclopramide, Prochlorperazine, or Dexamethasone or have a specific contraindication which does not allow use of these drugs as antiemetics or,
- b) Continued episodes of nausea and vomiting related to chemotherapy which have not responded to therapeutic doses of Metoclopramide, Prochlorperazine, or Dexamethasone.

**A maximum of 10 tablets per cycle of chemotherapy will be approved.**

**Grastofil** – see Filgrastim

**Guselkumab, autoinjector, prefilled syringe, 100mg/mL (Tremfya-JAN)**

### **Plaque Psoriasis**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet, or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Concurrent use of biologics not approved.

- Approvals will be for 100mg by subcutaneous injection at weeks 0,4 followed by maintenance dosing of 100mg every 8 weeks.
- Initial approval: 16 weeks.
- Renewal approval: 1 year

**The request must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and
  - Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and
  - Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

#### Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

#### Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial approval: 12 weeks, loading dose of 100mg at weeks 0, 4, and 8 weeks
- Maximum dose of 100mg every 8 weeks
- Renewal approval: 1 year. Confirmation of continued response required.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Hadlima** - see Adalimumab

**Hemangirol** – see Propranolol

**Hulio** - see Adalimumab

**Hydromorph Contin** - see Hydromorphone, controlled-release capsule

**Hydromorphone HCl, controlled-release capsule, 3mg, 4.5mg, 6mg, 9mg, 12mg, 18mg, 24mg, 30mg (Hydromorph Contin-PFR)**

For the treatment of patients with documented severe chronic pain that is not well controlled by short and long-acting Morphine and short-acting Hydromorphone products. Maximum reimbursable coverage is for twice daily dosing.

**Hydromorphone HCl, injection solution, 10mg/mL, 20mg/mL, 50mg/mL (Generic)**

Note: For Nursing Home Program, no Special Authorization is required.

For the treatment of severe chronic pain that is not well controlled by short and long-acting oral Morphine and Hydromorphone products:

For other patients upon written request or recommendation from a palliative care or pain clinic. A copy of the recommendation must accompany the Special Authorization.

**Hydromorphone HP** - see Hydromorphone, injection solution

**Hyrimoz** - see Adalimumab

**Ibrance** – see Palbociclib

**Ibrutinib, capsule, 140mg (Imbruvica-JAN)**

1. For the treatment of patients with previously untreated chronic lymphocytic leukemia (CLL) / small lymphocytic lymphoma (SLL) for whom fludarabine-based treatment is considered inappropriate due to high risk of relapse or refractory disease (includes 17p deletion, TP3 mutation, 11q deletion and unmutated IGHV) based on prognostic biomarkers.
2. For the treatment of patients with CLL/SLL who have received at least one prior therapy and are considered inappropriate for treatment of retreatment with a fludarabine-based

regimen.

3. For the treatment of patients with relapsed or refractory mantle cell lymphoma.

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

Claim Notes:

- Ibrutinib will not be reimbursed when used in combination with rituximab.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage under the High Cost Drug Program must apply to this program using the application that is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

#### **Icatibant, syringe, 30mg/3mL (Firazyr-SHR)**

For the treatment of acute attacks of hereditary angioedema (HAE) in adults with lab confirmed c1-esterase inhibitor deficiency (type I or type II) if the following conditions are met:

- Treatment of non-laryngeal attacks of at least moderate severity, or
- Treatment of acute laryngeal attacks

Limited to a single dose for self-administration per attack AND prescribed by physicians with experience in the treatment of HAE

**The Special Authorization form is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Iclusig** – see Ponatinib

**Idacio** - see Adalimumab

#### **Idelalisib, tablet, 100mg, 150mg (Zydelig-GIL)**

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

The Special Authorization form is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .

Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .

**Ilaris** – see Canakinumab

**Ilumya** – see Tildrakizumab

**Imatinib, tablet, 100mg, 400mg (Gleevec-NVR and generics)**

- a) For the treatment of patients who have documented evidence of Philadelphia chromosome positive (Ph+) chronic myeloid leukemia (CML), with an ECOG performance status of 0 - 2\*.
- b) For the treatment of adult patients with newly diagnosed Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ALL) when used as a single agent for induction and maintenance phase therapy.
- c) For the treatment of patients with C-Kit positive (CD117), metastatic or locally advanced, inoperable gastrointestinal stromal tumours (GIST) and who have an ECOG performance status of 0 - 2\*.
- d) For the adjuvant treatment of adult patients who are at intermediate to high risk of relapse following complete resection of Kit (CD117) positive GIST.

Must be prescribed by a hematologist or oncologist.

- Patients who are asymptomatic and those who are symptomatic and in bed less than 50% of the time.

Initial approval: 12 months

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

Patients requesting coverage under the High Cost Drug Program must apply to this program using the application that is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .

**Imbruvica** – see Ibrutinib

**Imitrex** - see Sumatriptan

**Increlex** – see Mecasermin

**Incruse Ellipta** – see Umeclidinium Bromide

**Indacaterol & Glycopyrronium & Mometasone, inhalation capsule, 150mcg & 50mcg & 160mcg (Enerzair Breezhaler-NVR)**

For the maintenance treatment of asthma in adult patients not adequately controlled with a maintenance combination of long-acting-beta<sub>2</sub>-agonist and a medium or high dose of an inhaled corticosteroid who experienced one or more asthma exacerbations in the previous 12 months.

**Indacaterol & Mometasone, inhalation capsule, 150mcg & 80mcg, 150mcg & 160mcg, 150mcg & 320mcg (Ateectura Breezhaler-NVR)**

For the treatment of asthma in patients who are not well controlled on a regular and adequate course of inhaled steroid therapy prior to the request for combination therapy.

**Infliximab, injection powder, 100mg/vial (Avsola-AGA; Inflectra-HOS; Renflexis-MSD)**

**Ankylosing Spondylitis**

For the treatment of patients with moderate to severe ankylosing spondylitis (Bath AS Disease Activity Index (BASDAI) score  $\geq 4$  on 10 point scale) who:

- a) have axial symptoms\* and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation or in whom NSAIDs are contraindicated OR
- b) have peripheral symptoms and who have failed to respond to, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.

\*Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to axial disease, do not require a trial of NSAIDs alone.

Claim Notes:

- Approvals will be for a maximum adult dose of 5mg/kg at 0,2 and 6 weeks then every 6 to 8 weeks.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial Approval: 6 months
- Renewal Approval: maximum of 12 months.

Requests for renewal must include information showing the beneficial effects of the treatment, specifically:

- a) a decrease of at least two points on the BASDAI scale, compared with pretreatment score OR

- b) patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as Health Assessment Questionnaire (HAQ) or ability to return to work).

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Ankylosing Spondylitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Crohn's Disease**

- 1) For the treatment of patients with moderate to severe Crohn's disease who have active disease and are refractory, intolerant or have contraindications to:
- Prednisone 40mg (or equivalent) daily for  $\geq 2$  weeks, AND
  - Azathioprine  $\geq 2$  mg/kg/day for  $\geq 3$  months, OR
  - Mercaptopurine  $\geq 1$  mg/kg/day for  $\geq 3$  months, OR
  - Methotrexate (SC or IM)  $\geq 15$  mg/week for  $\geq 3$  months

#### Clinical Notes:

- Refractory is defined as lack or loss of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.
- Consideration will be given for the approval of a biologic DMARD (disease modifying antirheumatic drug) without a trial of a traditional DMARD for patients who have an aggressive/severe disease course (e.g. extensive disease, a modified Harvey Bradshaw Index score  $> 16$ ) and are refractory, intolerant or have contraindications to systemic corticosteroids.

#### Claim notes:

- Initial approval is for 3 doses of 5mg/kg/dose administered at 0, 2, and 6 weeks. Renewal of coverage will require reassessment of the patient and submission of a new Crohn's Disease Special Authorization form. Continued coverage will be approved at a dose not exceeding 5mg/kg every 8 weeks.
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a gastroenterologist using the Crohn's Disease Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The**



**patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

- 2) For the treatment of fistulizing Crohn's Disease in patients who:
1. Have a Harvey Bradshaw Index score of 7 or more, AND
  2. Have an actively draining perianal or enterocutaneous fistula(e) that have recurred or persisted despite a course of appropriate antibiotic therapy (e.g. Ciprofloxacin with or without Metronidazole for a minimum of 3 weeks), AND
  3. Have not responded to or are intolerant to immunosuppressive therapy (Azathioprine, Mercaptopurine or Methotrexate) or where such therapy is contraindicated.

Claim notes:

- Initial approval is for 3 doses of 5mg/kg/dose administered at 0, 2, and 6 weeks. Renewal of coverage will require reassessment of the patient and submission of a new Crohn's Disease Special Authorization form. Continued coverage will be approved at a dose not exceeding 5mg/kg every 8 weeks.
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a gastroenterologist using the Crohn's Disease Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Plaque Psoriasis**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20$ mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score ; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 5 mg/kg at 0, 2, and 6 weeks then every 8 weeks for 12 weeks. If response criteria is met at 12 weeks, approval will be continued at a dose of 5 mg/kg every 8 weeks.
- Initial approval: 12 weeks.
- Renewal approval: 1 year.
- Concurrent use of biologics not approved.

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and
  - Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and
  - Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.

- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 5 mg/kg at 0, 2, and 6 weeks then every 8 weeks thereafter.
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Rheumatoid Arthritis**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks  
AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Approvals will be for a maximum adult dose of 3mg/kg/dose at 0, 2, and 6 weeks then every 8 weeks thereafter.
- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ulcerative Colitis**

For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score > 4, and a rectal bleeding subscore ≥ 2 and are:

- Refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks AND prednisone ≥ 40mg daily for two weeks or IV equivalent for one week) OR
- Corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Initial approval is for 3 doses of 5 mg/kg/dose administered at 0, 2, and 6 weeks.

Renewal of coverage will require reassessment of the patient and submission of a new Ulcerative Colitis Special Authorization form. Requests must include information demonstrating the beneficial effects of the treatment, specifically:

- a decrease in the partial Mayo score ≥ 2 from baseline, and
- a decrease in the rectal bleeding subscore ≥1.

- Renewal Approval: 1 year. The maximum approved dose is 5 mg/kg every 8 weeks.
- Combined use of more than one biologic DMARD will not be reimbursed.

Clinical Notes:

- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.

- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.
- Patients with severe disease (partial Mayo > 6) do not require a trial of 5-ASA

**The request for coverage must be made by a gastroenterologist using the Ulcerative Colitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Inflectra** – see Infliximab

**Inlyta** - see Axitinib

**Innohep** – see Low Molecular Weight Heparins

**Inqovi** – see Decitabine and Cedazuridine

**Inrebic** – see Fedratinib

**Inspiroto Respimat** – see Tiotropium/Olodaterol

**Insulin aspart, vial, 100unit/mL (Novorapid-NNO)**

For patients who are unable to switch to a biosimilar brand of insulin aspart due to insulin pump compatibility, the patient or healthcare provider can complete the online biosimilar switching exemption form for continued coverage of Novorapid.

[Application for Biosimilar Switching Exemptions](#)

**Insulin Detemir, cartridge, prefilled pen; 100 unit/mL (Levemir-NNO)**

- For the treatment of pediatric and adolescent patients with type 1 diabetes requiring insulin. Requests for pediatric and adolescent patients will be approved with an automatic Special Authorization tool within the electronic claims system.
- For the treatment of pregnant individuals with diabetes requiring insulin therapy. Requests for pregnant patients will require a written Special Authorization.

**Insulin Glargine, prefilled pen, 300 unit/mL (Toujeo Solostar & Toujeo Doublestar-AVN)**

For the treatment of patients who have been diagnosed with type 1 or type 2 diabetes requiring insulin and have previously used **all eligible open benefit** long-acting insulin analogues at optimal dosing

AND have experienced unexplained hypoglycemia at least once a month despite optimal management

**OR**

For the treatment of patients who have been diagnosed with type 1 or type 2 diabetes requiring high dose insulin.

**The request for coverage must be made using the Long Acting Insulin Analogues Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Insulin Regular, prefilled pen, 500 unit/mL (Entuzity-LIL)**

For the treatment of diabetes mellitus in patients with unacceptable glycemic control who require more than 200 units of insulin per day, with or without other therapies.

- Treatment should be initiated by a specialist with experience in treating severe insulin resistance.

**Interferon Beta-1A** – see Multiple Sclerosis Agents

**Interferon Beta-1B** – see Multiple Sclerosis Agents

**Invega Sustenna** – see Paliperidone

**Invega Trinza** – see Paliperidone

**Invokana** – see Canagliflozin

**Isavuconazole, capsule, 100mg (Cresemba-AVI)**

- For the treatment of adult patients with invasive aspergillosis who have a contraindication, intolerance or have failed to respond to oral voriconazole and caspofungin.
- For the treatment of adult patients with invasive mucormycosis.

Claim Notes:

- Must be prescribed by a hematologist or specialist in infectious diseases or medical microbiology.
- Initial requests will be approved for a maximum of 3 months.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Itraconazole, oral solution, 10mg/mL (Generic)**

For the treatment of immunocompromised adult patients with oral and/or esophageal

candidiasis

Clinical Note:

- As per the drug monograph, Itraconazole capsules and oral solution should not be used interchangeably due to differences in bioavailability.

**Ivabradine, tablet, 5mg, 7.5mg (Lancora-SER)**

For the treatment of adult patients with New York Heart Association (NYHA) class II or III stable heart failure when administered in combination with standard chronic heart failure therapies to reduce the incidence of cardiovascular death and hospitalization, who meet all of the following criteria:

- Left ventricular ejection fraction (LVEF) of  $\leq 35\%$
- Sinus rhythm with a resting heart rate  $\geq 77$  beats per minute (bpm)
- At least one hospitalization due to heart failure in the past year
- NYHA class II or III symptoms despite at least four weeks of treatment with the following:
  - a stable dose of an angiotensin converting enzyme inhibitor (ACEI) or an angiotensin II receptor blocker (ARB)
  - a stable dose of a beta blocker
  - an aldosterone antagonist

Clinical Notes:

1. Resting heart rate must be documented as  $\geq 77$  bpm on average using either an ECG on at least three separate visits or by continuous monitoring.
2. For patients who have not received four weeks of therapy with an ACEI/ARB, beta blocker and aldosterone antagonist due to an intolerance or contraindication, details must be provided.
3. Initiation and up-titration should be under the supervision of a physician experienced in the treatment of heart failure.

**Ivacaftor, tablet, 150mg (Kalydeco-VTX)**

For the treatment of cystic fibrosis in patients who meet the following criteria:

- the patient is at least 6 years old and has one of the following mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene: G551D, G1244E, G1349D, G178R, G551S, S1251N, S1255P, S549N, OR S549R; OR
- the patient is at least 18 years old with an R117H mutation in the CFTR gene.

Renewal requests will be considered in patients with documented response to treatment as evidenced by the following:

- a) In cases where the patient's sweat chloride levels prior to commencing therapy were above 60mmol/L:
  - the patient's sweat chloride level fell below 60mmol/L; or
  - the patient's sweat chloride level is 30% lower than the level reported in a

- previous test;
- b) In cases where the patient's sweat chloride levels prior to commencing therapy were below 60mmol/L:
- the patient's sweat chloride level is 30% lower than the level reported in a previous test; or
  - the patient demonstrates a sustained absolute improvement in FEV<sub>1</sub> of at least 5% when compared to the FEV<sub>1</sub> test conducted prior to the commencement of therapy. FEV<sub>1</sub> will be compared with the baseline pre-treatment level one month and three months after starting treatment.

**Clinical Notes:**

- The patient's sweat chloride level and FEV<sub>1</sub> must be provided with each request.
- A sweat chloride test must be performed within a few months of starting ivacaftor therapy to determine if sweat chloride levels are reducing.
  - If the expected reduction occurs, a sweat chloride test must be performed again 6 months after starting therapy to determine if the full reduction has been achieved. Thereafter, sweat chloride levels must be checked annually.
  - If the expected reduction does not occur, a sweat chloride test should be performed again one week later. If the criteria are not met, coverage will be discontinued.

**Claim Notes:**

- Approved dose: 150mg every 12 hours
- Approval period: 1 year

<sup>1</sup>Please note:baseline sweat chloride levels and FEV1 are not required to meet initial approval criteria for Kalydeco,but these parameters are used to evaluate the effect of Kalydeco at the time of renewal. To avoid delays, the prescriber should submit a copy of the mutation report, recent baseline sweat chloride levels before starting Kalydeco, and recent baseline FEV1 with the initial request for funding of Kalydeco. These baseline values will be used to evaluate the patient's response to therapy at the time of renewal and would be logistically difficult to obtain once treatment is initiated.

**Ivacaftor/lumacaftor – see Orkambi**

**Ixekizumab, autoinjector, syringe, 80mg/mL (Taltz-LIL)**

**Plaque Psoriasis**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of ≥ 20mg weekly (≥15mg if patient is ≥65 years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND



- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score ; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 160 mg at week 0, followed by 80mg at week 2, 4, 6, 8, 10 and 12. If response criteria is met at 12 weeks, approval will be continued at a dose of 80 mg every 4 weeks up to one year.
- Initial approval: 12 weeks.
- Renewal approval: 1 year.
- Concurrent use of biologics not approved.

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and

- Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and
- Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 160 mg by subcutaneous injection (two 80 mg injections) at Week 0, followed by 80 mg every 4 weeks.
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Jakavi** – see Ruxolitinib

**Jamteki** – see Ustekinumab

**Januvia** – see Sitagliptin

**Janumet** – see Sitagliptin & Metformin Hydrochloride

**Janumet XR** – see Sitagliptin & Metformin Hydrochloride

**Jardiance** – see Empagliflozin

**Jentadueto** – see Linagliptin & Metformin Hydrochloride

**Zejula** – see Niraparib

**Kalydeco** – see Ivacaftor

**Kesimpta** – see Ofatumumab

**Kevzara** – see Sarilumab

**Kisqali** – see Ribociclib

**Komboglyze** – see Saxagliptin & Metformin Hydrochloride

**Lactulose 667mg/mL syrup**

For the treatment of hepatic encephalopathy.

**Lanadelumab, vial, prefilled syringe, 300mg/2mL (Takhzyro-TAK)**

For the prevention of attacks of type I or II hereditary angioedema (HAE) in patients 12 years of age and older who have experienced at least three HAE attacks within any four-week period and required the use of an acute injectable treatment.

Discontinuation Criteria:

- No reduction in the number of HAE attacks for which acute injectable treatment was received during the first three months of treatment with lanadelumab compared to the number of attacks observed before initiating treatment with lanadelumab; or
- Increase in the number of HAE attacks for which acute injectable treatment was received compared to the number of attacks before initiating treatment with lanadelumab.

Clinical Note:

- The pre-treatment attack rate must be provided for those patients who are already receiving long-term prophylactic treatment for HAE and intend to transition to lanadelumab.

Claim Notes:

- The patient must be under the care of a physician experienced in the diagnosis and treatment of HAE
- Not to be used in combination with other long-term prophylactic treatment of HAE (e.g., C1 esterase inhibitor).
- Approvals will be for a maximum of 300 mg every two weeks.
- Initial approval period: 3 months.
- Renewal approval period: 6 months.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions using the DIN first, and then PDINs.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Lancora** – see Ivabradine

**Lansoprazole** – see Proton Pump Inhibitors

**Lansoprazole & Clarithromycin & Amoxicillin, 7-day package, 30mg & 500mg & 500mg (Generics)**

One week of therapy will be considered for individuals with documented duodenal or gastric ulcers and a recent documented positive helicobacter pylori test.

**Lapelga** – see Pegfilgrastim

**Larotrectinib, capsule, 25mg, 100mg; oral liquid, 20mg/mL (Vitrakvi-BAY)**

As monotherapy for the treatment of adult and pediatric patients with unresectable locally advanced or metastatic solid tumors who meet all of the following criteria:

- Tumors have a NTRK gene fusion without a known acquired resistance mutation
- No other satisfactory treatment options
- Not a candidate for surgery and/or radiation due to risk of substantial morbidity

Clinical Notes:

- Patients must have a good performance status.
- If brain metastases are present, patients must be asymptomatic.
- Treatment should be discontinued upon radiographic disease progression or unacceptable toxicity.
- Patients with prior disease progression on a NTRK inhibitor are not eligible.

Claim Notes:

- Approval period: 6 months

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Lemtrada** – see Multiple Sclerosis Agents

**Lenalidomide, capsule, 2.5mg, 5mg, 10mg, 15mg, 20mg, 25mg (Revlimid-CEL and generics)**

## **Multiple Myeloma**

For the treatment of newly diagnosed Multiple Myeloma, in combination with daratumumab and dexamethasone, for patients who are not suitable for autologous stem cell transplant and have a good performance status.

For the treatment of Multiple Myeloma when used in combination with dexamethasone, in patients who:

- Are not candidates for autologous stem cell transplant;  
AND
- Where the patient is either:
  - Refractory to or has relapsed after the conclusion of initial or subsequent treatments and who is suitable for further chemotherapy;
  - OR
  - Has completed at least one full treatment regimen therapy and is experiencing intolerance to their current chemotherapy.

For the Maintenance Treatment of patients with newly diagnosed Multiple Myeloma, following autologous stem-cell transplantation (ASCT), in patients who are with stable disease or better, with no evidence of disease progression.

## **Myelodysplastic Syndrome**

For the treatment of Myelodysplastic Syndrome (MDS) in patients with:

- Demonstrated diagnosis of MDS on bone marrow aspiration
- Presence of 5-Q31 deletion documented by appropriate genetic testing
- International Prognostic Scoring System (IPSS) risk category low or intermediate (Calculator available on [www.uptodate.com](http://www.uptodate.com) )
- Presence of symptomatic anemia (defined as transfusion dependent)
  - Initial approval period – 6 months  
Renewal criteria:
    - For patients who were transfusion-dependent and have demonstrated a reduction in transfusion requirements of at least 50%.
    - Renewal period – 1 year

**Pharmacare will reimburse a single capsule per day per person.  
Multiple strengths or multiple capsules per day will not be reimbursed.**

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

## **Lenvatinib, capsule, 4mg/dose, 8mg/dose, 12mg/dose (Lenvima-EIS)**

### **Advanced Endometrial Carcinoma**

Lenvatinib combined with pembrolizumab for the treatment of adult patients with advanced endometrial carcinoma that is not microsatellite instability high (MSI-H) or mismatch repair deficient (dMMR), who have disease progression following prior platinum-based systemic therapy and are not candidates for curative surgery or radiation.

### **Advanced and Metastatic Renal Cell Carcinoma**

Lenvatinib combined with pembrolizumab for the treatment of adult patients with advanced (not amenable to curative surgery or radiation) or metastatic renal cell carcinoma (RCC) who have had no prior systemic therapy for metastatic disease.

### **Differentiated Thyroid Cancer (Lenvima 10mg,14mg, 20mg and 24mg Compliance Pack)**

For the treatment of patients with locally recurrent or metastatic, progressive, differentiated thyroid cancer (DTC) who meet the following criteria:

- Pathologically confirmed papillary or follicular thyroid cancer, and
- Disease that is refractory or resistant to radioactive iodine therapy, and
- Radiological evidence of disease progression within the previous 13 months, and
- Previous treatment with no more than one tyrosine kinase inhibitor (TKI).

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

### **Hepatocellular Carcinoma (Lenvima 4mg, 8mg and 12mg Compliance Pack)**

For the first-line treatment of adult patients with unresectable metastatic hepatocellular carcinoma who meet all the following criteria:

1. Child-Pugh class status of A.
2. ECOG performance status of 0 or 1.
3. Less than 50% liver involvement and no invasion of the bile duct or main portal vein.
4. No brain metastases or prior liver transplantation.

Clinical Notes:

- Treatment should be continued until disease progression or unacceptable toxicity. Patients who are unable to tolerate lenvatinib may be switched to sorafenib if there is no disease progression and provided all other funding criteria are met.
- Patients with disease progression on lenvatinib are not eligible for reimbursement

of sorafenib.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Lenvima** – see Lenvatinib

**Letemovir, tablet, 240mg, 480mg (Prevymis-MER)**

For the prevention of cytomegalovirus (CMV) infection in adult CMV-seropositive recipients [R+] of an allogeneic hematopoietic stem cell transplant (HSCT) who have undetectable CMV viremia at baseline and meet one of the following criteria:

- umbilical cord blood as a stem cell source
- recipient of a haploidentical transplant
- recipient of T-cell depleted transplant
- treated with antithymocyte globulin (ATG) for conditioning
- requiring high-dose steroids or other immunosuppression for acute graft versus host disease (GVHD)
- treated with ATG for steroid-refractory acute GVHD
- documented history of CMV disease prior to transplantation

Clinical Note:

- High-dose steroids is defined as the use of greater than or equal to 1 mg/kg/day of prednisone or equivalent dose of another corticosteroid.

