



Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO)

2026 Formulary

List of Covered Drugs

PLEASE READ: THIS DOCUMENT CONTAINS INFORMATION
ABOUT THE DRUGS WE COVER IN THIS PLAN

Formulary ID#: 26314

This formulary was updated on 7/31/2026. For more recent information or other questions, please contact Troy Medicare for Pharmacy Member Service at 1-866-423-8065 (TTY users should call 711), Monday through Sunday, 24 hours a day, or visit <http://www.troymedicare.com>.

Important Message About What You Pay for Vaccines - Our plan covers most Part D vaccines at no cost to you. Call Member Services for more information.

Important Message About What You Pay for Insulin - You won't pay more than \$35 for a one-month supply of each insulin product covered by our plan, no matter what cost-sharing tier it's on. You won't pay more than \$10 for a one-month supply of generic insulin products covered by our plan on Tier 1.

Note to existing members: This formulary has changed since last year. Please review this document to make sure that it still contains the drugs you take. When this drug list (formulary) refers to “we,” “us,” or “our,” it means Troy Health, Inc. When it refers to “plan” or “our plan,” it means Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO). This document includes a list of the drugs (formulary) for our plan which is current as of August 2026. For an updated formulary, please contact us. Our contact information, along with the date we last updated the formulary, appears on the front and back cover pages. You must generally use network pharmacies to use your prescription drug benefit. Benefits, formulary, pharmacy network, and/or copayments/coinsurance may change on January 1, 2026, and from time to time during the year.

What is Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) Formulary?

In this document, we use the terms Drug List and formulary to mean the same thing. A formulary is a list of covered drugs selected by Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) in consultation with a team of health care providers, which represents the prescription therapies believed to be a necessary part of a quality treatment program. Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) will generally cover the drugs listed in our formulary as long as the drug is medically necessary, the prescription is filled at a Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) network pharmacy, and other plan rules are followed. For more information on how to fill your prescriptions, please review your Evidence of Coverage.

Can the Formulary (Drug List) change?

Most changes in drug coverage happen on January 1, but we may add or remove drugs on the Drug List during the year, move them to different cost-sharing tiers, or add new restrictions. We must follow the Medicare rules in making these changes. Updates to the formulary are posted monthly to our website here: www.troymedicare.com/prescription-drugs

Changes that can affect you this year: In the below cases, you will be affected by coverage changes during the year:

- **Immediate substitutions of certain new versions of brand name drugs and original biological products.** We may immediately remove a drug from our formulary if we are replacing it with a certain new version of that drug that will appear on the same or lower cost-sharing tier and with the same or fewer restrictions. When we add a new version of a drug to our formulary, we may decide to keep the brand name drug or original biological product on our formulary, but immediately move it to a different cost-sharing tier or add new restrictions. We can make these immediate changes only if we are adding a new generic version of a brand name drug, or adding certain new biosimilar versions of an original biological product, that was already on the formulary (for example, adding an interchangeable biosimilar that can be substituted for an original biological

product by a pharmacy without a new prescription). If you are currently taking the brand name drug or original biological product, we may not tell you in advance before we make an immediate change, but we will later provide you with information about the specific change(s) we have made. If we make such a change, you or your prescriber can ask us to make an exception and continue to cover for you the drug that is being changed. For more information, see the section below titled **“How do I request an exception to the Troy Medicare’s Formulary?”** Some of these drug types may be new to you. For more information, see the section below titled **“What are original biological products and how are they related to biosimilars?”**

- **Drugs removed from the market.** If a drug is withdrawn from sale by the manufacturer or the Food and Drug Administration (FDA) determines to be withdrawn for safety or effectiveness reasons, we may immediately remove the drug from our formulary and later provide notice to members who take the drug.
- **Other changes.** We may make other changes that affect members currently taking a drug. For instance, we may remove a brand name drug from the formulary when adding a generic equivalent or remove an original biological product when adding a biosimilar. We may also apply new restrictions to the brand name drug or original biological product, or move it to a different cost-sharing tier, or both. We may make changes based on new clinical guidelines. If we remove drugs from our formulary, add prior authorization, quantity limits and/or step therapy restrictions on a drug, or move a drug to a higher cost-sharing tier, we must notify affected members of the change at least 30 days before the change becomes effective. Alternatively, when a member requests a refill of the drug, they may receive a 30-day supply of the drug and notice of the change. If we make these other changes, you or your prescriber can ask us to make an exception for you and continue to cover the drug you have been taking. The notice we provide you will also include information on how to request an exception, and you can also find information in the section below entitled **“How do I request an exception to the Troy Medicare’s Formulary?”**