Claim Notes:

- Must be prescribed by a medical oncologist, hematologist, or infectious disease specialist or other physician with experience in the management of HSCT.
- Approvals will be for a maximum dose of 480mg per day.
- Approval period: 100 days per HSCT

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Levemir** – see Insulin Detemir

**Levetiracetam, oral solution, 100mg/mL (Generic)**

- For patients who require administration through a feeding tube.
- For patients 19 years of age and younger, who cannot use a tablet or capsule.
- Pediatric patients 12 and under will not require written Special Authorization.

**Levocarnitine, tablet, oral solution, 330mg, 100mg/mL (Carnitor-SIG and generic)**

- For the treatment of patients with primary systemic carnitine deficiency.
- For the treatment of patients with an inborn error of metabolism that results in secondary carnitine deficiency.

**Levodopa & Carbidopa, intestinal gel cassette, 20mg/mL & 5mg (Duodopa-ABV)**

For the treatment of patients with advanced levodopa-responsive Parkinson's Disease (PD) who meet all of the following criteria:

- Experiences severe disability with at least 25% of the waking day in the off state and/or ongoing levodopa-induced dyskinesias, despite having tried frequent dosing of levodopa (at least five doses per day).
- Received an adequate trial of maximally tolerated doses of levodopa, with demonstrated clinical response.
- Failed an adequate trial of the following adjunctive medications, if not contraindicated and/or contrary to the clinical judgement of prescriber: entacapone, a dopamine agonist, a monoamine oxidase-B (MAO-B) inhibitor and amantadine.
- Must be able to administer the medication and care for the administration port and infusion pump. Alternatively, trained personnel or care partner must be available to perform these tasks reliably.

Exclusion Criteria:

- Patients with a contraindication to the insertion of a PEG-J tube.
- Patients with severe psychosis or dementia.

Renewal Criteria:

- Patients continue to demonstrate a significant reduction in the time spent in the off state and/or ongoing levodopa-induced dyskinesias, along with an improvement in the related disability.

Clinical Note:

- Time in the off state, frequency of motor fluctuations, and severity of associated disability should be assessed by a movement disorder subspecialist and be based on an adequate and reliable account from longitudinal speciality care, clinical interview of a patient and/or care partner, or motor symptom diary.

Claim Notes:

- Must be prescribed by a movement disorder subspecialist who has appropriate training in the use of Duodopa and is practicing in a movement disorder clinic that provides ongoing management and support for patients receiving treatment with Duodopa.
- Approval period: 1 year.



**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Levofloxacin, tablet, 250mg, 500mg, 750mg (Generics)**

Note: For Cystic Fibrosis and Nursing Home Programs, no Special Authorization is required.

- a) For the treatment of infections in persons allergic to alternative agents. Up to 10 days of therapy will be considered.
- b) For the treatment of infections in patients with asthma or COPD not responding to first-line antibiotics. Up to 10 days of therapy will be considered.
- c) For the treatment of infections caused by organisms known to be resistant to alternative antibiotics. Up to 10 days of therapy will be considered.
- d) For the completion of treatment started in the hospital inpatient setting. Up to 7 days of therapy will be considered.

**Linagliptin, tablet, 5mg (Trajenta-BOE)**

For the treatment of type 2 diabetes as a third drug added on to metformin and a sulfonylurea for patients with inadequate glycemic control on optimal doses of metformin and a sulfonylurea, **and** for whom insulin is not an option.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Linagliptin & Metformin Hydrochloride, tablet, 2.5mg/500mg, 2.5mg/850mg, 2.5mg/1000mg (Jentadueto-BOE)**

For patients with type 2 diabetes for whom insulin is not an option and who are already stabilized on therapy with metformin, a sulfonylurea and linagliptin, to replace the individual components of linagliptin and metformin in these patients.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Linezolid, tablet, 600mg (Generics)**

- (a) For the treatment of proven VRE (Vancomycin-Resistant Enterococcus) infections. Must be prescribed in consultation with a specialist in infectious diseases. A copy of a C&S report demonstrating Vancomycin resistance must accompany the request. Up to 28 days of therapy will be considered.
- (b) For the treatment of proven MRSA (Methicillin-Resistant Staph. Aureus) and MRSE

(Methicillin-Resistant Staph. Epidermidis) infections in patients who are unresponsive or intolerant to Vancomycin. Must be prescribed in consultation with a specialist in infectious diseases. A copy of a C&S report demonstrating Vancomycin resistance must accompany the request. Up to 28 days of therapy will be considered.

**Lisdexamfetamine, capsule, chewable tablet, 10mg, 20mg, 30mg, 40mg, 50mg, 60mg (Vyvanse-TAK and generics)**

For treatment of patients with Attention Deficit Hyperactivity Disorder (ADHD) who have tried extended release methylphenidate, dexamphetamine or mixed salts amphetamine with unsatisfactory results.

Claim Note:

The maximum dose reimbursed is 60mg daily.

**Lonsurf** – see Trifluridine & Tipiracil

**Lorbrena** – see Lorlatinib

**Lorlatinib, tablet, 25mg, 100mg (Lorbrena-PFI)**

As monotherapy for the first-line treatment of adult patients with anaplastic lymphoma kinase (ALK)- positive locally advanced (not amenable to curative therapy) or metastatic non-small cell lung cancer.

Clinical Note:

1. Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.

Claim Notes:

1. Approval period: 1 year.
2. No further ALK inhibitor will be reimbursed following disease progression on lorlatinib.

\* Claims that exceed the maximum claim amount of \$9999.99 must be divided as separate transactions, using DIN first and PDIN\* second.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Losec** - see Proton Pump Inhibitors

## Low Molecular Weight Heparins

**Dalteparin, pre-filled syringe, 2500 iu, 5000 iu, 7500 iu, 10000 iu, 12500 iu, 15000 iu, 18000 iu; multi-dose vial (3.8mL), 25000 iu/mL (Fragmin-PFI)**

**Tinzaparin, vial, 10000 IU/mL, 20000 IU/mL; syringe, 2500 IU/0.25mL, 3500 IU/0.35mL, 4500 IU/0.45mL, 10000 IU/0.5mL, 14000 IU/0.7mL, 18000unit/0.9mL (Innohep-LEO)**

Note: For Nursing Home Program, no Special Authorization is required.

For the acute treatment of deep vein thrombosis (DVT) and/or pulmonary embolism (PE) for a maximum of 30 days.

For prophylaxis in hip replacement and hip fracture surgery, approval is limited to a maximum of 35 days.

For prophylaxis in knee replacement surgery, approval is limited to a maximum of 10 days.

For prophylaxis in high risk surgery, approval is limited to maximum of 10 days.

For the extended treatment of recurrent symptomatic venous thromboembolism (VTE) that has occurred while patients are on therapeutic doses of warfarin.

For the treatment and secondary prevention of symptomatic venous thromboembolism (VTE) or pulmonary embolism (PE) for a period of up to 6 months in patients with cancer.

**Lucentis** - see Ranibizumab

**Luspatercept, vial, 25mg, 75mg (Reblozyl)**

### **Beta-Thalassemia Anemia**

For the treatment of adult patients with RBC transfusion-dependent anemia associated with beta-thalassemia. Patients must be receiving regular transfusions, defined as:

- 6 to 20 RBC units in the 24 weeks prior to initiating treatment with luspatercept, AND
- No transfusion-free period greater than 35 days in the 24 weeks prior to initiating treatment with luspatercept.

Renewal Criteria:

- Patients must demonstrate an initial response, defined as a  $\geq 33\%$  reduction in transfusion burden (RBC units/time) compared to the pre-treatment baseline RBC transfusion burden, measured over 24 weeks prior to initiating treatment with luspatercept.

- For continued coverage, patients should maintain a reduction in transfusion burden of  $\geq 33\%$  compared to the pre-luspatercept transfusion burden.
- Luspatercept should be discontinued if a patient does not respond after nine weeks of treatment (three doses) at the maximum dose.

Claim Notes:

- The patient should be under the care of a specialist with experience in managing patients with beta-thalassemia.
- The maximum dose of luspatercept should not exceed 1.25mg/kg (or 120mg total dose) once every three weeks.
- Initial Approval: 6 months
- Renewal Approval: 1 year

**Myelodysplastic Syndromes**

For the treatment of adult patients with red blood cell (RBC) transfusion–dependent anemia associated with very low- to intermediate-risk MDS who have ring sideroblasts and who have failed or are not suitable for erythropoietin-based therapy.

Renewal Criteria:

- Patients should be RBC transfusion independent over a minimum of 16 consecutive weeks within the first 24 weeks of treatment initiation.
- For continued coverage, patients should be RBC transfusion independent over a minimum of 16 consecutive weeks within the previous approval period.

Claim Notes:

- Treatment should be initiated by a specialist with expertise in managing and treating patients with MDS.
- The maximum dose of luspatercept should not exceed 1.75mg/kg (or 168mg total dose) once every three weeks.
- Approval: 6 months

\* Claims that exceed the maximum claim amount of \$9999.99 must be divided as separate transactions, using DIN first and PDIN\* second.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

Lynparza – see Olaparib

**Macitentan, tablet, 10mg (Opsumit-JAN)**

For the treatment of patients with Group 1 pulmonary arterial hypertension (PAH) with a World Health Organization (WHO) functional class of at least II.

**Clinical Note:**

- The diagnosis of PAH should be confirmed by right heart catheterization.

**Claim Notes:**

- Must be prescribed by, or in consultation with, a physician experienced in the treatment of PAH.
- Combined use of more than one endothelin receptor antagonists will not be reimbursed.
- The maximum dose of macitentan that will be reimbursed is 10mg daily.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Magic Bullet** - see Bisacodyl

**Mavenclad** – see Cladribine

**Mayzent** – see Siponimod

**Mecasermin, vial, 10mg/mL (Increlex-IPS)**

For the treatment of growth failure in children and adolescents from 2 to 18 years with confirmed severe primary insulin-like growth factor-1 deficiency (SPIGFD) who meet the following criteria:

- Epiphyseal closure has not yet occurred; AND
- Have a confirmed diagnosis of SPIGFD, defined by:
  - a known genetic mutation recognized as a cause of SPIGFD, AND/OR
  - has clinical and biochemical features of SPIGFD

**Renewal Criteria:**

- Treatment with mecasermin must be discontinued upon the occurrence of any of the following:
  - Height velocity is less than 1cm per 6 months or less than 2cm per year, OR
  - Bone age is more than 16 years in boys and 14 years in girls.

**Claim Notes:**

- The patient must be under the care of a pediatric endocrinologist.
- Mecasermin must not be prescribed concomitantly with recombinant GH treatment.
- Approvals: 1 year

**Mekinist** – see Trametinib

**Mektovi** – see Binimetinib

**Mepolizumab, 100mg, vial, 100mg/mL, autoinjector, syringe (Nucala-GSK)**

As add-on maintenance treatment for adult patients with severe eosinophilic asthma, if the following criteria are met:

Initiation Criteria:

- Patient must have a documented diagnosis of asthma.
- Patient is inadequately controlled with high-dose inhaled corticosteroids, defined as greater or equal to 500mcg of fluticasone propionate or equivalent daily, and one or more additional asthma controller(s) (e.g., long-acting beta agonists).
- Patient has one of the following:
  - blood eosinophil count of  $\geq 300$  cells/ $\mu$ L within the past 12 months AND has experienced two or more clinically significant asthma exacerbations in the past 12 months, or
  - blood eosinophil count of  $\geq 150$  cells/ $\mu$ L AND is receiving maintenance treatment with oral corticosteroids (OCS).

Renewal Criteria:

- The effects of treatment should be assessed every 12 months to determine whether reimbursement should continue
- Reimbursement of treatment should be discontinued if:
  - the 12 months asthma control questionnaire score has not improved from baseline, when baseline represents the initiation of treatment, or
  - the asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or the number of clinically significant exacerbations has increased within the previous 12 months, or
  - in patients on maintenance treatment with OCS, there has been no decrease in the OCS dose in the first 12 months of treatment, or
  - in patients on maintenance treatment with OCS, the reduction in the dose of OCS achieved after the first 12 months of treatment is not maintained subsequently.

Clinical Notes:

- Mepolizumab should not be used in combination with other biologics used to treat asthma.
- A baseline assessment of asthma symptom control using a validated asthma control questionnaire must be completed prior to initiation of benralizumab treatment.
- Patients should be managed by a physician with expertise in treating asthma.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Metadol** - see Methadone

**Methadone, tablet, 1mg, 5mg, 10mg, 25mg (Metadol-PMS and generic)**

For the management of severe chronic or malignant pain that is not well controlled by short and long-acting Morphine and Hydromorphone as well as Fentanyl products.

**Methylphenidate HCl, controlled release capsule, 10mg, 15mg, 20mg, 30mg, 40mg, 50mg, 60mg, 80mg (Biphentin-PFR and generic)**

For the treatment of patients with Attention Deficit Hyperactivity Disorder who have tried extended-release methylphenidate with unsatisfactory results.

Claim Note: The maximum dose reimbursed is 80 mg daily

**Methylphenidate, extended release capsule, 25mg, 35mg, 45mg, 55mg, 70mg, 85mg, 100mg (Foquest-ELV)**

For treatment of patients with Attention Deficit Hyperactivity Disorder (ADHD) who have tried extended-release methylphenidate with unsatisfactory results.

Claim Note: The maximum dose reimbursed is 100mg daily.

**Mictoryl Pediatric** – see Propiverine

**Midostaurin, capsule, 25mg (Rydapt-NVR)**

For the treatment of adult patients with newly diagnosed FMS-like tyrosine kinase 3 (FLT3)-mutated acute myeloid leukemia (AML) when used in combination with standard cytarabine and daunorubicin (7+3) induction and cytarabine consolidation chemotherapy.

Claim Notes:

- Requests for midostaurin will not be considered when used as maintenance therapy, or as part of re-induction and/or re-consolidation.
- Requests for midostaurin in combination with idarubicin containing 7+3 induction and cytarabine consolidation chemotherapy will be considered.
- Approval period: Up to 6 cycles (maximum of 2 cycles of induction and 4 cycles of consolidation)

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a**

patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

**Mirabegron, extended release tablet, 25mg, 50mg (Myrbetriq-AST)**

For the treatment of overactive bladder (not stress incontinence) in patients who cannot tolerate or have an insufficient response to an adequate trial (eg. 3 months) of immediate release oxybutynin, solifenacin, tolterodine, or tolterodine XL.

**Modafinil, tablet, 100mg (Alertec-SHR and generics)**

For the treatment of patients with a confirmed sleep-laboratory diagnosis of narcolepsy or idiopathic CNS hypersomnia.

**Mometasone Furoate/Formoterol Fumarate Dihydrate, inhaler, 50mcg/5mcg, 100mcg/5mcg, 200mcg/5mcg (Zenhale-MSD)**

For the treatment of asthma in patients 12 years of age and older who are not well controlled on a regular and adequate course of inhaled steroid therapy prior to the request for combination therapy. Continuation of current coverage requires regular use of an adequate dose of this medication.

Maximum dose is 800mcg/20mcg (4 puffs) per day

**Monurol** – see Fosfomycin

**Morphine Sulfate, injection solution, 50mg/mL (Morphine Sulfate-SAB)**

For the treatment of severe chronic pain that is not well controlled by short and long-acting oral Morphine and Hydromorphone products:

- a) For patients covered by the Nursing Home Program without a Special Authorization.
- b) For other patients upon written request or recommendation from a palliative care or pain clinic. **A copy of the recommendation must accompany the Special Authorization.**

**Moxifloxacin, ophthalmic drops, 0.5% (Vigamox-ALC and generics)**

For the treatment/prevention of bacterial conjunctivitis associated with eye surgery.

**Moxifloxacin, tablet, 400mg (Generics)**

Note: For the Cystic Fibrosis Program, no Special Authorization is required.

- a) For the treatment of severe pneumonia in nursing home patients
- b) For the completion of therapy instituted in the hospital setting for the treatment of



severe community acquired pneumonia.

## Multiple Sclerosis Agents

**Dimethyl Fumarate, delayed release capsule, 120mg, 240mg (Tecfidera-BIG & generics)**

**Glatiramer Acetate, syringe, 20mg/mL (Glatect-PMS)**

**Interferon Beta-1A, pre-filled syringe, 30mcg (Avonex PS-BIG); pre-filled cartridge, 66mcg/1.5mL, 132mcg/1.5mL (Rebif-SRO); pre-filled syringe, 22mcg, 44mcg (Rebif-SRO)**

**Interferon Beta-1B, injection powder, 0.3mg (Betaseron-BAY)**

**Peginterferon Beta-1A, SC injection, 63/94mcg/0.5mL, 125mcg/0.5mL (Plegridy-BIG)**

**Teriflunomide, tablet, 14mg (Aubagio-GZY & generics)**

Note: For glatiramer acetate naïve patients whose glatiramer therapy is initiated July 26, 2021 or later, Glatect® formulation will be approved.

For the treatment of patients 18 years of age or older, diagnosed with relapsing-remitting and secondary progressive multiple sclerosis (if applicable), who have had two attacks within the past two years, and have an EDSS score of 6.5 or less.

**The request for coverage of any of the above medications must be made by a neurologist.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

## Fingolimod, capsule, 0.5mg (Gilenya-NVR and generics)

For the treatment of patients with Relapsing Remitting Multiple Sclerosis (RRMS) who meet **all** of the following criteria:

- a) Failure to respond to full and adequate courses\* of at least one disease modifying therapy (DMT) publicly insured under PEI Pharmacare as an initial therapy, or has intolerance\*\* to at least two initial publicly funded therapies.
- b) One or more clinically disabling relapses in the previous year.
- c) Significant increase in T2 lesion load compared with that from a previous MRI scan (i.e. 3 or more new lesions) or at least one gadolinium-enhancing lesion.
- d) Requested and followed by a neurologist experienced in the management of RRMS.
- e) Recent Expanded Disability Status Scale (EDSS) score of less than or equal to 5.5 (i.e. patients must be able to ambulate at least 100 meters without assistance)

\*Failure to respond to full and adequate courses: defined as a trial of at least 6 months of a publicly funded DMT **AND** experienced at least one disabling relapse (attack) while on a publicly funded DMT (MRI report does not need to be submitted with the request).

\*\*Intolerance is defined as: documented serious adverse effects or contraindications that are incomplete with further use of that class of drug.

Dosage: 0.5 mg once daily

Approval period: Up to 12 months

**Exclusion Criteria:**

- a) Do not fund combination therapy of Gilenya with other disease modifying therapies (e.g. Avonex, Betaseron, Rebif, Tysabri) nor in combination with Fampyra.
- b) Do not fund in patients with EDSS > 5.5
- c) Do not fund in patients who have had a heart attack or stroke in the last 6 months of funding request, history of sick sinus syndrome, atrioventricular block, significant QT prolongation, bradycardia, ischemic heart disease, or congestive heart failure
- d) Patients < 18 years of age
- e) Needle phobia or preference for oral therapy over injection in patients without clinical contraindication to interferon or glatiramer therapy
- f) Skin reactions at the site of injection do NOT qualify as a contraindication to interferon or glatiramer therapy

**Renewal:**

- a) Date and details of the most recent neurological examination and EDSS scores must be provided (exam must have occurred within that last 90 days).
- b) Patients must be stable or have experienced no more than 1 disabling attack/relapse in the past year; **AND**
- c) Recent Expanded Disability Status Scale (EDSS) score of less than or equal to 5.5 (i.e. patients must be able to ambulate at least 100 meters without assistance)

Dosage: 0.5 mg once daily

Renewal period: 12 months

**The request for coverage must be made by a neurologist.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at**

<http://www.princeedwardisland.ca/pharmacareforms>.

**Alemtuzumab, vial, 12mg/1.2mL (Lemtrada-GZY)**

For the management of adult patients with relapsing-remitting multiple sclerosis (RRMS), with active disease defined by clinical and imaging features, who have had an inadequate response to two other disease-modifying therapies (DMT's), except for when any other DMT is contraindicated or unsuitable, if the following clinical criteria are met:

- At least two attacks (first episode or relapse) in the previous two years, with at least one attack in the previous year;
- At least one relapse while on at least six months of two different disease modifying therapies within the last 10 years; except for when any other DMT is contraindicated or unsuitable,
- An Expanded Disability Status Scale (EDSS) score of five (5) or less;
- Prescribed by a specialist with experience in the treatment of multiple sclerosis

\* Claims that exceed the maximum claim amount of \$9999.99 must be divided as separate transactions, using DIN first and PDIN\* second.

**The request for coverage must be made by a neurologist.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Myrbetriq** – see Mirabegron

**Nabilone, capsules, 0.5mg, 1mg (Cesamet-VAL and generics)**

- For the treatment of severe nausea and vomiting associated with cancer chemotherapy in patients who have not been well controlled by standard stepwise antiemetic therapy.
- For the treatment of acquired immune deficiency syndrome (AIDS)-related anorexia associated with weight loss.

**Nalcrom** - see Sodium Cromoglycate

**Naltrexone, tablet, 50mg (Revia-TEV and generics)**

Note: For Substance Use Harm Reduction Drug Program, no Special Authorization is required.

For the treatment of alcohol use disorder.

**Naratriptan HCl, tablet, 1mg, 2.5mg (Amerge-GSK and generics)**

For the treatment of migraine headaches where other standard therapies, such as oral analgesics have failed **AND** the patient has not responded to Zolmitriptan or Rizatriptan.

**Coverage is limited to 6 tablets per 30 day period.** Anyone requiring more than 6 doses per 30 day period should be considered for migraine prophylaxis therapy if they are not already receiving such therapy.

### **Natalizumab, vial, 300mg & 15mL (Tysabri-BIG)**

#### Initial Request:

For the treatment of Relapsing-Remitting Multiple Sclerosis (RRMS) who meet all the following criteria:

- The patient's physician is a neurologist experienced in the management of relapsing-remitting multiple sclerosis (RRMS); **AND** the patient;
  - has a current EDSS less than or equal to 5.0; **AND**
  - has failed to respond to a full and adequate course<sup>1</sup> (at least six months) of at least ONE disease modifying therapy **OR** has contraindications/intolerance to at least TWO disease modifying therapies; **AND**
  - has had ONE of the following types of relapses in the past year:
    - the occurrence of one relapse with partial recovery during the past year **AND** has at least ONE gadolinium-enhancing lesion on brain MRI, **OR** significant increase in T2 lesion load compared to a previous MRI; **OR**
    - the occurrence of two or more relapses with partial recovery during the past year; **OR**
    - the occurrence of two or more relapses with complete recovery during the past year **AND** has at least ONE gadolinium- enhancing lesion on brain MRI, **OR** significant increase in T2 lesion load compared to a previous MRI.
- Approval period: 1 year.

#### Requirements for Initial Requests:

- the patient's physician provides documentation setting out the details of the patient's most recent neurological examination within ninety (90) days of the submitted request. This must include a description of any recent attacks, the dates, and the neurological findings.
- MRI reports do NOT need to be submitted with the initial request.

#### Renewal:

Date and details of the most recent neurological examination and EDSS scores must be provided (exam must have occurred within the last 90 days); **AND**

- Patients must be stable or have experienced no more than 1 disabling attack/relapse in the past year; **AND**
- Recent Expanded Disability Status Scale (EDSS) score less than or equal to 5.0.

1.Failure to respond to a full and adequate course is defined as a trial of at least one approved first line therapy for a minimum of 6 months AND experienced at least one disabling relapse (attack) while on this

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Netupitant & Palonosetron, capsule, 300mg/0.5mg (Akynzeo-PFR)**

In combination with dexamethasone for the prevention of acute and delayed nausea and vomiting in patients receiving:

- highly emetogenic chemotherapy or
- moderately emetogenic chemotherapy who have had inadequate symptom control using a 5-HT3 antagonist and dexamethasone in a previous cycle.

Clinical notes:

- Highly emetogenic chemotherapy (HEC) includes but it not limited to : cisplatin regimens, anthracycline and cyclophosphamide combination regimens, and regimens containing carmustine, mechlorethamine, streptozocin, dacarbazine, and cyclophosphamide > 1500mg/m<sup>2</sup>
- Patients who receive carboplatin-based regimens with AUC ≥ 4 are also eligible to receive netupitant/palonosetron in combination with dexamethasone for primary prevention of acute and delayed nausea and vomiting

**Neupro** – see Rotigotine

**Nexavar** - see Sorafenib

**Ngenla** – see Somatrogen

**Nilotinib, capsule, 150mg, 200mg (Tasigna-NVR)**

For the treatment of leukemia (CML, progressed or intolerant of imatinib)

- a) As a single second line agent for the treatment of adults with chronic or accelerated phase CML with resistance or intolerance to prior therapy.

These second line criteria include:

- Patients with CML in chronic phase who are intolerant to oral tyrosine kinase inhibitors (TKIs) (i.e. imatinib or dasatinib or both)
- Patients with CML in chronic phase who are resistant to imatinib
- Patients with CML that have progressed to accelerated phase while on imatinib therapy

- b) In any one patient, only two of the TKIs will be funded within these criteria during their lifetime.

- c) If a patient develops grade 3 or 4 toxicity to one of the TKIs used within 3 months of initiating therapy, access to a third agent will be funded.
- d) Sequential use of nilotinib and dasatinib is not permitted except in the circumstance described above (i.e. grade 3 or 4 toxicity).

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Nintedanib, capsule, 100mg, 150mg (Ofev-BOE)**

#### **Chronic Fibrosing Interstitial Lung Disease**

For the treatment of adult patients with chronic fibrosing interstitial lung disease (ILD) with a progressive phenotype and a forced vital capacity (FVC) greater than or equal to 45% of predicted.

#### **Renewal criteria:**

Patient must not demonstrate progression of a disease defined as an absolute decline in percent predicted FVC of greater than or equal to 10% over the preceding 12 months of treatment with nintedanib.

#### **Claim Notes:**

- Must be prescribed by, or in consultation with a physician experience in the treatment of ILD.
- Combination therapy of Ofev (nintedanib) and Esbriet (pirfenidone) will not be reimbursed
- Approval period: 1 year

#### **Idiopathic Pulmonary Fibrosis**

For the treatment of mild to moderate idiopathic pulmonary fibrosis in adult patients confirmed by a respirologist and a high-resolution CT scan within the previous 24 months. All other causes of restrictive lung disease should be excluded. Mild to moderate IPF is defined as forced vital capacity (FVC) greater than or equal to 50% of predicted.

#### **Initial renewal criteria (at 6 months):**

Patients must not demonstrate progression of disease defined as an absolute decline in percent predicted forced vital capacity (FVC) of  $\geq 10\%$  from initiation of therapy until renewal (initial 6 month treatment period). If a patient has experienced progression as defined above, then the results should be validated with a confirmatory pulmonary function test conducted 4 weeks later.

**Second and subsequent renewals (at 12 months and thereafter):**

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.

**Excluded criteria:**

Combination therapy of Ofev (nintedanib) and Esbriet (perfenidone) will not be reimbursed.

**Note:**

Patients who have experienced intolerance or failure to nintedanib or perfenidone will be considered for the alternate agent provided that the patient continues to meet the above coverage criteria.

**Requests for Nintedanib (Ofev-BOE) must be made using the Nintedanib/Pirfenidone Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Niraparib, capsule, 100mg; tablet, 100mg (Zejula-GSK)**

1. As monotherapy maintenance treatment of patients with newly-diagnosed ovarian, fallopian tube, or primary peritoneal cancer who are in response (complete or partial) to at least 4 cycles of first-line platinum-based chemotherapy. Eligible patients should have high-grade serous or endometrioid tumours classified as stage III or IV according to the International Federation of Gynecology and Obstetrics (FIGO) criteria.

**Clinical Notes:**

- Patients should have a good performance status.
  - Maintenance therapy with niraparib should begin within 12 weeks of completion of platinum-based chemotherapy and may continue for up to 3 years, or until disease progression or unacceptable toxicity, whichever occurs first.
  - Patients who have stable brain metastases are eligible for treatment with niraparib.
  - Patients who are unable to tolerate platinum-based chemotherapy (due to allergic reaction) and otherwise meet criteria, will be assessed on a case by case basis to determine eligibility for treatment with niraparib.
  - Niraparib in combination with bevacizumab is not funded
2. As monotherapy maintenance treatment for patients with relapsed, platinum-sensitive high grade serous epithelial ovarian, fallopian tube, or primary peritoneal cancer who have completed at least two previous lines of platinum-based chemotherapy, and have

achieved a complete or partial response to the most recent platinum-based chemotherapy regimen.

**Clinical Notes:**

- Platinum-sensitive disease is defined as disease progression occurring at least six months after completion of platinum-based chemotherapy.
- Patients should have a good performance status.
- Patients must have received at least 4 cycles of the most recent platinum-based chemotherapy before starting treatment with niraparib.
- Maintenance therapy with niraparib should begin within 12 weeks of the last chemotherapy treatment and may continue until disease progression or unacceptable toxicity, whichever occurs first.
- Patients who have stable brain metastases are eligible for treatment with niraparib.
- Patients who are unable to tolerate platinum-based chemotherapy (due to allergic reaction) and otherwise meet criteria, will be assessed on a case by case basis to determine eligibility for treatment with niraparib.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Nirmatrelvir & Ritonavir, tablet, 150mg & 100mg (Paxlovid-PFI)**

For the treatment of mild to moderate coronavirus disease 2019 (COVID-19) in adult patients with a positive COVID-19 test who are within 5 days of symptom onset and meet one of the following criteria:

- Severely immunosuppressed **due to one or more of the following conditions:**
  - Solid organ transplant
  - Receiving treatment for a malignant hematologic condition
  - Bone marrow transplant, stem cell transplant or transplant-related immunosuppressant use
  - Received an anti-CD20 therapy or B-cell depleting therapy (such as rituximab) in the previous two years
  - Severe primary immunodeficiencies
- Moderately immunosuppressed **due to one or more of the following conditions:**
  - Receiving treatment for cancer, including solid tumors
  - Receiving treatment with significantly immunosuppressing drugs (e.g., biologic in the past three months, oral immune-suppressing drug in the past month, oral steroid [20 mg per day of prednisone equivalent taken on an ongoing basis] in the past month, or immune-suppressing infusion or injection in the past three months)
  - Advanced HIV infection



- Moderate primary immunodeficiencies
- Renal conditions (i.e., hemodialysis, peritoneal dialysis, glomerulonephritis treated with a steroid, eGFR less than 15 mL/min/1.73m<sup>2</sup>)

Clinical Notes:

- COVID-19 testing to confirm diagnosis can be performed by polymerase chain reaction (PCR) or point-of-care test (POCT).
- Treatment should be initiated as soon as possible after a diagnosis of COVID-19 is confirmed.
- Patients are not eligible for coverage if they are asymptomatic or if more than 5 days have elapsed since symptom onset.
- Requests for patients who are moderately or severely immunosuppressed due to other conditions may be considered.

Claim Notes:

- Pharmacists must verify eligibility criteria prior to dispensing and provide a copy of the *Pharmacist Initiated Treatment of COVID-19 Paxlovid Special Authorization* form to Pharmacare.
- Approval period: 5 days.

Pharmacist Prescribers:

- Completion of the *Pharmacist Initiated Treatment of Covid-19 Paxlovid Special Authorization Form* is required. The completed form must be faxed to Pharmacare the day of dispensing.
- Pharmacies do not have to wait for special authorization approval by Pharmacare prior to dispensing.
- Please contact Pharmacare if considering Paxlovid coverage for patients who are moderately or severely immunosuppressed due to other conditions not defined above.

Note: Non-pharmacist prescribers are not required to submit a Special Authorization form when prescribing Paxlovid.

**Nivestym** – see Filgrastim

**Norfloxacin, tablet, 400mg (Generics)**

Note: For Nursing Home program no Special Authorization is required.

- a) For the treatment of urinary tract infections caused by *Pseudomonas aeruginosa*. Up to 10 days of therapy will be considered.
- b) For the treatment of urinary tract infections not responding to alternative therapy. Up to 10 days of therapy will be considered.
- c) For the treatment of urinary tract infections in persons allergic to alternative agents. Up to 10 days of therapy will be considered.
- d) Prophylaxis of chronic urinary tract infections in persons allergic to alternative agents or where prophylaxis with alternative agents has failed.

**(Note: Recommended dosage is 200mg at bedtime)**

**Novorapid** – see insulin aspart

**Nubeqa** – see Darolutamide

**Nucala** – see Mepolizumab

**Nusinersen, intrathecal vial, 2.4mg/mL (Spinraza-BIG)**

For patients diagnosed with 5q Spinal Muscular Atrophy (SMA) if the following clinical criteria are met:

- 1) Genetic documentation of 5q SMA homozygous gene deletion, homozygous mutation, or compound heterozygote, AND
- 2) Patients who:
  - are pre-symptomatic with two or three copies of SMN2, OR
  - have had disease duration of less than six months, two copies of SMN2, and symptom onset after the first week after birth and on or before seven months of age, OR
  - are under the age of 18 with symptom onset after six months of age, AND
- 3) Patient is not currently requiring permanent invasive ventilation\*, AND
- 4) A baseline assessment using an age-appropriate scale (the Hammersmith Infant Neurological Examination [HINE] Section 2, Children’s Hospital of Philadelphia Infant Test of Neuromuscular Disorders [CHOP INTEND], or Hammersmith Functional Motor Scale-Expanded [HFMSSE]) must be completed prior to initiation of nusinersen treatment.

Other patients with SMA type 2 or 3 who are over the age of 18 may be considered on a case by case basis.

For continued coverage, the patient must meet the following criteria:

- 1) There is demonstrated achievement or maintenance of motor milestone function (as assessed using age-appropriate scales: the [HINE] Section 2), CHOP INTEND, or HFMSSE since treatment initiation in patients who were pre- symptomatic at the time of treatment initiation; OR  
There is demonstrated maintenance of motor milestone function (as assessed using age-appropriate scales: the HINE Section 2, CHOP INTEND, or HFMSSE since treatment initiation in patients who were symptomatic at the time of treatment initiation;  
AND
- 2) Patient does not require permanent invasive ventilation\*.

Treatment should be discontinued if, prior to the fifth dose or every subsequent dose of nusinersen, the above renewal criteria are not met.

\* Permanent invasive ventilation is defined as the use of tracheostomy and a ventilator

due to progression of SMA that is not due to an identifiable and reversible cause.

Claim Notes:

- The patient must be under the care of a specialist experienced in the treatment of SMA.
- Approval Period: 1 year.
- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions using the DIN first, and then PDINs.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at [www.healthpei.ca/pharmacareforms](http://www.healthpei.ca/pharmacareforms)**

**Nyvepria** – see Pegfilgrastim

**Obeticholic, tablet, 5mg, 10mg (Ocaliva-INT)**

For the treatment of adult patients with primary biliary cholangitis (PBC) as either:

- combination therapy with ursodeoxycholic acid (UDCA) in patients who have experienced an inadequate response to a minimum of 12 months of UDCA treatment; or
- monotherapy in patients who have experienced unmanageable intolerance to UDCA.

Requirement for Initial Requests:

- Alkaline phosphatase (ALP) and bilirubin levels prior to initiation of treatment with obeticholic acid must be provided.

Renewal Criteria:

- Requests for renewal will be considered if the patient achieved:
  - a reduction in the ALP to less than 1.67 times the upper limit of normal (ULN); or
  - at least a 15% reduction in the ALP level from baseline (i.e. prior to initiation of treatment with obeticholic acid).

Clinical Notes:

- Diagnosis confirmed by positive antimitochondrial antibodies or liver biopsy results consistent with PBC.
- An inadequate response is defined as:
  - $ALP \geq 1.67$  times ULN, or
  - bilirubin  $>$  ULN and  $<$  2 times the ULN, or
  - evidence of compensated cirrhosis.
- For patients who experience unmanageable intolerance to UDCA, details must be provided.

Claim Notes:

- Must be prescribed by, or in consultation with, a gastroenterologist, hepatologist or other physician experienced in the treatment of PBC.
- Approval period: 12 months.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ocaliva** – see Obeticholic

**Ocrelizumab, vial, 300 mg/10 mL (Ocrevus- HLR)**

**Relapsing Remitting Multiple Sclerosis**

For the treatment of adult patients with relapsing remitting multiple sclerosis (RRMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Experienced one or more disabling relapses or new MRI activity in the last two years
- Ambulatory with or without aid (i.e. has a recent Expanded Disability Status Scale (EDSS) score of less than or equal to 6.5)
- Must be prescribed by a neurologist with experience in the diagnosis and management of multiple sclerosis.

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.

**Primary Progressive Multiple Sclerosis**

For the treatment of adult patients with early primary progressive multiple sclerosis (PPMS) who meet all of the following criteria:

- Confirmed diagnosis based on McDonald criteria
- Recent Expanded Disability Status Scale (EDSS) score between 3.0 and 6.5
- Recent Functional Systems Scale (FSS) score of at least 2 for the pyramidal functions component due to lower extremity findings
- Disease duration of 10 years for those with an EDSS of less than or equal to 5 or disease duration less than 15 years for those with an EDSS greater than 5
- Diagnostic imaging features characteristic of inflammatory activity

Clinical Note:

- Treatment should be discontinued for patients with an EDSS score of greater than or equal to 7.
- Must be prescribed by a neurologist with experience in the diagnosis and management of multiple sclerosis.

Claim Note:

- Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions using the DIN first, and then the PDIN.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ocrevus** – see Ocrelizumab

**Ocuflox** - see Ofloxacin

**Ofatumumab, prefilled pen, 20mg/0.4mL (Kesimpta-NVR)**

For the treatment of adult patients with relapsing remitting multiple sclerosis (RRMS) who meet all of the following criteria:

- An Expanded Disability Status Scale (EDSS) score of less than 6.0
- Evidence of active disease defined as at least one of the following:
  - One relapse during the previous year
  - Two relapses during the previous 2 years
  - A positive gadolinium (Gd)-enhancing MRI scan during the year before starting treatment with ofatumumab.

Renewal Criteria:

- EDSS score less than 6.0. Date and details of the most recent neurological examination and EDSS score must be provided (exam must have occurred within the last 90 days); AND
- Patients must be stable or have experienced no more than 1 disabling attack/relapse in the past year.

Claim Notes:

- Approval: 1 year.
- Combined use with other disease modifying therapies to treat multiple sclerosis will not be reimbursed.
- Must be prescribed by a neurologist with experience in the diagnosis and management of multiple sclerosis.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ofev** – see Nintedanib

**Olaparib, tablet, 100 mg, 150 mg (Lynparza- AZE)**

### **High-Risk Early Breast Cancer**

1. For the adjuvant treatment of adult patients with deleterious or suspected deleterious germline BRCA-mutated high-risk early breast cancer who have had upfront surgery followed by adjuvant chemotherapy and who meet one of the following criteria:
  - Triple negative breast cancer and either axillary node-positive or axillary node-negative with invasive primary tumor pathological size of at least 2 cm (> pT2 cm)
  - Hormone receptor positive, HER2-negative breast cancer with at least 4 pathologically confirmed positive lymph nodes.
  
2. For the adjuvant treatment of adult patients with deleterious or suspected deleterious germline BRCA-mutated high-risk early breast cancer who received neoadjuvant chemotherapy followed by surgery and who meet one of the following criteria:
  - Triple negative breast cancer with residual invasive disease in the breast and/or resected lymph nodes (nonpCR)
  - Hormone receptor positive, HER2-negative breast cancer with residual invasive disease in the breast, and/or the resected lymph nodes, and a CPS + EG score of 3 or higher.

#### Clinical Notes:

1. Patients must have completed neoadjuvant or adjuvant chemotherapy containing an anthracycline and/or taxane.
2. Treatment should be initiated within 12 weeks of completion of the last treatment (i.e., surgery, chemotherapy, or radiation therapy).
3. Patients must have a good performance status.
4. Treatment should be discontinued upon disease recurrence, unacceptable toxicity, or completion of 1 year of therapy, whichever occurs first.

#### Claim Notes:

- Requests for patients determined to be at high-risk for relapse using a disease scoring system other than CPS + EG will be considered.
- Approval period: 1 year

### **Newly Diagnosed, Advanced, BRCA-Mutated**

- As monotherapy maintenance treatment of patients with newly-diagnosed, advanced, BRCA-mutated (germline or somatic), high-grade epithelial ovarian, fallopian tube, or primary peritoneal cancer who are in response (complete or partial) to at least 4 cycles of first-line platinum-based chemotherapy.

#### Clinical Notes:

- Patients should have a good performance status.

- Maintenance therapy with olaparib should begin within 12 weeks of completion of platinum-based chemotherapy.
- Patients who are unable to tolerate platinum-based chemotherapy (due to allergic reaction) and otherwise meet criteria, will be assessed on a case by case basis to determine eligibility for treatment with olaparib.
- Treatment should continue until unacceptable toxicity, disease progression, or to a maximum of 2 years of therapy if no evidence of disease, whichever comes first.<sup>1</sup>
- Imaging is required for patients who are delayed in starting olaparib therapy, i.e. greater than 12 weeks after completion of platinum-based chemotherapy, or who have had a break in therapy for more than 14 days, to rule out progression prior to starting or re-starting olaparib.
- Olaparib in combination with bevacizumab is not funded. Patients already on bevacizumab maintenance at the time of olaparib funding may be switched to olaparib, as long as there is no evidence of progression on imaging and is within 12 weeks of completion of chemotherapy.

<sup>1</sup>Patients with a partial response or stable disease at 2 years may continue to receive olaparib at the discretion of the treating physician.

### **Relapsed, BRCA-Mutated**

- As monotherapy maintenance treatment for patients with platinum-sensitive, relapsed, BRCA-mutated (germline or somatic), high grade serous epithelial ovarian, fallopian tube, or primary peritoneal cancer who have completed at least two previous lines of platinum-based chemotherapy and are in radiologic response (complete or partial) to their most recent platinum-based chemotherapy regimen as per the SOLO-2 trial.
- Patients must have received at least four cycles of their most recent platinum-based chemotherapy before starting treatment with olaparib.

#### Clinical Notes:

- Treatment should continue until unacceptable toxicity or disease progression.
- Maintenance therapy with olaparib should begin within eight weeks of the last dose of platinum-based chemotherapy.
- Platinum-sensitive disease is defined as disease progression occurring at least six months after completion of platinum-based chemotherapy.
- Patients should have a good performance status.
- Patients who are unable to tolerate platinum-based chemotherapy (due to allergic reaction) and otherwise meet criteria, will be assessed on a case by case basis to determine eligibility for treatment with olaparib.

### **Metastatic Castrate-Resistant Prostate Cancer**

- For the treatment of patients with metastatic castration-resistant prostate cancer (mCRPC) who meet all of the following criteria:

- deleterious or suspected deleterious germline and/or somatic mutations in the homologous recombination repair (HRR) genes BRCA1, BRCA2 or ATM; and
- Disease progression on prior treatment with androgen-receptor-axis-targeted (ARAT) therapy.

Renewal Criteria:

Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

- Patients must have a good performance status.
- Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ofloxacin, ophthalmic solution, 0.3% (Ocuflox and generics)**

Note: For Nursing Home Program, no Special Authorization is required.

For the treatment of ophthalmic infections caused by susceptible bacteria and not responding to alternative agents.

**Olumiant** – see Baricitinib

**Omalizumab, vial, 150mg, prefilled syringe, 150mg (Xolair-NVR)**

For the treatment of patients ≥ 12 years of age with moderate to severe chronic idiopathic urticaria (CIU) who remain symptomatic (presence of hives and/or associated itching) despite optimum management with H1 antihistamines.

Initiation Criteria:

- Documentation of the most recent Urticaria Activity Score over 7 days (UAS7) to be provided on the submitted request.
- Approvals will be for a maximum dose of 300mg every four weeks.
- Initial approval period: 24 weeks.

Renewal Criteria:

- Requests for renewal will be considered if the patient has achieved:
  - Complete symptom control for less than 12 consecutive weeks; or
  - Partial response to treatment, defined as at least a ≥ 9.5 point reduction in baseline UAS7



Clinical Notes:

1. Moderate to severe CIU is defined as UAS7  $\geq$ 16.
2. Treatment cessation could be considered for patients who experience complete symptom control for at least 12 consecutive weeks at the end of a 24 week treatment period.
3. In patients who discontinue treatment due to temporary symptom control, re-initiation can be considered if CIU symptoms reappear.
4. Optimal management is defined as H1 antihistamines at up to 4 times the standard daily dose.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Omeprazole** - see Proton Pump Inhibitors

**Onabotulinumtoxina, injection, 200 units/vial (Botox-ALL)**

For the treatment of urinary incontinence due to neurogenic detrusor overactivity resulting from neurogenic bladder associated with multiple sclerosis (MS) or subcervical spinal cord injury (SCI) if the following conditions are met:

- patient failed to respond to behavioural modification and anticholinergics and/or is intolerant to anticholinergics.
- subsequent treatments are provided at intervals no less than every 36 weeks.
- Patients who fail to respond to initial treatment with onabotulinumtoxinA should not be retreated.

**Ondansetron, medicated film, 4mg, 8mg (Ondissolve-TAK and generic)**

For the treatment of emesis in cancer patients receiving highly emetogenic chemotherapy (i.e. containing Cisplatin); receiving moderately emetogenic chemotherapy (i.e. containing Cyclophosphamide, Doxorubicin, Epirubicin, or Melphalan); or receiving radiation therapy and who have:

- a) Experienced adverse effects to Metoclopramide, Prochlorperazine, or Dexamethasone or have a specific contraindication which does not allow use of these drugs as antiemetics or,
- b) Continued episodes of nausea and vomiting related to chemotherapy which have not responded to therapeutic doses of Metoclopramide, Prochlorperazine, or Dexamethasone.

**A maximum of 10 films per cycle of chemotherapy will be approved.**

**Ondansetron HCl, tablet, 4mg, 8mg (Zofran-GSK and generics)**

For the treatment of emesis in cancer patients receiving highly emetogenic chemotherapy (i.e. containing Cisplatin); receiving moderately emetogenic chemotherapy (i.e. containing

Cyclophosphamide, Doxorubicin, Epirubicin, or Melphalan); or receiving radiation therapy and who have:

- a) Experienced adverse effects to Metoclopramide, Prochlorperazine, or Dexamethasone or have a specific contraindication which does not allow use of these drugs as antiemetics or,
- b) Continued episodes of nausea and vomiting related to chemotherapy which have not responded to therapeutic doses of Metoclopramide, Prochlorperazine, or Dexamethasone.

**A maximum of 10 tablets per cycle of chemotherapy will be approved.**

**Only requests for the oral dosage forms are eligible for consideration.**

**Ondansetron, oral disintegrating tablets, 4mg, 8mg (Generic)**

For the treatment of emesis in cancer patients receiving highly emetogenic chemotherapy (i.e. containing Cisplatin); receiving moderately emetogenic chemotherapy (i.e. containing Cyclophosphamide, Doxorubicin, Epirubicin, or Melphalan); or receiving radiation therapy and who have:

- a) Experienced adverse effects to Metoclopramide, Prochlorperazine, or Dexamethasone or have a specific contraindication which does not allow use of these drugs as antiemetics or,
- b) Continued episodes of nausea and vomiting related to chemotherapy which have not responded to therapeutic doses of Metoclopramide, Prochlorperazine, or Dexamethasone.

**A maximum of 10 tablets per cycle of chemotherapy will be approved.**

**Ondissolve** – see Ondansetron

**Onureg** – see Azacitidine

**Opsumit** – see Macitentan

**Orkambi** – granule packet, 100 mg/125 mg, 150mg/188 mg; tablet, 100 mg/125 mg, 200 mg/125 mg

Coverage may be available for Orkambi for the treatment of cystic fibrosis patients who meet certain medical criteria and drug program enrollment.

Please contact the PEI Pharmacare Program office at 1-877-577-3737 for more information regarding coverage availability and the Special Authorization application process for this product.

**Osimertinib, tablet, 40mg, 80mg (Tagrisso-AZE)**

1. For the first-line treatment of patients with locally advanced (not amenable to curative intent therapy) or metastatic non-small cell lung cancer (NSCLC) whose tumors have epidermal growth factor receptor (EGFR) mutations (exon 19 deletions [exon 19 del] or exon 21 [L858R]).
2. In patients with locally advanced or metastatic epidermal growth factor receptor (EGFR) T790M mutation-positive non-small cell lung cancer (NSCLC) who have progressed on EGFR tyrosine kinase inhibitor (TKI) therapy.

Clinical Notes:

- Eligible patients should be previously untreated in the locally advanced or metastatic setting and have a good performance status.
  - Treatment should be discontinued upon clinically meaningful disease progression or unacceptable toxicity.
  - Prior treatment with EGFR TKI therapy is not required in patients with de novo T790M mutation-positive NSCLC.
3. For adjuvant therapy after tumour resection in patients with Stage IB-IIIa (AJCC 7th edition or equivalent) non-small cell lung cancer (NSCLC) whose tumors have epidermal growth factor receptor (EGFR) exon 19 deletions [exon 19 del] or exon 21 [L858R] substitution mutations.

Clinical Notes:

- Patients should have a good performance status.
- Treatment with osimertinib should continue for a total duration of 3 years, or until disease recurrence or unacceptable toxicity.
- Osimertinib treatment should be initiated within 10 weeks of complete surgical resection if adjuvant chemotherapy was not administered, or within 26 weeks if adjuvant chemotherapy was administered.
- Retreatment with osimertinib in the metastatic setting will be considered if disease recurrence is at least 6 months following completion of adjuvant therapy
- Program eligibility remains the same (Financial Assistance Drug Program, High Cost Drug Program, Nursing Home Drug Program, Catastrophic Drug Program).