Changes that will not affect you if you are currently taking the drug. Generally, if you are taking a drug on our 2026 formulary that was covered at the beginning of the year, we will not discontinue or reduce coverage of the drug during the 2026 coverage year except as described above. This means these drugs will remain available at the same cost-sharing and with no new restrictions for those members taking them for the remainder of the coverage year. You will not get direct notice this year about changes that do not affect you. However, on January 1 of the next year, such changes would affect you, and it is important to check the Drug List for the new benefit year for any changes to drugs.

The enclosed formulary is current as of July 2026. To get updated information about the drugs covered by Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO), please contact us. Our contact information appears on the front and

back cover pages. We will update the printable formularies each month and they will be available at <http://www.troymedicare.com/prescription-drugs>.

How do I use the Formulary?

There are two ways to find your drug within the formulary:

Medical Condition

The formulary begins on Page 10. The drugs in this formulary are grouped into categories depending on the type of medical conditions that they are used to treat. For example, drugs used to treat a heart condition are listed under the category, "Cardiovascular Agents - Treatment Of Conditions Affecting The Heart And Blood Vessels". If you know what your drug is used for, look for the category name in the list that begins on Page 10. Then look under the category name for your drug.

Alphabetical Listing

If you are not sure what category to look under, you should look for your drug in the Index that begins on Page 122. The Index provides an alphabetical list of all of the drugs included in this document. Both brand-name drugs and generic drugs are listed in the Index. Look in the Index and find your drug. Next to your drug, you will see the page number where you can find coverage information. Turn to the page listed in the Index and find the name of your drug in the first column of the list.

What are generic drugs?

Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) covers both brand-name drugs and generic drugs. A generic drug is approved by the FDA as having the same active ingredient as the brand-name drug. Generally, generic drugs cost less than brand-name drugs. There are generic drug substitutes available for many brand name drugs. Generic drugs usually can be substituted for the brand name drug at the pharmacy without needing a new prescription, depending on state laws.

What are original biological products and how are they related to biosimilars?

On the formulary, when we refer to drugs, this could mean a drug or a biological product. Biological products are drugs that are more complex than typical drugs. Since biological products are more complex than typical drugs, instead of having a generic form, they have alternatives that are called biosimilars. Generally, biosimilars work just as well as the original biological product and may cost less. There are biosimilar alternatives for some original biological products. Some biosimilars are interchangeable biosimilars and, depending on state laws, may be substituted for the original biological product at the pharmacy without needing a new prescription, just like generic drugs can be substituted for brand name drugs.

For discussion of drug types, please see the Evidence of Coverage, Chapter 5, Section 3.1, "The 'Drug List' tells which Part D drugs are covered."]

Are there any restrictions on my coverage?

Some covered drugs may have additional requirements or limits on coverage. These requirements and limits may include:

- **Prior Authorization (PA):** Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) requires you or your physician to get prior authorization for certain drugs. This means that you will need to get approval from us before you fill your prescriptions. If you do not get approval, we may not cover the drug.
- **Quantity Limits (QL):** For certain drugs, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) limits the amount of the drug that we will cover. For example, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) provides up to twelve (12) capsules per prescription for *gabapentin oral capsule 300 mg* per day. This may be in addition to a standard one-month or three-month supply.
- **Step Therapy (ST):** In some cases, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) requires you to first try certain drugs to treat your medical condition before we will cover another drug for that condition. For example, if Drug A and Drug B both treat your medical condition, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) may not cover Drug B unless you try Drug A first. If Drug A does not work for you, we will then cover Drug B.

You can find out if your drug has any additional requirements or limits by looking in the formulary that begins on Page 10. You can also get more information about the restrictions applied to specific covered drugs by visiting our website. We have posted online documents that explain our prior authorization and step therapy restrictions. You may also ask us to send you a copy. Our contact information, along with the date we last updated the formulary, appears on the front and back cover pages.

You can ask us to make an exception to these restrictions or limits or for a list of other, similar drugs that may treat your health condition. See the section, “**How do I request an exception to the Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) Formulary?**” on Page 6 for information about how to request an exception.

What if my drug is not on the Formulary?

If your drug is not included in this formulary (list of covered drugs), you should first contact Pharmacy Member Services and ask if your drug is covered.