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Oxeze** - see Formoterol

**Oxcarbazepine, tablet, 150mg, 300mg, 600mg (Trileptal-NVR and generics)**

For use in patients who have a diagnosis of epilepsy and have had an inadequate response to or are intolerant to at least 3 other formulary agents (prior or current use), including Carbamazepine.

**Ozanimod, capsule, 0.23mg & 0.46mg, 0.92mg (Zeposia-BMS)**

For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score > 4, and a rectal bleeding subscore  $\geq 2$  and are:

- Refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks AND prednisone  $\geq 40$ mg daily for two weeks or IV equivalent for one week)  
OR
- Corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Initial Approval: 12 weeks. Treatment has to be initiated in all patients with an initiation pack that lasts for 7 days.
  - Days 1-4 0.23 mg once daily
  - Days 5-7 0.46 mg once daily
  - Days 8 and thereafter 0.92 mg once daily.
- Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:
  - a decrease in the partial Mayo score  $\geq 2$  from baseline, and
  - a decrease in the rectal bleeding subscore  $\geq 1$ .
- Renewal Approval: 1 year. Maximum approved dose is 0.92mg once daily.
- Combined use of more than one biologic DMARD will not be reimbursed.

Clinical Notes:

- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.
- Patients with severe disease (partial Mayo > 6) do not require a trial of 5-ASA.

**The request for coverage must be made by a gastroenterologist using the Ulcerative Colitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a**

patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

**Ozempic** – see Semaglutide

**Palbociclib, capsules & tablets, 75mg, 100mg, 125mg (Ibrance-PFI)**

1. In combination with an aromatase inhibitor for the treatment of estrogen receptor positive, HER2 negative advanced breast cancer in postmenopausal women who:
  - have not received prior therapy for metastatic disease and
  - are not resistant to (neo) adjuvant non-steroidal aromatase inhibitor (NSAI) therapy and
  - do not have active or uncontrolled metastasis to the central nervous system.

Renewal criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

- Patients must have a good performance status.
- Resistance is defined as disease progression occurring during or within 12 months following NSAI therapy.
- Treatment should be discontinued up on disease progression or unacceptable toxicity.

Claim Notes:

Initial approval period: 1 year

Renewal approval period: 1 year

2. In combination with fulvestrant for the treatment of patients with hormone receptor (HR) positive, HER 2 negative advanced or metastatic breast cancer as initial endocrine-based therapy or following disease progression on endocrine therapy. Patients may have also received up to one prior line of chemotherapy for advanced disease. Patients should have a good performance status, without active or uncontrolled metastases to the central nervous system and in the case of women can be of any menopausal status (Perimenopausal and premenopausal women must be treated with an LHRH agonist).

Clinical Notes:

- Treatment should continue until unacceptable toxicity or disease progression.
- Patients who progress  $\leq$  12 months from (neo) adjuvant therapy are eligible for treatment with palbociclib plus fulvestrant.
- Patients who experience disease progression on prior CDK 4/6 inhibitor therapy, fulvestrant or everolimus are not eligible for treatment with palbociclib with fulvestrant.
- Patients currently receiving fulvestrant monotherapy, and who have not

progressed may have palbociclib added, provided they are CDK 4/6 inhibitor naïve and otherwise meet funding criteria.

- Patients who previously received everolimus plus exemestane will be eligible for funding of palbociclib plus fulvestrant on progression, provided that treatment was started prior to funding of CDK 4/6 + fulvestrant, patient must be CDK 4/6 naïve and otherwise meet funding criteria.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Paliperidone, injection, 50mg, 75mg, 100mg, 150mg (Invega Sustenna-JAN)**

Note: For Community Mental Health Drug Program, no Special Authorization is required.

For the treatment of schizophrenia or schizoaffective disorder in patients who have:

- a) A history of non adherence  
OR
- b) Inadequate control or significant side effects from two or more oral antipsychotic medications  
OR
- c) Inadequate control or significant side effects from at least one long acting depot antipsychotic agent.

Note: Must be requested and prescribed by a psychiatrist. Only doses up to 150 mg monthly will be approved.

### **Paliperidone, prefilled pen, 175mg/0.875mL, 263mg/1.315mL, 350mg/1.75mL, 525mg/2.625mL (Invega Trinza-JAN)**

Note: For Community Mental Health Drug Program, no Special Authorization is required.

For the maintenance treatment of schizophrenia and related psychotic disorders (not dementia related) in patients who have been stabilized on therapy with injectable paliperidone for at least four months.

### **Pamidronate Disodium, injection powder, 30mg, 60mg, 90mg vial (Generics)**

For the management of tumour-induced hypercalcemia following adequate saline rehydration or conditions associated with increased osteoclast activity.

**Pantoloc** - see Proton Pump Inhibitors

**Pantoprazole Magnesium** - see Proton Pump Inhibitors

**Pantoprazole Sodium** - see Proton Pump Inhibitors

**Pariet** - see Proton Pump Inhibitors

**Paxlovid** – see Nirmatrelvir & Ritonavir

**Pazopanib, tablet, 200mg (Votrient-GSK and generic)**

1. As a first-line treatment for patients with advanced or metastatic clear cell renal carcinoma and good performance status.
2. For the treatment of advanced or metastatic renal cell (clear cell) carcinoma (mRCC) in patients who are unable to tolerate sunitinib and who have an ECOG performance status of 0 or 1.

Renewal criteria:

Written confirmation that the patient has benefited from therapy and is expected to continue to do so.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Pegfilgrastim, prefilled syringe, 6mg/0.6mL (Fulphila-BGP, Lapelga-APX, Nyvepria-PFI, Ziextenzo-SDZ); prefilled syringe, 10mg/1.0mL (Lapelga-APX)**

For the prevention of febrile neutropenia in patients receiving myelosuppressive chemotherapy with curative intent who:

- are at high risk of febrile neutropenia due to chemotherapy regimen, co-morbidities or pre-existing severe neutropenia; or
- have had an episode of febrile neutropenia, neutropenic sepsis or profound neutropenia in a previous cycle of chemotherapy; or
- have had a dose reduction, or treatment delay greater than one week due to neutropenia.

Clinical Note:

- Patients with non-curative cancer receiving chemotherapy with palliative intent are not eligible for coverage of pegfilgrastim for prevention of febrile neutropenia.

**Must be requested and prescribed by a specialist in hematology or medical oncology.**

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Pentosan Polysulfate Sodium, capsule, 100mg (Elmiron-JAN)**

For the treatment of interstitial cystitis where other treatments have failed.

**Perampanel, tablet, 2mg, 4mg, 6mg, 8mg, 10mg, 12mg (Fycompa-EIS and generic)**

For the adjunctive treatment of refractory partial-onset seizures or primary generalized tonic-clonic seizures in patients who are currently receiving two or more antiepileptic drugs, and who have had an inadequate response to at least three other antiepileptic drugs.

**Perichlor** - see Chlorhexidine

**Peridex** - see Chlorhexidine

**Pilocarpine, tablet, 5mg (Salagen-PFI and generic)**

For oncology patients only, for the treatment of the symptoms of xerostomia due to salivary gland hypofunction caused by radiotherapy for cancer of the head and neck.

**Pirfenidone, capsule, 267mg, tablet, 267mg, tablet, 801mg (Esbriet-HLR and generics)**  
**Initial approval criteria:**

For the treatment of mild to moderate idiopathic pulmonary fibrosis in adult patients confirmed by a respirologist and a high-resolution CT scan within the previous 24 months. All other causes of restrictive lung disease should be excluded. Mild to moderate IPF is defined as forced vital capacity (FVC) greater than or equal to 50% of predicted.

**Initial renewal criteria (at 6 months):**

Patients must not demonstrate progression of disease defined as an absolute decline in percent predicted forced vital capacity (FVC) of  $\geq 10\%$  from initiation of therapy until renewal (initial 6 month treatment period). If a patient has experienced progression as defined above, then the results should be validated with a confirmatory pulmonary function test conducted 4 weeks later.

**Second and subsequent renewals (12 months and thereafter):**

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.

**Excluded criteria:**



Combination therapy of Ofev (nintedanib) and Esbriet (pirfenidone) will not be reimbursed.

**Note:**

Patients who have experienced intolerance or failure to nintedanib or pirfenidone will be considered for the alternate agent provided that the patient continues to meet the above coverage criteria.

**Requests for Pirfenidone (Esbriet-HLR) must be made using the Nintedanib/Pirfenidone Special Authorization form which is available from the Drug Programs office or on-line at**

**<http://www.princeedwardisland.ca/pharmacareforms> .**

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at**

**<http://www.princeedwardisland.ca/pharmacareforms> .**

**Plegridy – see Multiple Sclerosis Agents**

**Pomalidomide, capsule, 1mg, 2mg, 3mg, 4mg (Pomalyst-CEL and generics)**

For patients with relapsed and/or refractory multiple myeloma who have previously failed at least two treatments, including both bortezomib and lenalidomide and demonstrated disease progression on the last treatment.

Note: Pomalidomide may be an option in rare instances where bortezomib is not tolerated or contraindicated but in all cases, patients should have failed lenalidomide.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at**

**<http://www.princeedwardisland.ca/pharmacareforms>.**

**Pomalyst – see Pomalidomide**

**Ponatinib – tablet, 15mg (Iclusig-ARI)**

For the treatment of patients with chronic, accelerated or blast phase chronic myelogenous leukemia (CML) or Philadelphia chromosome positive acute lymphoblastic leukemia (Ph+ALL) who have:

- resistance or intolerance to two or more tyrosine kinase inhibitors (TKIs), OR
- confirmed T315i mutation positive disease.

Clinical Notes:

1. Patients must have an ECOG performance status of  $\leq 2$ .

2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Pulmozyme** – see Dornase Alfa

**Pradaxa** – see Dabigatran

**Praluent** – see Alirocumab

**Prasugrel, tablet, 10mg (Generic)**

For use in combination with ASA for patients with:

- ST-elevated myocardial infarction (STEMI) undergoing primary percutaneous coronary intervention (PCI) who have not received antiplatelet therapy prior to arrival in the catheterization lab. Treatment must be initiated in hospital.  
OR
- Acute coronary syndrome who failed on optimal clopidogrel and ASA therapy as defined by definite stent thrombosis, or recurrent STEMI or UA after prior revascularization via PCI.

Approval: up to 12 months

**Prevacid** - see Proton Pump Inhibitors

**Prevacid Fastab**

- pediatric patients 12 and under will not require written Special Authorization.
- see Proton Pump Inhibitors for criteria for all other requests

**Prevymis** – see Letemovir

**Prolia** – see Denosumab

**Propiverine, tablet, 5mg (Mictoryl Pediatric-DUI)**

For the treatment of overactive bladder with symptoms of urgency incontinence and/or urinary frequency and urgency in pediatric patients under 18 years of age.

**Propranolol, oral solution, 3.75mg/mL (Hemangirol-PFB)**

For the treatment of patients with proliferating infantile hemangioma that is:

- Life-or function-threatening OR
- Ulcerated with pain or not responding to simple wound care measures OR
- At risk of permanent scarring or disfigurement

**Proton Pump Inhibitors**

**Lansoprazole, delayed release capsule, 15mg, 30mg (Prevacid-ABB and generics);  
 Lansoprazole, delayed release tablet, 15mg, 30mg (Prevacid Fastab-ABB);  
 Omeprazole, capsule, 20mg (Losec-AZE and generics);  
 Omeprazole, delayed release tablet, 20mg (Losec-AZE and generics);  
 Pantoprazole Magnesium, enteric tablet, 40mg (Tecta-TAK and generics)  
 Pantoprazole Sodium, enteric tablet, 20mg, 40mg (Pantoloc-TAK and generics);  
 Rabeprazole, tablet, 10mg, 20mg (Pariet-JAN and generics)**

**\* Doses of Omeprazole 20mg daily, Pantoprazole Magnesium 40mg daily ,  
 Pantoprazole Sodium 20mg or 40mg up to one unit daily, and up to Rabeprazole  
 20mg daily DO NOT require a Special Authorization.**

For doses of Omeprazole and Rabeprazole greater than 20mg per day and greater than 40mg per day of Pantoprazole Magnesium, greater than one unit/day of Pantoprazole Sodium 20mg or 40mg, and all doses of Lansoprazole **WHERE** evidence is provided of resistance to two **recent** 12 week trials (ie within 6 months) of a standard dose (20mg daily) of Omeprazole, Rabeprazole, Pantoprazole Magnesium 40mg daily and greater than one unit per day of Pantoprazole Sodium 20mg or 40mg.

Up to 12 weeks of therapy will be considered for

- a) Gastric and Duodenal Ulcers
- b) Esophagitis

Long term therapy will be considered for

- c) Erosive Esophagitis
- d) Barrett’s Esophagitis
- e) Zollinger-Ellison Syndrome
  
- f) Helicobacter pylori Eradication – Up to 14 days of twice daily dosing for patients who are registered in an eligible Pharmacare Program, are symptomatic and have a documented positive Helicobacter Pylori test

**Protopic** - see Tacrolimus

**Pulmicort Nebuamps** - see Budesonide

**Qulipta** – see Atogepant

**Qinlock** – see Ripretinib

**Rabeprazole** - see Proton Pump Inhibitors

**Radicava** – see Edaravone

**Ranibizumab, vial, 2.3mg/ 0.23mL (Lucentis-NVR); vial, 2.3mg/0.23mL (Byooviz-BIG); vial, 2.4mg/0.23mL (Ranopto-TEV)**

**1. Neovascular Age-Related Macular Degeneration**

Initial Coverage:

For the treatment of patients with neovascular (wet) age-related macular degeneration (AMD) where all of the following apply:

- a) Best Corrected Visual Acuity (BCVA) is between 6/12 and 6/96 AND
- b) The lesion size is less than or equal to 12 disc areas in greatest linear dimension AND
- c) There is evidence of recent (< 3 months) presumed disease progression (blood vessel growth, as indicated by fluorescein angiography, optical coherence tomography (OCT), or recent visual acuity changes.

The interval between doses should not be shorter than one month. Administration is to be done by a qualified ophthalmologist experienced in intravitreal injections.

Criteria For Continued Coverage:

Treatment with ranibizumab should be continued only in people who maintain adequate response to therapy.

Ranibizumab should be discontinued if any of the following occur:

- a) Reduction in BCVA in the treated eye to less than 15 letters (absolute) on 2 consecutive visits in the treated eye, attributed to AMD in the absence of other pathology **OR**
- b) 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 **OR**
- c) There is evidence of deterioration of the lesion morphology despite optimum treatment over 3 consecutive visits.

Coverage will not be approved for patients:

- a) Receiving concurrent treatment with verteporfin.
- b) With permanent retinal damage as defined by the Royal College of Ophthalmology guidelines.

Coverage is limited to a maximum of one vial per eye in any 30-day period. Coverage must be renewed every 12 months.

## 2. **Diabetic Macular Edema (DME)**

Initial coverage:

For the treatment of visual impairment due to diabetic macular edema (DME) in patients who meet all of the following criteria:

- clinically significant centre-involving macular edema for whom laser photocoagulation is also indicated
- hemoglobin A1c test in the past 6 months with a value of less than or equal to 11%
- best corrected visual acuity of 20/32 to 20/400
- central retinal thickness greater than or equal to 250 micrometers

Renewal Criteria:

- confirm that a hemoglobin A1C test in the past 6 months had a value of less than or equal to 11%
- date of last visit and results of best corrected visual acuity at that visit
- date of last OCT and central retinal thickness on that examination
- if ranibizumab is being administered monthly, please provide details on the rationale.

Clinical Notes:

- Treatment should be given monthly until maximum visual acuity is achieved (i.e. stable visual acuity for three consecutive months). Thereafter, visual acuity should be monitored monthly.
- Treatment should be resumed when monitoring indicates a loss of visual acuity due to DME and continued until stable visual acuity is reached again for three consecutive months.
- Treatment should be discontinued if there is no improvement of retinal thickness or visual acuity after three consecutive treatments.
- Injection will be by a qualified ophthalmologist with experience in administering intravitreal injections.

Approval Period: 1 year

## 3. **Retinal Vein Occlusion (RVO)**

For the treatment of visual impairment due to macular edema secondary to central retinal vein occlusion (CRVO) or branch retinal vein occlusion (BRVO).

Clinical Notes:

- Treatment should be given monthly until maximum visual acuity is achieved (i.e. stable visual acuity for three consecutive months). Thereafter, visual acuity should be monitored monthly.

- Treatment should be resumed 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 months.
- Treatment should be discontinued if there is no improvement after 6 months of initial treatment.
- Injection will be by a qualified ophthalmologist with experience in administering intravitreal injections.

Approval Period: 1 year

#### 4. **Choroidal Neovascularization**

For the treatment of patients with visual impairment due to choroidal neovascularization secondary to pathologic myopia.

1. Treatment is initiated with a single intravitreal injection. Monitoring is recommended monthly for the first 2 months and at least every 3 months thereafter during the first year.
2. Treatment should be resumed 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.
3. Injection will be by a qualified ophthalmologist with experience in administering intravitreal injections

Approval Period: 1 year

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Rebif** - see Multiple Sclerosis Agents

**Reblozyl** – see Luspatercept

#### **Regorafenib, tablet, 40mg (Stivarga-BAY)**

1. For patients with metastatic and/or unresectable gastrointestinal stromal tumors (GIST) who have had disease progression on, or intolerance to, imatinib and sunitinib; AND has ECOG  $\leq$  1.
2. For the treatment of patients with unresectable hepatocellular carcinoma (HCC) who have experienced disease progression on sorafenib or lenvatinib and meet all of the following criteria:
  - Child-Pugh class status of A.
  - ECOG performance status of 0 or 1.

Clinical Notes:

- Treatment should continue until disease progression or unacceptable toxicity.
- Patients with disease progression on sorafenib must have tolerated a minimum dose of 400 mg per day for at least 20 of the last 28 days of treatment.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Renagel** - see Sevelamer

**Renflexis** – see Infliximab

**Repatha** – see Evolocumab

**Replagal – agalsidase alfa, vial, 1 mg/mL (Takeda)**

Coverage may be available for the treatment of Fabry Disease through the High Cost Drug Plan and Catastrophic Drug Plan, for eligible patients who meet the criteria set out in the Canadian Fabry Disease Treatment Guidelines.

The treatment guidelines are supported by the Canadian Fabry Disease Initiative (CFDI) and may be amended by the CFDI from time to time.

Please contact the PEI Pharmacare Program office at [1-877-577-3737](tel:1-877-577-3737) for more information regarding coverage availability and the Special Authorization application process for this product.

**Retevmo** – see Selpercatinib

**Rexulti** – see Brexpiprazole

**Revatio** - see Sildenafil

**ReVia** - see Naltrexone

**Revlimid** – see Lenalidomide

**Ribociclib, tablet, 200mg (Kisqali-NVR)**

1. In combination with an aromatase inhibitor for the treatment of patients with hormone receptor positive, HER2 negative or advanced or metastatic breast cancer who:

- have not received prior endocrine therapy for advanced or metastatic disease, and
- are not resistant to prior (neo) adjuvant non-steroidal aromatase inhibitor (NSAI) therapy and
- do not have active or uncontrolled metastases to the central nervous system

Renewal criteria:

Confirmation that the patient has responded to treatment and there is no evidence of disease progression.

Clinical Notes:

- Patients must have a good performance status
- Resistance is defined as disease progression occurring during or within 12 months following NSAI therapy
- Treatment should be discontinued upon disease progression or unacceptable toxicity

Claim Notes:

- Initial approval period: 1 year
- Renewal approval period: 1 year

2. In combination with fulvestrant for the treatment of patients with hormone receptor (HR) positive, HER2 negative advanced or metastatic breast cancer, as initial endocrine-based therapy or following disease progression on endocrine therapy. Patients may have also received up to one prior line of chemotherapy for advanced disease. Patients should have a good performance status, without active or uncontrolled metastases to the central nervous system and can be of any menopausal status (perimenopausal and premenopausal women must be treated with an LHRH agonist).

Clinical Notes:

- Treatment should continue until unacceptable toxicity or disease progression.
- Patients who progress  $\leq$  12 months from (neo) adjuvant therapy are eligible for treatment with ribociclib plus fulvestrant.
- Patients who experience disease progression on prior CDK 4/6 inhibitor therapy, fulvestrant or everolimus are not eligible for treatment with ribociclib with fulvestrant.
- Patients currently receiving fulvestrant monotherapy, and who have not progressed may have ribociclib added, provided they are CDK 4/6 inhibitor naïve and otherwise meet funding criteria.
- Patients who previously received everolimus plus exemestane will be eligible for funding of ribociclib plus fulvestrant on progression, provided that treatment was started prior to funding of CDK 4/6 + fulvestrant, patient must be CDK 4/6 naïve and otherwise meet funding criteria.

Claim Notes:



- Initial approval period: 1 year
- Renewal approval period: 1 year

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Rifaximin, tablet, 550mg (Zaxine-LUP)**

For reducing the risk of overt hepatic encephalopathy (HE) recurrence (i.e., 2 or more episodes), if the following clinical criteria are met:

Clinical Criteria:

- Patients are unable to achieve adequate control of HE recurrence with maximal tolerated dose of lactulose alone.
- Must be used in combination with maximal tolerated doses of lactulose.
- For patients not maintained on lactulose, information is required regarding the nature of the patient's intolerance to lactulose.

**Rinvoq** – see Upadacitinib

**Riociguat, tablet, 0.5mg, 1mg, 1.5mg, 2mg, 2.5mg (Adempas-BAY and generic)**

For the treatment of inoperable chronic thromboembolic pulmonary hypertension (CTEPH, World Health Organization [WHO] Group 4) or persistent or recurrent CTEPH after surgical treatment in adult patients (>18 years of age) with WHO Functional Class (FC) II or III pulmonary hypertension (PH).

Should be prescribed by a clinician with experience in the diagnosis and treatment of CTEPH.

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ripretinib, tablet, 50mg (Qinlock-MDP)**

For the treatment of adult patients with advanced gastrointestinal stromal tumours (GIST) who have progression on or intolerance to imatinib, sunitinib, and regorafenib.

Clinical Notes:

- Patients must have a good performance status and no active central nervous system metastases.

- Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Risankizumab, prefilled syringe, 75mg/0.83mL, 150mg/mL; prefilled pen, 150mg/mL; vial, 600mg/10mL; prefilled cartridge, 360mg/2.4mL (Skyrizi-ABV)**

### **Crohn's Disease**

**(600mg/10mL vial and 360mg/2.4mL prefilled cartridge)**

For the treatment of patients with moderate to severe Crohn's disease who have active disease and are refractory, intolerant or have contraindications to:

- Prednisone 40mg (or equivalent) daily for ≥ 2 weeks, AND
- Azathioprine ≥ 2 mg/kg/day for ≥ 3 months, OR
- Mercaptopurine ≥ 1 mg/kg/day for ≥ 3 months, OR
- Methotrexate (SC or IM) ≥ 15 mg/week for ≥ 3 months

#### Clinical Notes:

- Refractory is defined as lack or loss of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.
- Consideration will be given for the approval of a biologic DMARD (disease modifying antirheumatic drug) without a trial of a traditional DMARD for patients who have an aggressive/severe disease course (e.g. extensive disease, a modified Harvey Bradshaw Index score > 16) and are refractory, intolerant or have contraindications to systemic corticosteroids.

#### Claim Notes:

- Initial approval is for 600mg administered by IV infusion at week 0, 4 and 8. Renewal of coverage will require reassessment of the patient and submission of a new Crohn's Disease Special Authorization form. Continued coverage will be approved at a dose of 360mg administered by subcutaneous infusion at week 12, and every 8 weeks thereafter.
- Initial Approval: 16 weeks
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a gastroenterologist using the Crohn's**

Disease Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

### **Plaque Psoriasis**

**(75mg/0.83mL prefilled syringe and 150mg/mL prefilled syringe/pen)**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals.
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20$ mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5-point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet, or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 150mg administered at week 0, week 4, and every 12 weeks thereafter. If response criteria is met at 16 weeks, approval will be continued to a maximum dose of 150mg every 12 weeks up to one year.
- Initial approval: 16 weeks.
- Renewal approval: 1 year.
- Concurrent use of biologics not approved.

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Risdiplam, powder for oral solution, 0.75mg/mL (Evrysdi-HLR)**

For patients diagnosed with 5q Spinal Muscular Atrophy (SMA) under the care of a specialist with experience in the diagnosis and management of SMA, if the following clinical criteria are met:

- Genetic documentation of 5q SMA homozygous gene deletion or compound heterozygote, AND
- Patients who:
  - are symptomatic and have genetic documentation of two or three copies of the SMN2 gene, AND
  - aged between 2 months and 7 months (inclusive), OR
  - aged 8 months up to 25 years and are non-ambulatory
- Patient is not currently requiring permanent invasive ventilation\*, AND
- A baseline assessment using an age-appropriate scale (the Hammersmith Infant Neurological Examination [HINE] Section 2, Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders [CHOP INTEND], or Hammersmith Functional Motor Scale-Expanded [HFMSE]) must be completed prior to initiation of risdiplam treatment.
- For continued coverage, the patient must meet the following criteria:  
There is demonstrated achievement or maintenance of motor milestone function (as assessed using age-appropriate scales: the [HINE] Section 2, CHOP INTEND, or HFMSE) after treatment initiation in patients aged between 2 months and 2 years at the time of treatment initiation; OR
  - There is demonstrated maintenance of motor milestone function (as assessed using age-appropriate scales: the HINE Section 2, CHOP INTEND, or HFMSE) after treatment initiation in patients aged between 2 years and 25 years at the time of treatment initiation; AND
  - Patient does not require permanent invasive ventilation\*.

The decision to discontinue reimbursement should be based on 2 assessments separated by no longer than a 12-week interval.

#### **Claim Notes:**

- Approval: 12 months

\* Permanent invasive ventilation is defined as the use of tracheostomy and a ventilator due to progression of SMA that is not due to an identifiable and reversible cause.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Rituximab, vial, 10mg/mL (Riabni-AMG, Riximyo-SDZ, Ruxience-PFI, Truxima-TEV)**

For Rituximab-naive adult patients whose rituximab therapy is initiated after August 30, 2021, a rituximab biosimilar will be the product approved.

For the treatment of patients with:

1. Rheumatoid arthritis who have a severe intolerance or other contraindication to an anti-TNF agent or failed an adequate trial of an anti-TNF agent.
2. Vasculitis who have a severe intolerance or other contraindication to cyclophosphamide or failed an adequate trial of cyclophosphamide.
3. Other autoimmune diseases whom have failed previous treatments.

Clinical Note: A detailed description of previously failed treatments must be provided.

Claim Notes:

1. Must be prescribed by a specialist.
2. Initial approval period: 6 months. Confirmation of response is required.
3. Claims that exceed the maximum claim amount of \$9,999.99 must be divided and submitted as separate transactions using the DIN first, and then PDINs.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Risedronate, tablet, 30mg (Generic)**

For the treatment of Paget's disease of the bone for a maximum 2 month period. One additional 2 month course of treatment may be considered after a drug holiday of at least 60 days.

**Risperdal Consta** - see Risperidone prolonged release injection

**Risperidone, prolonged release injection, 12.5mg/2mL, 25mg/2mL, 37.5mg/2mL, 50mg/2mL (Risperdal Consta-JAN)**

Note: For Community Mental Health Drug Program, no Special Authorization is required. For the treatment of schizophrenia or schizoaffective disorder in patients who have:

a) A history of non-adherence.

**OR**

b) Inadequate control or significant side-effects from two or more oral antipsychotic medications.

**OR**

- c) Inadequate control or significant side-effects from at least one long-acting depot antipsychotic agent.

**NOTE:**

- **Must be requested and prescribed by a psychiatrist.**
- **Only doses up to 50mg every two weeks will be approved.**

**Rivastigmine** - see Cholinesterase Inhibitors (ChEI)

**Riximyo** – see Rituximab

**Rotigotine, transdermal patch, 2mg, 4mg, 6mg, 8mg (Neupro-UCB)**

For the treatment of the signs and symptoms of Parkinson's Disease in patients who are experiencing motor fluctuations despite optimal treatment with Levodopa/Carboxylase therapy upon written request or recommendation of a neurologist. A copy of the recommendation must accompany the Special Authorization.

**Rozlytrek** – see Entrectinib

**Rufinamide, tablet, 100mg, 200mg, 400mg (Banzel-EIS and generic)**

For the adjunctive treatment of seizures associated with Lennox-Gastaut syndrome for patients who meet all of the following criteria:

- are under the care of a physician experienced in treating Lennox-Gastaut syndrome-associated seizures, AND
- are currently receiving two or more antiepileptic drugs, AND
- in whom less costly antiepileptic drugs are ineffective or not appropriate.

**Ruxience** – see Rituximab

**Ruxolitinib, tablet, 5mg, 10mg, 15mg, 20mg (Jakavi-NVR)**

**Acute Graft-Versus-Host Disease**

For the treatment of steroid-refractory or steroid-dependent acute graft-versus-host disease (aGvHD) in adult and pediatric patients aged 12 years and older who meet all the following criteria:

- Clinically diagnosed grade II to IV aGvHD according to the NIH criteria (Harris et al. [2016]).
- Confirmed diagnosis of corticosteroid-refractory or corticosteroid-dependent aGvHD.

Renewal criteria:

- Achieved an overall response (i.e., CR, VGPR, PR, or stable disease with significant reduction in steroid doses), according to standard NIH criteria at day 28.

- For subsequent renewals, patients should be assessed for treatment response every 2 to 3 months, until the occurrence of any of the discontinuation criteria listed below.

Clinical Notes:

- Treatment should be discontinued upon the occurrence of any of the following:
  - progression of aGvHD, defined as worsening of aGvHD symptoms or occurrence of new aGvHD symptoms
  - unacceptable toxicity
  - addition of systemic therapies (other than calcineurin inhibitors) for aGvHD after day 28
  - recurrence or relapse of underlying hematological malignancy.

Claim Notes:

- Must be prescribed by clinicians who have experience in the diagnosis and management of patients with aGvHD.
- Must not be added to patients' concurrent treatment of systemic therapies for the treatment of aGvHD other than steroids with or without calcineurin inhibitors.
- Approval: 6 months

**Chronic Graft-Versus-Host Disease**

For the treatment of chronic graft-versus-host disease (cGvHD) in adults and pediatric patients aged 12 years and older who have inadequate response to corticosteroids or other systemic therapies who meet all the following criteria:

- Clinically diagnosed cGvHD staging of moderate to severe based on NIH consensus criteria
- Confirmed diagnosis cGvHD with inadequate response to corticosteroids or other systemic therapies

Renewal criteria:

- Achieved an overall response (i.e., CR or PR, or stable disease with significant reduction in steroid doses), according to NIH criteria, after 24 weeks of therapy.

Clinical Notes:

- Treatment should be discontinued upon the occurrence of any of the following:
  - Progression of cGvHD, defined as worsening of cGvHD symptoms or occurrence of new cGvHD symptoms
  - recurrence or relapse of underlying hematological malignancy

Claim Notes:

- Must be prescribed by clinicians who have experience in the diagnosis and management of patients with cGvHD.
- Must not be added to patients' concurrent treatment of systemic therapies other than steroids with or without calcineurin inhibitors.

- Initial Approval: 6 months

### **Myelofibrosis**

For patients with intermediate to high risk symptomatic Myelofibrosis (MF) as assessed using the Dynamic International Prognostic Scoring System (DIPSS) Plus or patients with symptomatic splenomegaly. Patients should have ECOG performance status of  $\leq 3$  and be either previously untreated or refractory to other treatment.

### **Polycythemia Vera**

For the treatment of patients with polycythemia vera who have demonstrated resistance or intolerance to hydroxyurea (HU).

#### Renewal Criteria:

- Written confirmation that the patient has responded to treatment and there is no evidence of disease progression.

#### Clinical Notes:

- Patients must have a good performance status.
- Treatment should be discontinued upon disease progression or unacceptable toxicity.
- Resistance is considered if, after at least 3 months of HU therapy at the maximum tolerated dose, patients experience at least one of the following:
  - Need for phlebotomy to maintain hematocrit (HCT)  $< 45\%$
  - Uncontrolled myeloproliferation (i.e., platelet count  $> 400 \times 10^9/L$  and white blood cell count  $> 10 \times 10^9/L$ )
  - Failure to reduce massive splenomegaly by greater than 50%, as measured by palpation
- Intolerance to HU is considered if patients experience at least one of the following:
  - Absolute neutrophil count  $< 1.0 \times 10^9/L$ , platelet count  $< 100 \times 10^9/L$  or hemoglobin  $< 100g/L$  at the lowest dose of HU required to achieve a response (a response to HU is defined as HCT  $< 45\%$  without phlebotomy, and/or all of the following: platelet count  $< 400 \times 10^9/L$ , white blood cell count  $< 10 \times 10^9/L$ , and nonpalpable spleen).
  - Presence of leg ulcers or other unacceptable HU-related non-hematological toxicities (defined as grade 3 or 4 or, more than one week of grade 2) such as mucocutaneous manifestations, gastrointestinal symptoms, pneumonitis, or fever.
  - Toxicity requiring permanent discontinuation of HU, interruption of HU until toxicity resolved, or hospitalization due to HU toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**



**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ruzurgi** – see Amifampridine

**Rybelsus** – see Semaglutide

**Rydapt** – see Midostaurin

**Rymti** – see Etanercept

**Sabril** – see Vigabatrin

**Sacubitril & Valsartan, tablet, 24mg & 26mg, 49mg & 51mg, 97mg & 103mg (Entresto-NVR)**

For the treatment of patients with New York Heart Association (NYHA) class II or III heart failure to reduce the incidence of cardiovascular death and heart failure hospitalization, who meet all of the following criteria:

- Left ventricular ejection fraction (LVEF) of <40%
- NYHA class II or III symptoms despite at least four weeks of treatment of the following:
  - a stable dose of an angiotensin-converting enzyme inhibitor (ACEI) or an angiotensin II receptor antagonist (ARB); and
  - a stable dose of a beta-blocker and other recommended therapies, including an aldosterone antagonist.
- Plasma B-type natriuretic peptide (BNP)  $\geq$  150pg/mL or N-terminal prohormone B-type natriuretic peptide (NTproBNP)  $\geq$  600 pg/mL.

Clinical Notes:

1. A plasma BNP  $\geq$  100 pg/mL or NT-proBNP  $\geq$  400 pg/mL will be considered if the patient has been hospitalized for heart failure within the past 12 months.
2. For patients who have not received four weeks of therapy with a beta blocker or aldosterone antagonist due to an intolerance or contraindication, details must be provided.

**Salagen** - see Pilocarpine

**Salmeterol Xinafoate, aerosol powder disk, 50µg/dose (Serevent Diskus-GSK)**

- a) For the treatment of asthma when used in patients on concurrent steroid therapy.
- b) For the treatment of COPD, see Chronic Obstructive Pulmonary Disease.

**Note: Patients using this product must also have access to a short-acting beta-2 agonist bronchodilator for the relief of acute symptoms.**

**Salmeterol & Fluticasone, aerosol inhalation, 25mcg & 125mcg per dose, 25mcg & 250mcg per dose (Advair-GSK); inhaled powder disk, 50mcg & 100mcg per dose, 50mcg & 250mcg per dose, 50mcg & 500mcg per dose (Advair Diskus- GSK and generics)**

- a) For the treatment of asthma in patients who are not well controlled on a regular and adequate course of inhaled steroid therapy prior to the request for combination therapy.
- b) For the treatment of COPD, see Chronic Obstructive Pulmonary Disease.

**Note: Patients using this product must also have access to a short-acting beta-2 agonist bronchodilator for the relief of acute symptoms.**

**Saphnelo** – see Anifrolumab

**Sapropterin, powder for oral solution, 100 mg, 500 mg (Reddy-Sapropterin-RCH)**

For the ongoing treatment of hyperphenylalaninemia (HPA) due to tetrahydrobiopterin (BH4)-responsive phenylketonuria (PKU) in conjunction with a phenylalanine (Phe)-restricted diet in patients who meet all of the following criteria:

- Confirmed diagnosis based on genetic testing.
- Response to sapropterin provided by the manufacturers initial 6 month trial through the Patient Support Program (PSP) 'Reddy-Sapropterin Support Program for Patients with HPA due to PKU'.
- Baseline blood Phe levels greater than 360 umol/L despite compliance with a low protein diet and formulas (nonpregnant patients require at least 2 baseline levels and pregnant patients require at least 1 baseline level during a 3 to 6 month time frame).
- Achievement of the following during a 6-month trial of treatment:
  - For pregnant or non-pregnant patients, normal sustained blood Phe levels of 120 umol/L to 360 umol/L; or
  - For non-pregnant patients, sustained blood Phe reduction of at least 30% compared to baseline if the baseline blood Phe level is less than 1200 umol/L; or
  - For non-pregnant patients, sustained blood Phe reduction of at least 50% compared to baseline if the baseline blood Phe level is greater than 1200 umol/L.
  - For non-pregnant patients, documented increase in dietary protein tolerance based on targets set between the clinician and patient.

Renewal Criteria:

- Confirmation of continued response to sapropterin based on Phe levels achieved during the 6-month trial. Two Phe levels taken at least 1 month apart must be provided.

Clinical Notes:

1. Patients must be initiated on treatment and followed in a specialized clinic with expertise in the diagnosis and management of PKU.
2. Phe blood levels and Phe tolerance levels must be provided.
3. Pregnant patients who have maintained a decrease in Phe levels below 360 umol/L during the 6-month trial period will be eligible for coverage of sapropterin for the duration of the pregnancy.
4. Confirmation of compliance with a low protein diet is required before initiation and in conjunction with ongoing use.

Claim Notes:

- Approvals will be for a maximum of 20 mg/kg per day.
- Renewals for sapropterin in pregnant patients will not be considered.
- Approval period: 1 year

\* Claims that exceed the maximum claim amount of \$9999.99 must be divided as separate transactions, using DIN first and PDIN\* second.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Sarilumab, syringe, 150mg/1.14mL, 200mg/1.14mL; pen, 150mg/1.14mL, 200mg/1.14mL (Kevzara-AVN)**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks AND Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks;

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.

- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Approval for adults is 200 mg once every 2 weeks given as a subcutaneous injection. Reduction of dose to 150 mg once every 2 weeks is recommended for management of neutropenia, thrombocytopenia, and elevated liver enzymes.
- Must be prescribed by a rheumatologist.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Satralizumab, prefilled syringe, 120mg/mL (Enspryng-HLR)**

For the treatment of patients 12 years of age and older with neuromyelitis optica spectrum disorder (NMOSD) who meet all of the following criteria:

- Are anti-aquaporin4 (AQP4) seropositive.
- Must have had at least one relapse of NMOSD in the previous 12 months:
  - despite an adequate trial of other accessible preventive treatments<sup>1</sup> for NMOSD, OR
  - because the patient cannot tolerate other preventive treatments<sup>1</sup> for NMOSD
- Patients must have an EDSS score of 6.5 points or less.
- Satralizumab should not be initiated during a NMOSD relapse episode.

Renewal:

- Requests for renewal will be considered for patients who maintain an EDSS score of less than 8 points.

Clinical Note:

- Must be prescribed by a neurologist with expertise in treating NMOSD.

Claim Notes:

1. Combined use of more than one biologic drug will not be reimbursed.
2. Approvals will be for a maximum of 120mg at week 0, 2 and 4, then 120 mg every

four weeks thereafter.

<sup>1</sup>Other accessible preventative treatments include, but are not limited to, monoclonal antibodies and other immunosuppressants.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Saxagliptin, tablet, 2.5mg, 5mg (Onglyza-AZE and generics)**

For the treatment of type 2 diabetes as a third drug added on to metformin and a sulfonylurea for patients with inadequate glycemic control on optimal doses of metformin and a sulfonylurea, **and** for whom insulin is not an option.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Saxagliptin & Metformin Hydrochloride, 2.5mg/500mg, 2.5mg/850mg/2.5mg/1000mg (Komboglyze-AZE)**

For patients with type 2 diabetes for whom insulin is not an option and who are already stabilized on therapy with metformin, a sulfonylurea and saxagliptin, to replace the individual components of saxagliptin and metformin in these patients.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Scemblix** – see Asciminib

**Secukinumab, syringe or pen, 150mg/mL (Cosentyx-NVR)**

**Ankylosing Spondylitis**

For the treatment of patients with moderate to severe ankylosing spondylitis (Bath AS Disease Activity Index (BASDAI) score  $\geq 4$  on 10 point scale) who:

- a) have axial symptoms\* and who have failed to respond to the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation or in whom NSAIDs are contraindicated OR
- b) have peripheral symptoms and who have failed to respond to, or have contraindications to, the sequential use of at least 2 NSAIDs at the optimum dose for a minimum period of 3 months observation and have had an inadequate response to an optimal dose or maximal tolerated dose of a DMARD.

\*Patients with recurrent uveitis (2 or more episodes within 12 months) as a complication to

axial disease, do not require a trial of NSAIDs alone.

Claim Notes:

- Approvals will be for a maximum adult dose of 150 mg at 0, 1, 2, and 3 weeks followed by monthly maintenance dosing of 150 mg starting at week 4.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial Approval: 6 months
- Renewal Approval: maximum of 12 months.

Requests for renewal must include information showing the beneficial effects of the treatment, specifically:

- a) a decrease of at least two points on the BASDAI scale, compared with pretreatment score OR
- b) patient and expert opinion of an adequate clinical response as indicated by a significant functional improvement (measured by outcomes such as Health Assessment Questionnaire (HAQ) or ability to return to work).

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Ankylosing Spondylitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Plaque Psoriasis**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score ; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 300 mg at 0, 1, 2, and 3 weeks followed by monthly maintenance dosing starting at week 4, up to 12 weeks. If response criteria is met at 12 weeks, approval will be continued to a maximum dose of 300 mg.
- Initial approval: 12 weeks.
- Renewal approval: 1 year.
- Concurrent use of biologics not approved.

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Psoriatic Arthritis**

- 1) For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; and
  - Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and
  - Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

Clinical notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.

- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of 150mg at weeks 0, 1, 2 and 3 followed by monthly maintenance dosing of 150mg starting at week 4. For patients who are anti-TNF $\alpha$  inadequate responders and continue to have active psoriatic arthritis, consider using the 300 mg dose. For psoriatic arthritis patients with coexistent moderate to severe plaque psoriasis, use the dosing and administration recommendations for plaque psoriasis (i.e. 300 mg at weeks 0, 1, 2, and 3, followed by monthly maintenance dosing starting at week 4).
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Seebri Breezhaler** – see Glycopyrronium Bromide

**Selexipag, tablet, 200mcg, 400mcg, 600mcg, 800mcg, 1000mcg, 1200mcg, 1400mcg, 1600mcg (Uptravi-ACT)**

For the long-term treatment of idiopathic pulmonary arterial hypertension (PAH), heritable PAH, PAH associated with connective tissue disorders, and PAH associated with congenital heart disease, in adult patients with World Health Organization (WHO) functional class (FC) II to III to delay disease progression, if the following clinical criterion and conditions are met:

- Inadequate control with a first and second-line PAH therapy
- Prescribed by a clinician with experience in the diagnosis and treatment of PAH

**NOTE:**

Combination therapy with prostacyclin or prostacyclin analogs therapies will not be covered

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Selinexor, tablet, 20mg (Xpovio-FTI)**

In combination with bortezomib and dexamethasone for the treatment of adult patients with multiple myeloma and who have received at least one prior therapy.



Clinical Notes:

1. Prior treatment with bortezomib/proteasome inhibitor is permitted if all the following criteria are met:
  - Best response achieved with bortezomib/proteasome inhibitor was at least a partial response.
  - Bortezomib/proteasome inhibitor not discontinued for grade 3 or higher toxicity
  - Bortezomib/proteasome inhibitor treatment-free interval has been at least six months.
2. Treatment should continue until disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Selpercatinib, capsule, 40mg, 80mg (Retevmo-LIL)**

**Medullary Thyroid Cancer**

For the treatment of patients 12 years and older with unresectable locally advanced or metastatic RET-mutant medullary thyroid cancer (MTC) who have progressed on, are intolerant to, or have a contraindication to first-line therapy.

Clinical Notes:

- Discontinuation for unacceptable toxicity or loss of clinical benefit.
- Patients should have a good performance status.
- Monotherapy only.
- Confirm RET mutation prior to initiating therapy.
- Patients with prior progression on a RET inhibitor are ineligible.

**Differentiated Thyroid Carcinoma (DTC)**

For the treatment of adult patients with locally advanced or metastatic RET fusion-positive differentiated thyroid carcinoma (DTC) not amenable to surgery or radioactive iodine therapy, following prior treatment with lenvatinib.

Clinical Notes:

- Discontinuation for unacceptable toxicity or loss of clinical benefit.
- Patients should have a good performance status.
- Monotherapy only.
- Confirm RET mutation prior to initiating therapy.
- Patients with prior progression on a RET inhibitor are ineligible.

### **Non-Small Cell Lung Cancer**

For the treatment of adult patients with metastatic RET fusion-positive non-small cell lung cancer (NSCLC) as first-line treatment or after prior systemic therapy.

#### Clinical Notes:

- Discontinuation for unacceptable toxicity or loss of clinical benefit.
- Patients should have a good performance status.
- Monotherapy only.
- Confirm RET mutation prior to initiating therapy.
- Patients with prior progression on a RET inhibitor are ineligible.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Semaglutide, tablet, 3mg, 7mg, 14mg (Rybelsus-NNO)**

For the treatment of type 2 diabetes in combination with metformin and a sulfonylurea, when diet and exercise plus dual therapy with metformin and a sulfonylurea do not achieve adequate glycemic control.

**The request for coverage must be made using the Semaglutide Special Authorization Request Form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

### **Semaglutide, pen injector, 0.25-0.5mg, 1mg (Ozempic-NNO)**

For the treatment of type 2 diabetes in combination with metformin and a sulfonylurea, when diet and exercise plus dual therapy with metformin and a sulfonylurea do not achieve adequate glycemic control.

#### Claim Note:

- Approvals will be for a maximum of 1 prefilled pen every 4 weeks.

**The request for coverage must be made using the Semaglutide Special Authorization Request Form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Serc** - see Betahistine

**Serevent** - see Salmeterol

**Serevent Diskus** - see Salmeterol

**Sevelamer carbonate, tablet, 800mg (Accel-Sevelamer)**

For the treatment of hyperphosphatemia (>1.8 mmol/L) in patients with end stage renal disease (eGFR<15mL/min) who have:

- Inadequate control of phosphate levels on a calcium based phosphate binder, or
- Hypercalcemia (corrected for albumin), or
- Calciphylaxis (calcific arteriolopathy)

**NOTE**

Initial approval for 6 months, renewed at 1 year intervals with demonstration of clinically meaningful improvement of phosphate levels (lab values must be provided).

**Sevelamer hcl, tablet, 800mg (Renagel-GZY)**

For the treatment of hyperphosphatemia (>1.8 mmol/L) in patients with end stage renal disease (eGFR<15mL/min) who have:

- Inadequate control of phosphate levels on a calcium based phosphate binder, or
- Hypercalcemia (corrected for albumin), or
- Calciphylaxis (calcific arteriolopathy)

**NOTE**

Initial approval for 6 months, renewed at 1 year intervals with demonstration of clinically meaningful improvement of phosphate levels (lab values must be provided).

**Sildenafil, tablet, 20mg (Revatio-PFI and generics)**

For the treatment of patients with World Health Organization (WHO) functional class III idiopathic pulmonary arterial hypertension (IPAH) who do not demonstrate vasoreactivity on testing or who do demonstrate vasoreactivity on testing but fail a trial of calcium channel blockers.

For the treatment of patients with World Health Organization (WHO) functional class III pulmonary arterial hypertension (PAH) associated with connective tissue diseases who do not respond to conventional therapy.

Diagnosis of PAH should be confirmed by cardiac catheterization.

Coverage must be requested and medications prescribed by a Pulmonary Arterial Hypertension (PAH) specialist.

Claim Note:

The maximum dose of sildenafil that will be reimbursed is 20mg three times daily.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Siliq** – see Brodalumab

**Simlandi** - see Adalimumab

**Simponi** – see Golimumab

**Siponimod, tablet, 0.25mg, 2mg (Mayzent-NVR)**

Initiation Criteria:

For the treatment of patients with active secondary progressive multiple sclerosis, who meet all the following criteria:

- a history of relapsing-remitting multiple sclerosis (RRMS)
- an Expanded Disability Status Scale (EDSS) score of 3.0 to 6.5
- documented EDSS progression during the two years prior to initiating treatment with siponimod ( $\geq 1$  point if EDSS < 6.0;  $\geq 0.5$  points if EDSS  $\geq 6.0$  at screening).

Renewal Criteria:

- Patients who do NOT exhibit evidence of disease progression since the previous assessment. Disease progression is defined as:
  - an increase in the EDSS score of greater than or equal to 1 point if the EDSS score was 3.0 to 5.0 at siponimod initiation
  - OR
  - an increase of greater than or equal to 0.5 points if the EDSS score was 5.5 to 6.5 at siponimod initiation

Patients who do NOT exhibit one of the following:

- progression to an EDSS score of equal to or greater than 7.0 at any time during siponimod treatment
- confirmed worsening of at least 20% on the timed 25-foot walk (T25W) since initiating siponimod treatment

Clinical Notes:

- Patients should be assessed for a response to siponimod every six months.

Claim Notes:

- The patient is under the care of a neurologist with experience in the diagnosis and management of multiple sclerosis.
- Siponimod should not be used in combination with other disease-modifying treatments (DMTs) used to treat multiple sclerosis.
- Approval period: 1 year

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Sitagliptin, tablet, 25mg, 50mg, 100mg (Januvia-MSD and generics)**

For the treatment of type 2 diabetes as a third drug added on to metformin and a sulfonyleurea for patients with inadequate glycemic control on optimal doses of metformin and a sulfonyleurea, **and** for whom insulin is not an option.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Sitagliptin & Metformin Hydrochloride, tablet, 50mg/500mg, 50mg/850mg, 50mg/1000mg (Janumet-MSD and generics)**

For the treatment of type 2 diabetes for patients for whom insulin is not an option and who are already stabilized on therapy with metformin, a sulfonyleurea and sitagliptin, to replace the individual components of sitagliptin and metformin.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Sitagliptin & Metformin Hydrochloride, extended release tablet, 50mg/500mg, 50mg/1000mg, 100mg/1000mg (Janumet XR-MSD and generics)**

For the treatment of type 2 diabetes for patients for whom insulin is not an option and who are already stabilized on therapy with metformin, a sulfonyleurea and sitagliptin, to replace the individual components of sitagliptin and metformin.

**The request for coverage must be made using the DPP-4/SGLT2 Inhibitors Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms> .**

**Skyrizi** – see Risankizumab

**Sodium Cromoglycate, capsule, 100mg (Nalcrom-AVN)**

For the treatment of patients who experience severe reactions to foods which cannot be avoided.

**Somatrogon, prefilled pen, 24 mg/1.2 ml, 60 mg/1.2 ml (Ngenla-PFI)**

For the long-term treatment of pediatric patients who have growth failure due to an inadequate secretion of endogenous growth hormone (growth hormone deficiency [GHD]) only if the following conditions are met:

**Initiation Criteria:**

Pre-pubertal children who are at least 3 years of age, and who are diagnosed with either isolated GHD, or growth hormone insufficiency as part of multiple pituitary hormone deficiency.

**Discontinuation Criteria:**

Treatment with somatrogen must be discontinued upon the occurrence of any of the following:

1. Height velocity is less than 2 cm per year and bone age is more than 16 years in boys and 14 years in girls
2. Closure of the epiphyseal growth plates

**Clinical Notes:**

1. Patient height and weight must be provided with all requests.
2. Confirmation there is no evidence of epiphyseal growth plate closure and a copy of the bone age report must be provided with all requests.
3. Bone age assessments may be based on the Greulich Pyle Atlas, Tanner-Whitehouse, or other appropriate methods of assessment.

**Claim Notes:**

- Must be prescribed by, or in consultation with, an endocrinologist.
- Approvals will be for a maximum of 0.66 mg/kg weekly.
- Approval period: 1 year

**Sorafenib, tablet, 200mg (Nexavar-BAY)**

- For use as a single agent second line treatment in patients with documented evidence of histologically confirmed advanced or metastatic clear cell renal cell carcinoma, considered to be intermediate or low risk (according to Memorial Sloan-Kettering (MSKCC) prognostic score, see below), have an ECOG performance status of 0 or 1 and progressed after prior cytokine therapy (or intolerance) within the previous 8 months. In any one patient all of the following conditions must be met:
- Sorafenib may be a second line option only after cytokine therapy.
- Sorafenib may not be used after another tyrosine kinase inhibitor (i.e.Sunitinib) as sequential therapy.  
In the event of severe toxicity within the first 8 weeks of therapy, a switch to another tyrosine kinase inhibitor (i.e.Sunitinib) may be allowed.
- For use in patients with Child-Pugh Class A advanced hepatocellular carcinoma, who have progressed on trans-arterial chemoembolization (TACE) or are not suitable for the TACE procedure and have an ECOG performance status of 0 to 2. Renewal of coverage requires no further progression of the patient's disease as evidenced by radiological or scan

results. Copies of the results must accompany the Special Authorization.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Spinraza** – see Nusinersen

**Spiriva** - see Tiotropium

**Spiriva Respimat** – see Tiotropium

**Sporanox** - see Itraconazole

**Sprycel** - see Dasatinib

**Stalevo** – see Carbidopa & Levodopa & Entacapone

**Stelara** – see Ustekinumab

**Stiripentol, capsules, powder for inhalation, 250mg, 500mg (Diacomit-BIO)**

For use in combination with clobazam and valproate as adjunctive therapy of refractory generalized tonic-clonic seizures in patients with severe myoclonic epilepsy in infancy (Dravet syndrome), whose seizures are not adequately controlled with clobazam and valproate alone.

The patient must be under the care of a neurologist or a pediatrician.

**Patients must also apply for coverage through the High Cost Drug Program. The patient application is available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Stivarga** – see Regorafenib

**Sucroferric oxyhydroxide, chewable tablet, 500mg (Velphoro-VFM)**

For the treatment of hyperphosphatemia (>1.8 mmol/L) in patients with end stage renal disease (eGFR<15mL/min) who have:

- Inadequate control of phosphate levels on a calcium based phosphate binder, or
- Hypercalcemia (corrected for albumin), or
- Calciphylaxis (calcific arteriopathy)

Clinical Notes:

- Initial approval for 6 months, renewed at 1 year intervals with demonstration of clinically meaningful improvement of phosphate levels (lab values must be provided).

**Sumatriptan, nasal spray, 5mg, 20mg; injection 6mg/0.5mL (Imitrex DF-GSK and generics)**

For the treatment of migraine headaches where other standard therapies, such as oral analgesics have failed **AND** the patient has not responded to Zolmitriptan or Rizatriptan.

**Coverage for the injectable form will only be considered if the tablet and nasal dosage forms are not appropriate.**

**Coverage is limited to 6 sprays or 6 syringes per 30 day period.** Anyone requiring more than 6 doses per 30 day period should be considered for migraine prophylaxis therapy if they are not already receiving such therapy.

**Sunitinib, capsule, 12.5mg, 25mg, 50mg (Sutent-PFI and generic)**

- a) For use as a single agent first line treatment in patients with documented evidence of histologically confirmed advanced or metastatic clear cell renal cell carcinoma who have an ECOG performance status of 0 or 1. In any one patient all of the following conditions must be met:
- Sunitinib may be a first line option.
  - Sunitinib may not be used after another tyrosine kinase inhibitor (i.e. Sorafenib) as sequential therapy.

In the event of severe toxicity within the first 8 weeks of therapy, a switch to another tyrosine kinase inhibitor (i.e.Sorafenib) may be allowed.

- b) For use as a single agent for the treatment of advanced gastrointestinal stromal tumor (GIST) patients after failure of Imatinib due to intolerance or resistance.
- c) For the treatment of patients with progressive, unresectable, locally advanced or metastatic, well or moderately differentiated pancreatic neuroendocrine tumours.

**Clinical Notes:**

1. Patients must have a good performance status.
2. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**



**Sutent** - see Sunitinib

**Symbicort Turbuhaler** - see Formoterol & Budesonide

**Synjardy** – see Empagliflozin & Metformin

**Tacrolimus, topical ointment, 0.1% (Protopic-AST)**

For intermittent use in adults with moderate to severe atopic dermatitis who have failed or are intolerant to a site appropriate strength of corticosteroid therapy (i.e. low potency on face versus intermediate to high potency for trunk and extremities).

**Tacrolimus, topical ointment, 0.03% (Protopic-AST)**

For use in children greater than 2 years of age with refractory atopic dermatitis for a period of up to 12 months.

**Tafamidis, tablet, 61mg (Vyndamax-PFI)**

**Tafamidis meglumine, capsule, 20mg (Vyndaqel-PFI)**

For the treatment of cardiomyopathy in adult patients with documented hereditary or wildtype transthyretin-mediated amyloidosis (ATTR) who meet all of the following criteria:

- New York Heart Association (NYHA) class I to III heart failure.
- At least one prior hospitalization for heart failure or clinical evidence of heart failure that required treatment with a diuretic.
- Has not previously undergone a heart or liver transplant.
- Does not have an implanted cardiac mechanical assist device (CMAD).

Discontinuation Criteria:

The patient has:

- NYHA class IV heart failure, or
- received an implanted CMAD, or
- received a heart or liver transplant.

Clinical Notes :

1. Wild-type ATTR-cardiomyopathy (CM) consists of all of the following:
  - absence of a variant transthyretin (TTR) genotype
  - evidence of cardiac involvement by echocardiography with end-diastolic interventricular septal wall thickness greater than 12mm
  - positive findings on technetium-99m pyrophosphate (Tc-99m-PYP) scintigraphy with single-photon emission computerized tomography (SPECT) scanning OR presence of amyloid deposits in biopsy tissue (fat aspirate, salivary gland, median nerve connection tissue sheath, or cardiac tissue); and

TTR precursor protein identification by immunohistochemistry, scintigraphy, or mass spectrometer

2. Hereditary ATTR-CM consists of all of the following:
- presence of a variant TTR genotype associated with CM and presenting with a CM phenotype
  - evidence of cardiac involvement by echocardiography with end-diastolic interventricular septal wall thickness greater than 12mm
  - positive findings on technetium-99m pyrophosphate (Tc-99m-PYP) scintigraphy with single-photon emission computerized tomography (SPECT) scanning OR presence of amyloid deposits in biopsy tissue (fat aspirate, salivary gland, median nerve connective tissue sheath, or cardiac tissue)

Claim Notes:

- The patient must be under the care of a physician with experience in the diagnosis and treatment of ATTR-CM.
- Combination therapy with other interfering ribonucleic acid drugs or transthyretin stabilizers used to treat ATTR-CM will not be reimbursed.
- Initial approval period: 9 months.
- Renewal approval period: 1 year.

**Patients must also apply for coverage by the High Cost Drug Program. The patient application is available from the Drug Program Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Tafinlar** – see Dabrafenib

**Tagrisso** – see Osimertinib

**Takhzyro** – see Lanadelumab

**Taltz** – see Ixekizumab

**Tarceva** – Erlotinib

**Tasigna** – see Nilotinib

**Tazarotene/Halobetasol propionate, lotion, 0.01%/0.045% (Duobrii – BLO)**

Patients must have a clinical diagnosis of moderate to severe plaque psoriasis and an inadequate response to a topical high-potency corticosteroid.

**Tecfidera** – see Multiple Sclerosis Agents

**Tecta** - see Proton Pump Inhibitors

**Temodal** – see Temozolomide

**Temozolomide, capsule, 5mg, 20mg, 100mg, 140mg, 250mg (Temodal–MSD and generics)**

For the treatment of brain tumors (Malignant glioma)

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Testim** – see Testosterone

**Testosterone, transdermal gel, 25mg/2.5gm packet, 50mg/5gm packet (AndroGel-BGP); 50mg/5gm tube (Testim-PAL)**

For the treatment of congenital and acquired primary or secondary hypogonadism in males with a specific diagnosis of;

**Primary** - Cryptorchidism, Klinefelter's, orchiectomy, and other established causes.

**Secondary** - Pituitary-hypothalamic injury due to tumors, trauma, radiation. Testosterone deficiency should be clearly demonstrated by clinical features and confirmed by two separate biochemical tests before initiating any testosterone therapy. Limited to 5 g/day gel.

**Older males with non-specific symptoms of fatigue, malaise or depression who have low testosterone (T) levels do not satisfy these criteria.**

**Testosterone Undecanoate, capsule, 40mg (Generics)**

For patients with a documented deficiency in whom treatment with depo-testosterone products have been unsuccessful, intolerable or are medically contraindicated.

**Tezepelumab, prefilled pen, prefilled syringe, 210mg/1.91mL (Tezspire-AZE)**

For the treatment of severe asthma in patients 12 years and older who are inadequately controlled with high-dose inhaled corticosteroids (ICS), and one or more additional asthma controller(s) (e.g., long-acting beta-agonist), and have experienced 2 or more clinically significant asthma exacerbations in the past 12 months.

Initial Discontinuation Criteria:

- Baseline asthma control questionnaire score has not improved at 12 months since initiation of treatment, or
- No decrease in the daily maintenance OCS dose in the first 12 months of treatment, or

- The number of clinically significant asthma exacerbations has increased within the previous 12 months.

Subsequent Discontinuation Criteria:

- Asthma control questionnaire score achieved after the first 12 months of therapy has not been maintained subsequently, or
- The reduction in the daily maintenance dose of OCS achieved after the first 12 months of treatment is not maintained or improved subsequently, or
- The number of clinically significant asthma exacerbations has increased within the previous 12 months.

Clinical Notes:

- A baseline assessment of asthma symptom control using a validated asthma control questionnaire must be provided.
- A baseline and annual number of clinically significant asthma exacerbations must be provided.
- High dose ICS is defined as  $\geq 500$  mcg of fluticasone propionate or equivalent daily dose.
- A significant clinical exacerbation is defined as worsening of asthma such that the treating physician elected to administer systemic glucocorticoids for at least 3 days or the patient visited an emergency department or was hospitalized.

Claim Notes:

- Must be prescribed by a respirologist, clinical immunologist, allergist or internist experienced in the treatment of severe asthma.
- Combined use of tezepelumab with other biologics used to treat asthma will not be reimbursed.
- Approvals will be for a maximum of 210mg subcutaneous injection every 4 weeks.
- Approval period: 1 year.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Tezspire** – see Tezepelumab

**Tildrakizumab, prefilled syringe, 100mg/mL (Ilumya-SUN)**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of  $>10\%$  and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of  $\geq 20$ mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND

- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score ; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum of 100mg at week 0, week 4, and every 12 weeks thereafter. If response criteria is met at 16 weeks, approvals will be continued to a maximum dose of 100mg every 12 weeks up to one year.
- Initial approval: 16 weeks.
- Renewal approval: 1 year.
- Concurrent use of biologics not approved.

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Thyrogen** - see Thyrotropin

**Thyrotropin, injection, 0.9mg.mL (Thyrogen-GZY)**

For use as a single agent in patients who have documented evidence of thyroid cancer, who have undergone appropriate surgical and/or medical management, and require on-going evaluation to monitor for recurrence and metastatic disease. This includes:

- a) Primary use in patients with inability to raise an endogenous TSH level ( $\geq 25$  mu/L) with thyroid hormone withdrawal.

- b) Primary use in cases of documented morbidity in patients for whom severe hypothyroidism could be life threatening, such as unstable angina, recent myocardial infarction, class III to IV congestive heart failure, or uncontrolled psychiatric illness.
- c) Secondary use in patients with previous thyroid hormone withdrawal resulting in a documented life-threatening event.

**(This criteria is for patients of the Catastrophic Drug Program, only)**

- d) As a single agent for the preparation of radioiodine remnant ablation in patients with papillary or follicular thyroid cancer who have undergone thyroidectomy as treatment for thyroid cancer. Thyrotropin may be used in new patients or patients with previously incomplete remnant ablation or who have a recurrence of thyroid cancer and require therapeutic remnant ablation.

**Ticagrelor, tablet, 90mg (Brilinta – AZE and generic)**

To be taken in combination with ASA 75mg -150mg daily<sup>a</sup> for patients with acute coronary syndrome (i.e. ST elevation myocardial infarction (STEMI), non-ST elevation myocardial infarction (NSTEMI), or unstable angina (UA), as follows:

**STEMI<sup>bc</sup>**

- STEMI patients undergoing primary PCI

**NSTEMI or Unstable Angina<sup>bc</sup>**

- Presence of high risk features irrespective of intent to perform revascularization:
  - High GRACE risk score (>140)
  - High TIMI risk score (5-7)
  - Second ACS within 12 months
  - Complex or extensive coronary artery disease e.g. diffuse three vessel disease
  - Definite documented cerebrovascular or peripheral vascular disease
  - Previous CABG

OR

- Undergoing PCI + high risk angiographic anatomy<sup>d</sup>

**Notes:**

- a) Co-administration of ticagrelor with high maintenance dose ASA (>150mg daily) is not recommended.
- b) In the PLATO study more patients on ticagrelor experienced non CABG related major bleeding than patients on clopidogrel, however, there was no difference between the rate of overall major bleeding, between patients treated with ticagrelor and those treated with clopidogrel. As with all other antiplatelet

treatments the benefit/risk ratio of antithrombotic effect vs. bleeding complications should be evaluated.

- c) Ticagrelor is contraindicated in patients with active pathological bleeding, in those with a history of intracranial hemorrhage and moderate to severe hepatic impairment.
- d) High risk angiographic anatomy is defined as any of the following: left main stenting, high risk bifurcation stenting (i.e., two-stent techniques), long stents  $\geq$  38 mm or overlapping stents, small stents  $\leq$  2.5 mm in patients with diabetes.

**Approval will be for a maximum of 12 months.**

**Ticlopidine HCL, tablet, 250mg (Generics)**

- For the secondary prevention of the ischemic stroke or transient ischemic attack (TIA) in patients with a documented severe allergy to ASA (manifested by anaphylactic reaction, asthma, or nasal polyps) or who experience a recurrent thrombotic event (stroke, symptoms of TIA) while taking ASA; or
- For the prevention of thrombosis in patients post intra coronary stent implantation for a period of up to six months.

**GI intolerance to ASA is not considered a criterion for coverage of Ticlopidine,** although severe cases (e.g. gastric ulceration or bleeds) may be considered.

**Tinzaparin** – see Low Molecular Weight Heparins

**Tiotropium** - see Chronic Obstructive Pulmonary Disease

**Tipranavir, capsule, 250mg (Aptivus-BOE)**

For the treatment of adult patients with HIV-1 infection who are treatment experienced, have demonstrated failure to multiple protease inhibitors and in whom no other protease inhibitor is a treatment option.

**Tizanidine HCl, tablet 4mg (Generic)**

For the second- line treatment for those individuals with spasticity resulting from traumatic brain injury, multiple sclerosis, spinal cord injury or cerebral vascular accident and are intolerant to or have had ineffective results from Baclofen and/or benzodiazepines.

**Tocilizumab, IV Vial, 80mg/4l, 200mg/10mL, 400mg/20mL, 162mg/0.9mL (Actemra-HLR)**

**Giant Cell Arteritis**

For the treatment of adult patients with new onset or relapsed giant cell arteritis (GCA) in combination with glucocorticoids (at initiation of therapy, or with relapse).

Initial coverage will be for 16 weeks.

- Reassessment should occur between 12 weeks and 16 weeks of therapy to determine response.

Renewal requests:

- Confirmation of response to treatment (i.e absence of flares AND normalization of C-reactive protein (CRP) to <1mg/dL)

Clinical Note:

- Flare is defined as the recurrence of signs or symptoms of GCA and/or erythrocyte sedimentation rate (ESR) greater or equal to 30 mm/hr attributable to GCA.

Claim Note:

- Must be prescribed by, or in consultation with, a rheumatologist or other physician experienced in the treatment of GCA.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Subcutaneous injection: Approvals will be for 162 mg every week
- Duration of therapy will be limited to 52 weeks per treatment course

Authorization may be granted following any new episode of the disease, according to the treatment terms and conditions previously mentioned for the initial episode.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Juvenile Idiopathic Arthritis**

For the treatment of active systemic juvenile idiopathic arthritis (sJIA), in patients 2 years of age or older, who have responded inadequately to non-steroidal anti-inflammatory drugs (NSAIDs) and systemic corticosteroids (with or without methotrexate) due to intolerance or lack of efficacy.

- Must be prescribed by, or in consultation with, a pediatric rheumatologist.
- Coverage will be approved for an IV dose of 12 mg/kg for patients weighing less than 30kg or 8 mg/kg for patients weighing greater than or equal to 30kg to a maximum of 800mg, administered every two weeks.
- Continued coverage will be dependent on a positive patient response as determined by a pediatric rheumatologist.
- Initial approval period: 16 weeks. Renewal period: 1 year

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**



### **Polyarticular Juvenile Idiopathic Arthritis**

For patients who have had an inadequate response to one or more disease modifying antirheumatic drugs (DMARDs).

- Must be prescribed by, or in consultation with, a pediatric rheumatologist.
- Coverage will be approved for an IV dose of 10 mg/kg for patients weighing less than 30kg or 8 mg/kg for patients weighing greater than or equal to 30kg to a maximum of 800mg, administered every four weeks.
- Continued coverage will be dependent on a positive patient response as determined by a pediatric rheumatologist.
- Initial approval period: 16 weeks. Renewal period: 1 year

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Rheumatoid Arthritis**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks  
AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks;

IV formulation: approvals for adults is 4mg/kg/dose every four weeks, with a maximum maintenance dose escalation up to 8mg/kg, to a maximum of 800mg per infusion.

SC formulation: approvals for adults is 162mg every other week for patients less than 100kg, with a maximum maintenance dose escalation to weekly dosing permitted. Patients equal to or greater than 100kg will be approved for 162mg every week, with no dose escalation permitted

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.
- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration

of treatments specified above.

- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Tofacitinib, tablet, 5mg, 10mg (Xeljanz-PFI and generics); extended release tablet, 11mg (Xeljanz XR-PFI)**

**Rheumatoid Arthritis**

For the treatment of severely active rheumatoid arthritis, in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant to:

Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks  
AND

Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks

NOTE:

Must be prescribed by a rheumatologist.

Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Rheumatoid Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a**

patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.

### **Ulcerative Colitis**

For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score > 4, and a rectal bleeding subscore  $\geq 2$  and are:

- Refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks AND prednisone  $\geq 40\text{mg}$  daily for two weeks or IV equivalent for one week) OR
- Corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year.

Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:

- a decrease in the partial Mayo score  $\geq 2$  from baseline, and
- a decrease in the rectal bleeding subscore  $\geq 1$ .

Clinical Notes:

- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.
- Patients with severe disease (partial Mayo > 6) do not require a trial of 5-ASA

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial approval is for a maximum dose of 10 mg twice daily for 16 weeks.
- Renewal of coverage will require reassessment of the patient and submission of a new Ulcerative Colitis Special Authorization form.
- Renewal: 1 year at a maximum dose of 10 mg twice daily

**The request for coverage must be made by a gastroenterologist using the Ulcerative Colitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

Topamax – see Topiramate

**Topiramate, 15mg, 25mg, sprinkle capsule (Topamax-JAN)**

For patients who require topiramate, cannot take the tablet form, and require sprinkle capsules for proper administration

**Tucatinib, tablet, 50mg 150mg (Tukysa-SGC)**

In combination with trastuzumab and capecitabine for the treatment of patients with locally advanced unresectable or metastatic HER2-positive breast cancer who have received prior treatment with trastuzumab, pertuzumab and a HER2-targeted antibody-drug conjugate (e.g., trastuzumab emtansine or trastuzumab deruxtecan), where at least one was given in the advanced or metastatic setting.

**Clinical Notes:**

1. Patients should have a good performance status.
2. Treatment should be discontinued upon disease progression, unacceptable toxicity, or if both trastuzumab and capecitabine are discontinued.

\* Claims that exceed the maximum claim amount of \$9999.99 must be divided as separate transactions, using DIN first and PDIN\* second.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Tukysa – see Tucatinib**

**Toujeo Solostar– see insulin Glargine**

**Toujeo Doublestar – see insulin Glargine**

**Tracleer - see Bosentan**

**Trajenta – see Linagliptin**

**Trametinib, tablet, 0.5mg, 2mg (Mekinist-NVR)**

**Adjuvant Melanoma**

In combination with dabrafenib for the adjuvant treatment of patients with cutaneous melanoma who meet all of the following criteria:

- Stage IIIA (limited to lymph node metastases of greater than 1 mm) to stage IIID disease (AJCC 8th edition)

- BRAF V600-mutation positive
- Completely resected disease including in-transit metastases

Clinical Notes:

1. Patients must have a good performance status.
2. Treatment should continue until disease recurrence, unacceptable toxicity, or up to a maximum of 12 months.

Claim Notes:

1. Requests will be considered for patients with regional lymph nodes with micrometastases after sentinel lymph node biopsy.
2. Requests will not be considered for patients who received adjuvant immunotherapy for greater than three months. Patients may switch to BRAF targeted therapy within the first three months of initiating immunotherapy to complete a total of 12 months of adjuvant treatment.
3. Approval period: up to 12 months

**Metastatic Melanoma**

For the treatment of patients with BRAF V600 mutation-positive unresectable or metastatic melanoma when used alone or in combination with dabrafenib.

Clinical Notes:

1. Patients must have a good performance status.
2. If brain metastases are present, patients should be asymptomatic or have stable symptoms.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Trelegy Ellipta** – see Fluticasone & Umeclidinium & Vilanterol

**Tremfya** – see Guselkumab

**Tretinoin, capsule, 10mg (Vesanoid - ROC and generic)**

Open benefit if written by an oncologist upon notification to Pharmacare.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Triamcinolone Hexacetonide, ampule, 20mg/mL (Trispan-MED)**

For the treatment of Juvenile Idiopathic Arthritis.

**Trientine, capsule, 250mg (Waymade-Trientine-WMD & Mar-Trientine-MAR)**

For the treatment of Wilson's disease in patients who have experienced intolerance or have a contraindication to d-penicillamine.

Clinical Notes:

- Intolerance is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

1. Treatment must be initiated by clinicians experienced in the management of Wilson's disease for adult patients 18 years of age or older.
2. Treatment must be initiated and renewed by clinicians experienced in the management of Wilson's disease for patients less than 18 years of age.

**Trifluridone & Tipiracil, tablet, 15mg & 6.14mg, 20mg & 8.19mg (Lonsurf-TAI)**

For the treatment of adult patients with metastatic gastric cancer or adenocarcinoma of the gastroesophageal junction who meet the following criteria:

- Previously treated with at least two prior lines of chemotherapy including a fluoropyrimidine, a platinum, and either a taxane or irinotecan and if appropriate, with HER2-targeted therapy.
- Patients should have a good performance status.

Clinical notes:

- Trifluridine/tipiracil should be used in combination with best supportive care
- Treatment should be discontinued upon disease progression or unacceptable toxicity
- Requests will be considered for patients who have an intolerance or contraindication to platinum-based therapy

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Triheptanoin, oral liquid, 8.3kcal/mL (Dojolvi-UGX)**

For the treatment of adult and pediatric patients with an acute life-threatening long-chain fatty acid oxidation disorder (LC-FAOD) who meet the following criteria:

- patients with a confirmed diagnosis of LC-FAOD and acute life-threatening events who require alternative therapy to conventional even-chain medium-chain triglyceride (MCT) supplementation, OR
- patients without a confirmed diagnosis of LC-FAOD presenting with acute life-threatening events consistent with LC-FAOD who require alternative therapy to conventional even-chain MCT supplementation.

Claim Notes:

1. Triheptanoin should only be prescribed by clinicians experienced in the management of LC-FAOD.
2. Approval: 1 year. Confirmation of continued response required.

\* Claims that exceed the maximum claim amount of \$9999.99 must be divided as separate transactions, using DIN first and PDIN\* second.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Trikafta** – see Elexacaftor & Tezacaftor & Ivacaftor & Ivacaftor

**Trileptal** - see Oxcarbazepine

**Trispan** – see Triamcinolone Hexacetonide

**Trosec** - see Trospium

**Trospium, tablet, 20mg (Trosec-SNV)**

For the treatment of over-active bladder (not stress incontinence) after a reasonable trial (e.g. 3 months) of immediate release oxybutynin, solifenacin, or tolterodine is not tolerated.

**Truxima** – see Rituximab

**Turdoze Genuair** – see Acridinium Bromide

**Tysabri** – see Natalizumab

**Ultibro Breezhaler** – see Indacaterol & Glycopyrronium

**Upadacitinib, extended release tablet, 15 mg, 30 mg (Rinvoq-ABV)**

### **Atopic Dermatitis**

For the treatment of moderate to severe atopic dermatitis (AD) in patients 12 years of age and older who meet all of the following criteria:

Patients must have had an adequate trial (with a documented refractory disease), or were intolerant (with documented intolerance), or are ineligible for each of the following therapies:

- maximally tolerated medical topical therapies for AD combined with phototherapy (where available), and;
- maximally tolerated medical topical therapies for AD combined with at least 1 of the 4 systemic immunomodulators (methotrexate, cyclosporine, mycophenolate mofetil, or azathioprine).

AND

Baseline Physician Global Assessment score of 3 or greater and Eczema Area and Severity Index (EASI) of 7.1 or greater.

Renewal criteria:

- Requests for renewal must provide proof of beneficial clinical effect defined as a 75% or greater improvement from baseline in the Eczema Area and Severity Index (EASI-75) score six months after treatment initiation.
- Proof of maintenance of EASI-75 response from baseline must be provided for subsequent authorizations.

Clinical Note:

- Not to be used in combination with phototherapy or any immunomodulatory drugs (including biologics) or a Janus kinase inhibitor treatment for moderate-to-severe AD.

Claim Notes:

- The patient must be under the care of a dermatologist, allergist, clinical immunologist, or pediatrician who has expertise in the management of moderate to severe AD.
- Approvals will be for a maximum of 30 mg daily.
- Initial approval period: 6 months.
- Renewal approval period: 1 year.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Psoriatic Arthritis**

For the treatment of predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs at a maximum tolerated dose for a minimum of two weeks each.

For the treatment of predominantly peripheral psoriatic arthritis who are refractory or intolerant or have contraindications to:



- Sequential use of at least two NSAIDs at a maximum tolerated dose for a minimum of two weeks each; and
- Methotrexate (oral or parenteral) at a dose of  $\geq 20$ mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; and
- Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use with other biologic drugs will not be reimbursed.
- Approvals will be for a maximum of 15 mg daily.
- Initial approval 16 weeks.
- Renewal approval: 1 year. Confirmation of continued response is required.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Rheumatoid Arthritis**

For the treatment of moderately to severely active rheumatoid arthritis, alone or in combination with methotrexate or other disease-modifying antirheumatic drugs (DMARDs), in adult patients who are refractory or intolerant or have contraindications to:

- Methotrexate (oral or parenteral), alone or in combination with another DMARD, at a dose of  $\geq 20$  mg weekly ( $\geq 15$ mg if patient is  $\geq 65$  years of age), for a minimum of 12 weeks AND
- Methotrexate in combination with at least two other DMARDs, such as hydroxychloroquine and sulfasalazine, for a minimum of 12 weeks.

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Optimal treatment response to DMARDs may take up to 24 weeks, however coverage of a biologic therapy can be considered if no improvement is seen after 12 weeks of triple DMARD use.

- If patient factors (e.g. intolerance) prevent the use of triple DMARD therapy, these must be described and dual therapy with DMARDs must be tried.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Combined use with other biologic drugs or janus kinase inhibitors will not be reimbursed.
- Approvals will be for a maximum of 15 mg daily.
- Initial Approval: 6 months
- Renewal Approval: 1 year. Confirmation of continued response is required.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Uptravi** – see Selexipag

**Urispas** - see Flavoxate

**Ustekinumab, syringe, 45mg/0.5mL, 90mg/mL (Jamteki-JPC; Stelara-JAN; Wezlana-AMG); vial, 45mg/0.5mL (Wezlana-AMG)**

For ustekinumab-naïve patients whose ustekinumab therapy is initiated after August 26, 2024, an ustekinumab biosimilar will be the product approved.

Patients with existing PEI Pharmacare coverage for Stelara® will need to switch to a biosimilar version before August 31, 2025, or by the renewal date of their current special authorization, whichever is earlier, to maintain coverage through PEI Pharmacare.

**Plaque Psoriasis**

For patients with severe, debilitating chronic plaque psoriasis who meet all of the following:

- Body surface area (BSA) involvement of >10% and/or significant involvement of the face, hands, feet or genitals;
- Failure to, contraindication to or intolerant of methotrexate (oral or parenteral) at a dose of ≥ 20mg weekly (≥15mg if patient is ≥65 years of age) for a minimum of 12 weeks or cyclosporine for a minimum of 6 weeks; AND
- Failure to, intolerant of or unable to access phototherapy.

Continued coverage is dependent on evidence of improvement, specifically:

- A >75% reduction (provide baseline and current score) in the Psoriasis Area and Severity Index (PASI) score; or
- A >50% reduction (provide baseline and current score) in PASI with a > 5 point improvement in DLQI (Dermatology Life Quality Index); or
- Significant reduction in BSA involved, with consideration of important regions such as the face, hands, feet or genitals.

Clinical Note:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Failure is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Treatment should be discontinued if a response has not been demonstrated after 16 weeks.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim notes:

- Approvals will be for a maximum adult dose of up to 90 mg at 0, 4, and 16 weeks. If response criteria is met at 16 weeks, approval will be continued to a maximum dose of up to 90 mg every 12 weeks up to one year.
- Initial approval: 16 weeks.
- Renewal approval: 1 year.
- Concurrent use of biologics not approved.

**Requests must be made by a dermatologist using the Plaque Psoriasis Special Authorization form which is available from the Drug Programs office or on-line at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ustekinumab, syringe, 45mg/0.5mL, 90mg/mL (Jamteki-JPC; Wezlana-AMG); vial, 45mg/0.5mL (Wezlana-AMG)**

**Psoriatic Arthritis**

- 1) For the treatment of patients with predominantly axial psoriatic arthritis who are refractory, intolerant or have contraindications to the sequential use of at least two NSAIDs for a minimum of two weeks each.
- 2) For the treatment of predominantly peripheral psoriatic arthritis who are refractory, intolerant or have contraindications to:
  - Sequential use of at least two NSAIDs for a minimum of two weeks each; AND

- Methotrexate (oral or parenteral) at a dose of  $\geq 20\text{mg}$  weekly ( $\geq 15\text{mg}$  if patient is  $\geq 65$  years of age) for a minimum of 8 weeks; AND
- Leflunomide for a minimum of 10 weeks or sulfasalazine for a minimum of 3 months

Clinical Notes:

- For patients who do not demonstrate a clinical response to oral methotrexate, or who experience gastrointestinal intolerance, a trial of parenteral methotrexate must be considered.
- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.

Claim Notes:

- Must be prescribed by a rheumatologist.
- Concurrent use of biologics not approved.
- Initial period 6 months.
- Approvals will be for a maximum of 45mg subcutaneously at Weeks 0 and 4, and maintenance dosing of 45mg subcutaneously every 12 weeks. For patients  $>100\text{kg}$ , doses of 90mg may be considered.
- Renewal approval: 1 year. Confirmation of continued response required.

**The request for coverage must be made by a rheumatologist or prescriber with a specialty in rheumatology, using the Psoriatic Arthritis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Ustekinumab, vial, 45mg/0.5mL; syringe, 45mg/0.5mL, 90mg/mL (Wezlana-AMG); vial, 130mg/26mL (Wezlana I.V.-AMG)**

**Crohn's Disease**

For the treatment of patients with moderate to severe Crohn's disease who have active disease and are refractory, intolerant or have contraindications to:

- Prednisone 40mg (or equivalent) daily for  $\geq 2$  weeks, AND
- Azathioprine  $\geq 2 \text{ mg/kg/day}$  for  $\geq 3$  months, OR
- Mercaptopurine  $\geq 1 \text{ mg/kg/day}$  for  $\geq 3$  months, OR

- Methotrexate (SC or IM)  $\geq$  15 mg/week for  $\geq$  3 months

Clinical Notes:

- Refractory is defined as lack or loss of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.
- Consideration will be given for the approval of a biologic DMARD (disease modifying antirheumatic drug) without a trial of a traditional DMARD for patients who have an aggressive/severe disease course (e.g. extensive disease, a modified Harvey Bradshaw Index score  $>$  16) and are refractory, intolerant or have contraindications to systemic corticosteroids.

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use with other biologic drugs or janus kinase (JAK) inhibitors will not be reimbursed.
- Initial reimbursement will be for a single intravenous dose of up to 520mg at Week 0 and a subcutaneous dose of 90mg at Week 8 and 16. Subsequent reimbursement for maintenance dosing is 90mg subcutaneously every 8 weeks.
- Initial Approval: 16 weeks
- Renewal Approval: 1 year

**The request for coverage must be made by a gastroenterologist using the Crohn's Disease Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Ulcerative Colitis**

For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score  $>$  4, and a rectal bleeding subscore  $\geq$  2 and are:

- Refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks AND prednisone  $\geq$  40mg daily for two weeks or IV equivalent for one week) OR
- Corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year.

Clinical Notes:

- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.
- Patients with severe disease (partial Mayo > 6) do not require a trial of 5-ASA

Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Combined use of more than one biologic DMARD will not be reimbursed.
- Initial reimbursement will be for a single intravenous dose of up to 520mg at Week 0 and a subcutaneous dose of 90mg at Week 8 and 16. Subsequent reimbursement for maintenance dosing is 90mg subcutaneously every 8 weeks.
- Initial Approval: 16 weeks
- Renewal Approval: 1 year

**The request for coverage must be made by a gastroenterologist using the Ulcerative Colitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Vabysmo** – see Faricimab

**Valcyte** - see Valganciclovir

**Valganciclovir, tablet, 450mg (Valcyte-CAG and generics)**

- a) For the treatment of cytomegalovirus (CMV) retinitis in patients with AIDS.
- b) For the prevention of cytomegalovirus (CMV) disease in solid organ transplant patients at risk (where either the donor or the recipient is CMV +).

**Valganciclovir, oral suspension, 50mg/mL (Valcyte-XPI)**

Requests for oral suspension will be considered for patients when oral tablets are not an option, for the following indications:

- a) For the treatment of cytomegalovirus (CMV) retinitis in patients with AIDS.
- b) For the prevention of cytomegalovirus (CMV) disease in solid organ transplant patients at risk (where either the donor or the recipient is CMV +).

## **Vandetanib, tablet, 100mg, 300mg (Caprelsa-GZY)**

For the treatment of symptomatic and/or progressive medullary thyroid cancer (MTC) in patients with unresectable locally advanced or metastatic disease.

Treatment should be for patients with a good performance status and should continue until disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

## **Vedolizumab, prefilled pen, prefilled syringe (108 mg/0.68 mL), vial, 300mg (Entyvio-TAK)**

### **Crohn's Disease**

For the treatment of patients with moderate to severe Crohn's disease who have active disease and are refractory, intolerant or have contraindications to:

- Prednisone 40mg (or equivalent) daily for  $\geq 2$  weeks, AND
- Azathioprine  $\geq 2$  mg/kg/day for  $\geq 3$  months, OR
- Mercaptopurine  $\geq 1$  mg/kg/day for  $\geq 3$  months, OR
- Methotrexate (SC or IM)  $\geq 15$  mg/week for  $\geq 3$  months

#### Clinical Notes:

- Refractory is defined as lack or loss of effect at the recommended doses and for duration of treatments specified above.
- Intolerant is defined as demonstrating serious adverse effects to treatments. The nature of intolerance(s) must be clearly documented.
- Consideration will be given for the approval of a biologic DMARD (disease modifying antirheumatic drug) without a trial of a traditional DMARD for patients who have an aggressive/severe disease course (e.g. extensive disease, a modified Harvey Bradshaw Index score  $> 16$ ) and are refractory, intolerant or have contraindications to systemic corticosteroids.

#### Claim notes:

- Intravenous infusion: Initial approval for adults is for induction doses of 300mg at weeks 0, 2, and 6.
- Subcutaneous injection: Approvals will be for a maximum of 108 mg every two weeks following at least two intravenous infusions of vedolizumab. Renewal of coverage will require reassessment of the patient and submission of a new Crohn's Disease Special Authorization form. Continued coverage may be approved at a dose not exceeding 300mg every 8 weeks.

- Renewal Approval: 1 year. Confirmation of continued response is required.
- Combined use of more than one biologic DMARD will not be reimbursed.

**The request for coverage must be made by a gastroenterologist using the Crohn's Disease Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

### **Ulcerative Colitis**

For the treatment of adult patients with moderately to severely active ulcerative colitis who have a partial Mayo score > 4, and a rectal bleeding subscore  $\geq 2$  and are:

- Refractory or intolerant to conventional therapy (i.e. aminosalicylates for a minimum of four weeks AND prednisone  $\geq 40$ mg daily for two weeks or IV equivalent for one week) OR
- Corticosteroid dependent (i.e. cannot be tapered from corticosteroids without disease recurrence; or have relapsed within three months of stopping corticosteroids; or require two or more courses of corticosteroids within one year.

#### Claim Notes:

- Must be prescribed by a gastroenterologist or physician with a specialty in gastroenterology.
- Intravenous infusion: Initial approval is for induction doses of 300mg at weeks 0, 2, and 6, then 300 mg every eight weeks.
- Subcutaneous injection: Approvals will be for a maximum of 108 mg every two weeks following at least two intravenous infusions of vedolizumab. Renewal of coverage will require reassessment of the patient and submission of a new Ulcerative Colitis Special Authorization form.
- Continued coverage will be approved at a dose not exceeding 300mg every 8 weeks.

Renewal requests must include information demonstrating the beneficial effects of the treatment, specifically:

- a decrease in the partial Mayo score  $\geq 2$  from baseline, and
  - a decrease in the rectal bleeding subscore  $\geq 1$ .
- Combined use of more than one biologic DMARD will not be reimbursed.

#### Clinical Notes:

- Refractory is defined as lack of effect at the recommended doses and for duration of treatments specified above.



- Intolerant is defined as demonstrating serious adverse effects or contraindications to treatments as defined in product monographs. The nature of intolerance(s) must be clearly documented.
- Patients with severe disease (partial Mayo > 6) do not require a trial of 5-ASA

**The request for coverage must be made by a gastroenterologist using the Ulcerative Colitis Special Authorization form available from the Drug Programs office or online at <http://www.princeedwardisland.ca/pharmacareforms>**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Velaglucerase alfa, vial, 400 unit (VPRIV-SHR)**

For patients with Gaucher disease type 1 (GD1) who meet established clinical criteria.

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Velphoro – see SuCroferric oxyhydroxide**

**Vemurafenib, tablet, 240mg (Zelboraf-HLR)**

As a first line, single agent for the treatment of BRAF V600 mutation positive unresectable or metastatic melanoma in patients with an ECOG performance status (PS) of 0 or 1. For BRAF V600 mutation positive patients who have progressed after first line treatment prior to vemurafenib availability, funding or vemurafenib as a second line agent may be considered.

**OR**

For use in combination with cobimetinib, for the treatment of patients with previously untreated BRAF V600 mutation-positive unresectable stage III or IV melanoma who have a good performance status. Treatment should continue until unacceptable toxicity or disease progression. If brain metastases are present, patients should be asymptomatic or have stable symptoms.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Venclexta – see Venetoclax**

## Venetoclax, tablet, starter pack, 10mg, 50mg 100mg (Venclexta-ABV)

1. Monotherapy:
  - As monotherapy in patients with chronic lymphocytic leukemia (CLL) who have received at least one prior therapy and who have failed a B-cell receptor inhibitor (BCRi).
  - Patients should have good performance status and treatment should be continued until disease progression or unacceptable toxicity.
2. Combination therapy:
  - As combination therapy with rituximab for the treatment of adult patients with chronic lymphocytic leukemia (CLL) who have received at least one prior therapy, irrespective of their 17p deletion status.
  - Patients should be continued until disease progression or unacceptable toxicity up to a maximum of two years, whichever comes first.

### Clinical Notes:

- Patients currently receiving and responding to venetoclax monotherapy, but who have not achieved an adequate response are eligible to have rituximab added to venetoclax. The funded duration of venetoclax therapy from the point rituximab addition will be up to a maximum of 2 years.
- Patients may be re-treated with venetoclax plus rituximab if they responded to and completed two years of therapy with at least 12 months of progression-free interval.
- Patients with relapsed CLL will be eligible for sequencing venetoclax + rituximab and ibrutinib in second or third line settings, for either intolerance or disease progression, providing patients have not received prior treatment with either option and meet all other funding criteria.

3. Combination therapy:
  - In combination with obinutuzumab for the treatment of adult patients with previously untreated chronic lymphocytic leukemia (CLL) or small lymphocytic lymphoma (SLL) who are fludarabine ineligible.

### Clinical Notes:

- Treatment should be given for a total of 12 months (six 28-day cycles in combination with obinutuzumab, followed by six months of monotherapy), or until disease progression or unacceptable toxicity, whichever occurs first.
- Retreatment with a venetoclax based regimen is funded if relapse is greater than 12 months from completion of venetoclax in combination with obinutuzumab.
- Either ibrutinib or acalabrutinib is funded as a subsequent treatment option, provided all other funding criteria are met.

4. Combination therapy:

- In combination with azacitidine for the treatment of patients with newly diagnosed acute myeloid leukemia (AML) who are 75 years of age or older, or who have comorbidities that preclude the use of intensive induction chemotherapy.

Clinical Notes:

- Treatment should continue until disease progression or unacceptable toxicity.
- All newly diagnosed AML patients who are ineligible for induction chemotherapy are eligible regardless of cytogenetic risk.,
- On a time-limited need, patients who are currently receiving azacitidine for newly diagnosed AML may have venetoclax added to their treatment provided there is no disease progression and patient otherwise meets criteria.

Claim Notes:

- Patients who have been previously treated with a hypomethylating agent or chemotherapy for the treatment of myelodysplastic syndromes (MDS) are not eligible for treatment with venetoclax in combination with azacitidine.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Verkazia** – see Cyclosporine

**Verzenio** – see Abemaciclib

**Vesanoid** – see Tretinoin

**Vigabatrin, tablet, 500mg (Sabril-LUD)**

- Vigabatrin is an alternative treatment option for patients who have had an inadequate response or intolerance to other antiepileptic drug combinations
- A restricted benefit status is appropriate due to the risk of ophthalmological adverse effects associated with vigabatrin

**Vigamox** – see Moxifloxacin

**Vismodegib, capsule, 150mg (Erivedge-HLR)**

For the treatment of locally advanced BCC (including basal cell nevus syndrome i.e. Gorlin syndrome who are 18 years of age and older) in patients who are inappropriate for surgery and radiotherapy based on a discussion/evaluation with other members of the multi-disciplinary team OR

As a single agent for the treatment of measurable metastatic basal cell carcinoma (BCC)

Clinical Note:

1. Patients must have an ECOG performance status of  $\leq 2$

Note: Vismodegib (Erivedge) is only available through a controlled distribution program called the Erivedge Pregnancy Prevention Program (EPPP). Under this program, only prescribers and pharmacies registered with the program are able to prescribe and dispense the product, respectively. In addition, Vismodegib can only be dispensed to patients who are registered and meet all the conditions of the EPPP.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Vitrakvi** – see Larotrectinib

**Vocabria** – see Cabotegravir

**Volibris** – see Ambrisentan

**Voriconazole, tablet, 50mg, 200mg (Vfend-PFI and generics)**

**Candidemia:** For the treatment of culture proven invasive candidiasis with documented resistance to fluconazole.

**Aspergillosis, invasive:** For the management of invasive aspergillosis. Initial requests will be approved for a maximum of 3 months.

**Must be prescribed in consultation with a specialist in infectious diseases or medical microbiology.**

**Votrient** – see Pazopanib

**VPRIV** – see Velaglucerase Alfa

**Vyepti** – see Eptinezumab

**Vyndaqel** – see Tafamidis meglumine

**Vyndamax** – see Tafamidis

**Vyvanse** – see Lisdexamfetamine

**Wezlana** – see Ustekinumab

**Xalkori** – see Crizotinib

**Xeljanz** – see Tofacitinib

**Xeljanz XR** – see Tofacitinib

**Xolair** – see Omalizumab

**Xospata** – see Gilteritinib

**Xpovio** – see Selinexor

**Xtandi** – see Enzalutamide

**Yuflyma** - see Adalimumab

**Zanubrutinib, capsule, 80mg (Brukinsa-BIG)**

For the treatment of adult patients with relapsed or refractory Waldenstrom macroglobulinemia who have received at least one prior therapy and have not experienced disease progression on a Bruton's tyrosine kinase inhibitor.

**Clinical Notes:**

1. Patients must meet at least one criterion for treatment as per IWWM consensus panel.
2. Patients must have a good performance status and no evidence of disease transformation.
3. Treatment should be discontinued upon disease progression or unacceptable toxicity.

**Prescriptions written by PEI oncologists do not require written Special Authorization.**

**Patients requesting coverage through the High Cost Drug Program must submit a patient application, available from the Drug Programs Office or online at <http://www.princeedwardisland.ca/pharmacareforms>.**

**Zaxine** – see Rifaximin

**Zelboraf** – see Venurafenib

**Zeposia** – see Ozanimod

**Ziextenzo** – see Pegfilgrastim

**Ziprasidone hydrochloride. Capsule, 20mg, 40mg, 60mg, 80mg (Zeldox-PFI and generic)**

For the treatment of schizophrenia and schizoaffective disorders in patients who have a contraindication to a trial of at least TWO less expensive antipsychotic agents because of intolerance or lack of response.

**Zithromax** - see Azithromycin

**Zofran** - see Ondansetron

**Zydelig** – see Idelalisib

**Zykadia** – see Ceritinib

**Zymar** – see Gatifloxacin

**Zytiga** – see Abiraterone

**Zyvoxam** - see Linezolid

## **APPENDIX B**

### **Links to Drug Program Forms**

#### **Special Authorization Forms**

[Ankylosing Spondylitis Special Authorization Form](#)

[Crohn's Disease Special Authorization Form](#)

[Diabetes Glucose Sensor Program](#)

[DPP-4/SGLT2 Inhibitors](#)

[Enfuvirtide Special Authorization Form](#)

[Idiopathic Pulmonary Fibrosis Special Authorization Form](#)

[Long Acting Insulin Analogues Special Authorization Form](#)

[Plaque Psoriasis Special Authorization Form](#)

[Psoriatic Arthritis Special Authorization Form](#)

[Rheumatoid Arthritis Special Authorization Form](#)

[Standard Special Authorization Form](#)

[Ulcerative Colitis Special Authorization Form](#)

#### **Program Application Forms**

[Catastrophic Drug Program Application Form](#)

[Diabetes Glucose Sensor Program](#)

[Diabetes Referral Form](#)

[Erythropoietin Program Approval Form](#)

[Family Health Benefits Drug Program - Application Form](#)

[High Cost Drug Program - Application Form](#)

[Home Oxygen Program - Application Form](#)

[Insulin Pump Program](#)

[Ostomy Supplies Program Application Form](#)

[Ostomy Supplies Program Registration Form for Health Care Providers](#)

## APPENDIX C List of Manufacturer Abbreviations

AAA	AA Pharmaceuticals Inc.
ABB	Abbott Laboratories Ltd.
ABC	Abbott Diabetes Care
ABV	Abbvie Corporation
ACC	Accel Pharma
ACH	Accord Healthcare
ACS	Acerus Pharmaceuticals Corp.
ALC	Alcon Canada Inc.
ALH	Altius Healthcare
ALL	Allergan Inc.
ALY	Amylyx Pharmaceuticals
AMB	Ambicare Pharmaceuticals In.
AMD	Amdipharm Limited
AMG	Amgen Canada Inc.
ANB	ANB Canada
ANG	Angita Pharma.
APX	Apotex Inc.
ARO	Auro Pharma Inc
ARN	Accelera Pharma Canada Inc.
ASN	Aspen Pharma Trading Ltd.
AST	Astellas Pharma Canada, Inc.
AVI	Avir Pharma Inc.
ATL	Laboratoire Atlas
ATN	Atnahs Pharma UK Ltd.
AVN	Sanofi-Aventis
AZE	AstraZeneca Canada Inc.
BAX	Baxter Corporation
BAY	Bayer Inc.
BDD	Bayer Healthcare, Diabetes Care Division
BIG	Biogen Idec Canada Inc.
BGP	BGP Pharma Ulc.
BIG	Beigene (Canada) ULC
BIN	Bionime Corporation
BLO	Bausch & Lomb Inc.
BMS	Bristol-Myers Squibb Canada
BOE	Boehringer Ingelheim (Canada) Ltd.
BOX	Biocodex S.A.
CDC	Church & Dwight Canada Corp.
CEL	Celgene Inc.
CIP	Cipher Pharmaceuticals
CHE	Cheplapharm Arzneimittel GMBH.
COV	Covis Pharma Canada Ltd.
DUI	Duchesnay Inc.
D&C	D&C Mobility Solutions Inc.
EIS	Eisai Limited
ELV	Elvium Life Sciences
END	Endomedical
ERF	Erfa Canada Inc.
EPM	Essential Pharma
ETH	Ethypharm Inc.
FEI	Ferring Inc.
FKB	Fresenius Kabi Canada



FTI	Forus Therapeutics
GAC	Galderma Canada Inc.
GIL	Gilead Sciences, Inc.
GMD	GenMed, Division of Pfizer Canada
GMP	Generic Medical Partners
GSK	GlaxoSmithKline Inc.
GLM	Glenmark Generic
GZY	Sanofi Genzyme
ICL	Indivior Canada Ltd.
HLR	Hoffmann-La Roche Limited
HOS	Hospira Healthcare Corporation
JAC	Jacobus Pharmaceutical Company Inc.
JAN	Janssen Inc..
JJM	Johnson & Johnson - Merck Consumer Pharmaceuticals of Canada
JPC	Jamp Pharma
KNI	Knight Therapeutics Inc.
KYE	Kye Pharmaceuticals
LBI	Leadiant Biosciences Inc.
LEO	Leo Pharma Inc.
LIL	Eli Lilly Canada Inc.
LTH	Labtician Thea
LSN	Life Scan Canada Ltd.
LUD	Lundbeck Canada Inc.
LUP	Lupin Pharma
MRA	MantraPharma
MAR	Marcan Pharmaceuticals Inc.
MCL	McNeil Consumer Healthcare
MDN	MDA Inc.
MDU	Medunik Canada
MSR	Medisure Canada Inc.
MDA	3M Pharmaceuticals
MDP	Deciphera Pharmaceuticals
MFI	Medical Futures Inc.
MNT	Mint Pharmaceuticals
MJS	Mead Johnson Canada, Division of Bristol-Myers Squibb Canada Inc.
MRS	Merus Labs
MSD	Merck Frosst Canada Ltd.
MTP	Methapharm Inc.
MYL	Mylan Pharmaceuticals
NAT	Natco Pharma
NRA	Nora Pharma Inc.
NNO	Novo Nordisk Canada Inc.
NVR	Novartis Pharmaceuticals Canada Inc.
ODN	Odan Laboratories Ltd.
OMG	Omega Laboratories Ltd.
ORG	Organon Canada
OTS	Otsuka Canada Pharmaceuticals
PAL	Paladin Labs Inc.
PEN	Pendopharm, Division of Pharmascience Inc.
PFI	Pfizer Canada ULC
PSL	Pharma Stullin
PFR	Purdue Pharma
PGA	Proctor & Gamble Inc.

PMS	Pharmascience Inc.
PRZ	Pharmaris Canada
RAN	Ranbaxy Pharmaceuticals Canada Inc.
RCH	Dr. Reddy's Laboratory
RIV	Laboratoire Riva Inc.
ROC	Roche Diagnostics
ROG	Rougier Pharma Inc., Division of Ratiopharm Inc.
ROS	Ross Laboratories, Division of Abbott Laboratories Ltd.
SDZ	Sandoz Canada Inc.
SGC	Seagen Inc.
SNN	Santen Inc.
SEV	Servier Canada Inc.
SHR	Shire Biochem Inc.
SIV	Sivem Pharmaceutical
SLP	Searchlight Pharma Inc.
SNE	Smith & Nephew Inc.
SNS	Sanis Health Inc.
SRO	Serono Canada Inc.
STE	Sterimax Inc.
STR	Strides Pharma.
SNV	Sunovion Pharmaceuticals Canada
TAK	Takeda Canada Inc.
TAR	Taro Pharmaceuticals Inc.
TAV	Tanvex BioPharma USA
TPG	Tillotts Pharma
TRT	TerSera Therapeutics
TEV	Teva Canada Ltd.
TLG	Teligent
TRI	Trividia Health
UCB	UCB Canada Inc.
UGX	Ultragenyx Pharmaceutical
UJC	Upjohn Canada ULC
VAL	Valeant Canada Limited
VII	VIIV Healthcare ULC
WES	WellSpring Pharmaceutical Canada
XPI	Xediton Pharmaceuticals Inc.

## Appendix D Insulin Pump Program Approved Vendors List

*Revised June 1, 2023*

### Medtronic of Canada Insulin Pumps and Supplies

Device Name	Model Number	Description
MiniMed 630G-Insulin Pump	MMT-1754K	Pump 3.0 reservoir capacity <ul style="list-style-type: none"> <li>Black</li> </ul>
MiniMed 670G-Insulin Pump	MMT-1762KCN*	Pump 3.0 reservoir capacity <ul style="list-style-type: none"> <li>Black</li> </ul>
MiniMed 770G-Insulin Pump	MMT- 1891 CN	Pump 3.0 reservoir capacity <ul style="list-style-type: none"> <li>Black</li> </ul>
MiniMed 780G- Insulin Pump	MMT- 1895 CN	Pump 3.0 reservoir capacity <ul style="list-style-type: none"> <li>Black</li> </ul>
Reservoir for 5 series MiniMed® Paradigm® Pump (1.8 mls reservoir)	MMT-326A	1.8 mls reservoir for use in 5 series Paradigm Insulin Pump (10 reservoirs /box)
Reservoir for 7 Series MiniMed® Paradigm® Pump (3.0mls reservoir)	MMT-332A	3.0 mls reservoir for use in 7 series Paradigm Insulin Pump only (10 reservoirs /box)
Medtronic Extended Reservoir 3.0 ML ***	MMT-342	3.0 mls reservoirs (10 reservoirs per box)
Quick –Serter	MMT-305QS600	Insertion device for Quickset infusion sets
Device Name	Model Number	Description
Medtronic Extended wear infusion set 6MM – box of 3 sets ***	MMT-431AH	6mm cannula with 60cm (23") tubing. 3 infusion sets per box. Up to 7 day wear time
Medtronic Extended wear infusion set 9MM – box of 3 sets ***	MMT-441AH	9mm cannula with 60cm (23") tubing. 3 infusion sets per box. Up to 7 day wear time
Medtronic Extended wear infusion set 9MM – box of 3 sets ***	MMT-442AH	9mm cannula with 80cm (32") tubing. 3 infusion sets per box. Up to 7 day wear time
Medtronic Extended infusion set 6MM – box of 1 set ***	MMT-431AJ	6mm cannula with 60cm (23") tubing. 1 infusion set per box. Up to 7 day wear time
Medtronic Extended infusion set 9MM – box of 1 set ***	MMT-441AJ	9mm cannula with 60cm (23") tubing. 1 infusion set per box. Up to 7 day wear time
Medtronic Extended infusion set 9MM – box of 1 set***	MMT-442AJ	9mm cannula with 80cm (32") tubing. 1 infusion set per box. Up to 7 day wear time
MiniMed Mio 30 infusion set	MMT 905A600	13mm cannula infusion set with 60cm (23 ") tubing GRAY (10 per box)
	MMT 906A600	13mm cannula infusion set with 110cm (43 ") tubing GRAY (10 per box)
Quickset® Infusion Sets <ul style="list-style-type: none"> <li>6mm or 9mm cannula</li> </ul>	MMT-394A600	6 mm teflon cannula infusion set with 45cm (18") tubing (10 cannula and 10 tubing / box)
	MMT-399A600	6 mm teflon cannula infusion set with 60cm (23")

Quickset® Infusion Sets(cont'd) <ul style="list-style-type: none"> <li>6mm or 9mm cannula</li> </ul>		tubing (10 cannula and 10 tubing / box)
	MMT-387A600	6 mm teflon cannula infusion set with 80cm (32") tubing (10 cannula and 10 tubing / box)
	MMT-398A600	6 mm teflon cannula infusion set with 110cm (43") tubing (10 cannula and 10 tubing / box)
	MMT-397A600	9 mm teflon cannula infusion set with 60cm (23") tubing (10 cannula and 10 tubing / box)
	MMT-386A600	9 mm teflon cannula infusion set with 80cm (32") tubing (10 cannula and 10 tubing / box)
	MMT-396A600	9 mm teflon cannula infusion set with 110cm (43") tubing (10 cannula and 10 tubing / box)
<b>Device Name</b>	<b>Model Number</b>	<b>Description</b>
MiniMed Mio Advance Infusion Sets <ul style="list-style-type: none"> <li><i>P cap connectors</i> - Compatible with Medtronic insulin pumps only</li> <li><i>Luer Lock connectors</i>- Compatible with Non-Medtronic durable insulin pumps only</li> </ul>	MMT-242600	6mm cannula infusion set with Length - 23 " tubing, P-cap connector, (10 cannula and 10 tubing / box) Color- Clear
	MMT-243A600	9mm cannula infusion set with Length - 23 " tubing, P-cap connector, (10 cannula and 10 tubing / box) Color- Clear
	MMT-244A600	9mm cannula infusion set with Length - 43 " tubing, P-cap connector, (10 cannula and 10 tubing / box) Color- Clear
	MMT-247600	6mm cannula infusion set with Length - 23 " tubing, Luer lock connector, (10 cannula and 10 tubing / box) Color- Clear
	MMT-248600	9mm cannula infusion set with Length - 43 " tubing, Luer lock connector, (10 cannula and 10 tubing / box) Color- Clear
mio™ Infusion Sets <ul style="list-style-type: none"> <li>All in one infusion set and insertion device</li> <li>6mm or 9mm cannula</li> </ul>	MMT-921A600	6 mm teflon cannula infusion set with 45cm (18") tubing PINK (10 cannula and 10 tubing / box)
	MMT-941A600	6 mm teflon cannula infusion set with 45cm (18") tubing BLUE (10 cannula and 10 tubing / box)
	MMT-923A600	6 mm teflon cannula infusion set with 60cm (23") tubing PINK (10 cannula and 10 tubing / box)
	MMT-943A600	6 mm teflon cannula infusion set with 60cm (23") tubing BLUE (10 cannula and 10 tubing / box)
	MMT-965A600	6 mm teflon cannula infusion set with 80cm (32") tubing CLEAR (10 cannula and 10 tubing / box)
	MMT-975A600	9 mm teflon cannula infusion set with 80cm (32") tubing CLEAR (10 cannula and 10 tubing / box)
Mio 30™	MMT-905A600	13 mm cannula infusion set with 60cm (23") tubing GRAY 10/Box
	MMT-906A600	13 mm cannula infusion set with 110cm (43") tubing GRAY 10/Box
Silhouette® Infusion Sets <ul style="list-style-type: none"> <li>13mm or 17mm cannula</li> </ul>	MMT-371	Silhouette 43" Full Set 10/Box
	MMT-373	Silhouette 23" Full Set 10/Box
	MMT-368A600	13 mm teflon cannula infusion set with 45cm (18") tubing (10 cannula and 10 tubing / box)
	MMT-381A600	13 mm teflon cannula infusion set with 60cm

		(23") tubing (10 cannula and 10 tubing / box)
	MMT-383A600	13 mm teflon cannula infusion set with 80cm (32") tubing (10 cannula and 10 tubing / box)
	MMT-382A600	13 mm teflon cannula infusion set with 110cm (43") tubing (10 cannula and 10 tubing / box)
	MMT-378A600	17 mm teflon cannula infusion set with 60cm (23") tubing (10 cannula and 10 tubing / box)
	MMT-384A600	17 mm teflon cannula infusion set with 80cm (32") tubing (10 cannula and 10 tubing / box)
	MMT-377A600	17 mm teflon cannula infusion set with 110cm (43") tubing (10 cannula and 10 tubing / box)
	MMT- 369600	Silhouette Cannula only, 13 mm 10 cannulas / box
	MMT- 370600	Silhouette Cannula only, 17 mm 10 cannulas / box
Sure-T Infusion Sets <ul style="list-style-type: none"> <li>• Needle infusion set</li> <li>• (90° angle of insertion)</li> </ul>	MMT-862A	6mm <u>needle</u> infusion set with 45cm (18") tubing (10 needles & 10 tubing / box)
	MMT-864A	6mm <u>needle</u> infusion set with 60cm (23") tubing (10 needles & 10 tubing / box)
	MMT-866A	6mm <u>needle</u> infusion set with 80cm (32") tubing (10 needles & 10 tubing / box)
	MMT-874A	8mm <u>needle</u> infusion set with 60cm (23") tubing (10 needles & 10 tubing / box)
<b>Additional supplies – Skin Preparation and Skin Tape</b>		
Skin Prep Wipes	HMS-59420425	Skin Prep Adhesive wipes (box of 50)
Skin Tact Wipes	HMS-180	Skin Tact Wipes (50 / box)
Tape Dressing	MMT-134A	Polyskin Tape Dressing (100 / box)
Adhesive patch	MMT-172	Acutek Non-Sterile Sof-set Adhesive Patch (50/box)
Transparent dressing	MMT-174	IV 3000 1-Hand with Strips and Label (100 / box)
Transparent dressing	HMS-175	IV 3000 Adhesive patch Large (50/box)
Transparent dressing	HMS-66800786	IV 3000 Adhesive tape 1/3" x 2 3/4" transparent dressing (box of 30)
Adhesive remover wipes	403120	Universal Adhesive Remover Wipes (50 / box)

### Insulet Canada Insulin Pumps and Supplies

<b>Insulin Pumps</b>		
<b>Device Name</b>	<b>Model Number</b>	<b>Description</b>
Omnipod Insulin Management System Starter Kit (ENGLISH)	SKT-CAT45E	<ul style="list-style-type: none"> <li>• 1 Personal Diabetes Manager</li> <li>• 1 USB Cable</li> <li>• 1 User guide</li> <li>• 1 Carrying case</li> <li>• 1 Software CD</li> <li>• Choice of PDM gel skin cover (7 colors available)</li> </ul>
Omnipod Insulin Management System Starter Kit (FRENCH)	SKT-CAT45F	
Omnipod DASH® Insulin Management System – Personal Diabetes Manager (PDM) Starter Kit [Bilingual]	SKT-CAN-D001-MM	

<b>Pump Supplies</b>		
<b>Device Name</b>	<b>Model Number</b>	<b>Description</b>
Omnipod Insulin management System (POD)	POD-ZXR425	Internal insulin reservoir and pumping mechanism <ul style="list-style-type: none"> <li>• Small and lightweight</li> <li>• Strong adhesive</li> </ul>
Omnipod DASH® Insulin Management System - PODs	POD-BLE-C1-529	<ul style="list-style-type: none"> <li>• Stores personalized settings</li> <li>• Built-in insertion components</li> <li>• Durable, waterproof exterior</li> <li>• Customizable reminders</li> </ul>

### Tandem Diabetes Care Canada Insulin Pumps and Supplies

<b>Insulin Pumps</b>		
<b>Device Name</b>	<b>Model Number</b>	<b>Description</b>
t:slim X2 insulin pump with Control-IQ technology v7.4	1005611	Insulin Pump with 5 year warranty
t:slim X2 insulin pump with Basal-IQ technology v6.4	1006419	Insulin Pump with 5 year warranty
Accessory Kit Included with Pump Purchase (based on pump model)	<p><b>For version 7.4</b> 1005583 EN 1005585 FR</p> <p><b>For version 6.4</b> 1006816 EN 1006730 FR</p>	<p>Accessory Kit includes</p> <ul style="list-style-type: none"> <li>• t:slim™ USB Cable (6ft.)</li> <li>• t:slim Wall Power USB Adapter</li> <li>• t:slim Car Power USB Adapter</li> <li>• Cartridge Removal Tool</li> <li>• Pump Screen Protector</li> <li>• t:case™ Pump Case, Black</li> <li>• User Guide and Instructions (EN or FR)</li> </ul>

<b>Pump Supplies</b>		
<b>Device Name</b>	<b>Model Number</b>	<b>Description / Units or Measure per box</b>
t:slim Cartridge (300 units)	1002541	300 units (3mls) cartridge for insulin, 10 cartridges per box
AutoSoft 90 infusion sets	1002817	6mm cannula infusion set with 23" (60cm) tubing, grey (10 cannula and tubing/box)
	1002818	6mm cannula infusion set with 43" (110cm) tubing, grey (10 cannula and tubing/box)
	1002819	9mm cannula infusion set with 23" (60cm) tubing, grey (10 cannula and tubing/box)
	1002820	9mm cannula infusion set with 43" (110cm) tubing, grey (10 cannula and tubing/box)
	1002821	6mm cannula infusion set with 23" (60cm) tubing, pink (10 cannula and tubing/box)
	1002822	9mm cannula infusion set with 23" (60cm) tubing, pink (10 cannula and tubing/box)
	1002823	6mm cannula infusion set with 23" (60cm) tubing, blue (10 cannula and tubing/box)
	1002824	9mm cannula infusion set with 23" (60cm) tubing, blue (10 cannula and tubing/box)

<b>Pump Supplies (cont'd)</b>		
<b>Device Name</b>	<b>Model Number</b>	<b>Description / Units or Measure per box</b>
AutoSoft 30 infusion sets	1002825	13mm cannula infusion set with 23" (60cm) tubing (10 cannula and tubing/box)
	1002826	13mm cannula infusion set with 23" (110cm) tubing (10 cannula and tubing/box)
VariSoft infusion sets	1002827	13mm cannula infusion set with 23" (60cm) tubing (10 cannula and tubing/box)
	1002828	13mm cannula infusion set with 32" (80cm) tubing (10 cannula and tubing/box)
	1002830	17mm cannula infusion set with 23" (60cm) tubing (10 cannula and tubing/box)
	1002832	17mm cannula infusion set with 43" (110cm) tubing (10 cannula and tubing/box)
TruSteel infusion sets	1002833	6mm needle infusion set with 23" (60cm) tubing (10 needles and tubing/box)
	1002834	6mm needle infusion set with 32" (80cm) tubing (10 needles and tubing/box)
	1002835	8mm needle infusion set, 23" (60cm) tubing (10 needles and tubing/box)
	1002836	8mm needle infusion set, 32" (80cm) tubing (10 needles and tubing/box)
<b>Skin preparation</b>		
<b>Product Name</b>	<b>Model Number</b>	<b>Description / Units or Measure per box</b>
3M Tegaderm Transparent Dressing	RP-1624W	100 dressings per box
Smith & Nephew Skin Prep wipes	MMT-173	50 wipes per box

## **Appendix E Eligible Ostomy Supplies List**

This list details eligible categories, and examples of products within each category. This list may not be exhaustive of all examples within each category.

### **Skin wafers & Pouches**

#### Hollister

- Ceraplus
- New Image
- Premier
- Karaya
- Pouchkins
- Hollihesive

#### Coloplast

- Sensura Mio
- Sensura
- Assura
- Easiflex

#### Convatec

- Natura
- Esteem synergy
- Esteem
- Activelife
- Little Ones
- Salts
- Confidence
- Harmony

### **Adhesive removers**

- Brava
- Wipeaway
- AllKare
- Niltac
- Adapt
- Universal

### **Skin barrier wipes**

- Peri-prep sensitive
- Brava
- AllKare
- Silesse
- Restore

### **Stoma powders, pastes and barrier rings**

- Adapt



- Karaya
- Stomahesive
- Eakin Cohesive
- Stomapaste
- Secuplast
- Brava

**Ostomy belts**

- Ostomy appliance belt
- Adjustable ostomy belt
- Brava
- Adapt

## Appendix F Eligible Diabetes Glucose Sensor Supplies

Sensor (pseudoDIN)	Wear time per sensor	Maximum Annual # of device	Packaged	Annual maximum # of dispenses	Quantity/ Day Supply	Maximum Reimbursable Price
Medtronic Guardian Sensor (3) (97799158)	7 days	55 sensors	5 per box	11 boxes per year	5 sensors every 35 days	79.80 per sensor
Medtronic Guardian Sensor (4) (97798971)	7 days	55 sensors	5 per box	11 boxes per year	5 sensors every 35 days	79.80 per sensor
Dexcom G6 (97799136)	10 days	39 sensors	3 per box	13 boxes per year	3 sensors every 30 days	99.67 per sensor
Dexcom G7 (97798972)	10 days	39 sensors	1 per box	39 boxes per year	3 sensors every 30 days	75.00 per sensor
Libre 2 (97799075)	14 days	26 sensors	1 per box	26 boxes per year	2 sensors every 28 days	90.00 per sensor
Medtronic Guardian Link transmitter for Minimed 670G (97799154)	365 days	1 transmitter	1 per box	1 per year	1 transmitter every 365 days	800 per transmitter
Medtronic Guardian Link transmitter for Minimed 770G (97799071)	365 days	1 transmitter	1 per box	1 per year	1 transmitter every 365 days	800 per transmitter
Medtronic Guardian 4 transmitter for Minimed 780G (97798969)	365 days	1 transmitter	1 per box	1 per year	1 transmitter every 365 days	800 per transmitter
Medtronic Guardian Connect transmitter (97799152)	365 days	1 transmitter	1 per box	1 per year	1 transmitter every 365 days	599 per transmitter
Dexcom G6 Transmitter (97799135)	90 days	4 transmitters	1 per box	4 per year	1 transmitter every 90 days	29 per transmitter

**Optional readers (for Libre) and receivers (for Dexcom) should be acquired by the patients directly through the manufacturer's Customer Care Line.**

**For patients who do not have a compatible smart phone, a G7 receiver may be dispensed through the Diabetes Glucose Sensor Program at no cost to the patient utilizing DIN 97798973. The pharmacy must call the Glucose Sensor Program Administrative Officer at 1-833-335-0538 to have the receiver added to the patients profile if one is needed.**

Household Income Range	Co-payment per dispense period of benefit*
\$0 to \$20,000	\$0.00
\$20,001 to \$40,000	\$10.00
\$41,001 to \$50,000	\$20.00
\$50,001 to \$100,000	\$60.00
\$100,001 or greater	\$80.00

\*co-payment adjustments for those with third-party insurance based on the *Drug Cost Assistance Regulations*