If you learn that Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) does not cover your drug, you have two options:

- You can ask Member Services for a list of similar drugs that are covered by our plan. When you receive the list, show it to your doctor and ask them to prescribe a similar drug that is covered by Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP).

- You can ask Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) to make an exception and cover your drug. See below for information about how to request an exception.

How do I request an exception to the Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) Formulary?

You can ask us to make an exception to our coverage rules. There are several types of exceptions that you can ask us to make.

- You can ask us to cover a drug even if it is not on our formulary. If approved, this drug will be covered at a pre-determined cost-sharing level, and you would not be able to ask us to provide the drug at a lower cost-sharing level.
- You can ask us to waive coverage restrictions or limits on your drug. For example, for certain drugs, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) limits the amount of the drug that we will cover. If your drug has a quantity limit, you can ask us to waive the limit and cover a greater amount.
- You can ask us to cover a formulary drug at a lower cost-sharing level unless the drug is on the specialty tier. If approved, this would lower the amount you must pay for your drug.

Generally, we will only approve your request for an exception if the alternative drugs included on the plan's formulary, the lower cost-sharing drug or additional utilization restrictions would not be as effective in treating your condition and/or would cause you to have adverse medical effects.

You or your prescriber should contact us to ask for a tiering or formulary exception, including an exception to a coverage restriction. **When you request an exception, your prescriber will need to explain the medical reasons why you need the exception.**

Generally, we must make our decision within 72 hours of getting your prescriber's supporting statement. You can ask for an expedited (fast) decision if you believe, and we agree, that your health could be seriously harmed by waiting up to 72 hours for a decision. If we agree, or if your prescriber asks for a fast decision, we must give you a decision no later than 24 hours after we get your prescriber's supporting statement.

What can I do if my drug is not on the formulary or has a restriction?

As a new or continuing member in our plan you may be taking drugs that are not on our formulary. Or, you may be taking a drug that is on our formulary but has a coverage restriction, such as prior authorization. You should talk to your prescriber about requesting a coverage decision to show that you meet the criteria for approval, switching to an alternative drug that we cover, or requesting a formulary exception so that we will cover the drug you take. While you and your doctor determine the right course of action for you, we may cover your drug in certain cases during the first 90 days you are a member of our plan.

For each of your drugs that is not on our formulary or has a coverage restriction, we will cover a temporary 30-day supply. If your prescription is written for fewer days, we'll allow refills to provide up to a maximum 30-day supply of medication. If coverage is not approved, after your first 30-day supply, we will not pay for these drugs, even if you have been a member of the plan less than 90 days.

If you are a resident of a long-term care facility and you need a drug that is not on our formulary or if your ability to get your drugs is limited, but you are past the first 90 days of membership in our plan, we will cover a 31-day emergency supply of that drug while you pursue a formulary exception.

For More Information

For more detailed information about your Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO) prescription drug coverage, please review your Evidence of Coverage and other plan materials.

If you have questions about Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare (HMO), please contact us. Our contact information, along with the date we last updated the formulary, appears on the front and back cover pages.

If you have general questions about Medicare prescription drug coverage, please call Medicare at 1-800-MEDICARE (1-800-633-4227) 24 hours a day / 7 days a week. TTY users should call 1-877-486-2048. Or, visit <http://www.medicare.gov>.

Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy Medicare Formulary

The formulary below provides coverage information about the drugs covered by Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP). If you have trouble finding your drug in the list, turn to the Index that begins on Page 122.

The first column of the chart lists the drug name. Brand-name drugs are capitalized (e.g., TRESIBA FLEXTOUCH SUBCUTANEOUS SOLUTION PEN-INJECTOR 200 UNIT/ML) and generic drugs are listed in lower-case italics (e.g., *insulin lispro (1 unit dial) subcutaneous solution pen-injector 100 unit/ml*).

The information in the Requirements/Limits column tells you if Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) has special requirements for coverage of your drug.

- **Prior Authorization (PA):** Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) requires you or your physician to get prior authorization for certain drugs. This means that you will need to get approval from us before you fill your prescriptions. If you do not get approval, we may not cover the drug.
- **Quantity Limits (QL):** For certain drugs, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) limits the amount of the drug that the plan will cover. Each quantity limit is defined in our formulary below as a quantity of each (EA) dosage form (e.g., capsule, patch, tablet, etc.) or milliliters (mL) for solutions or

suspensions that we will cover within a specific time period. For example, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) provides 360 per prescription for *gabapentin oral capsule 300 mg* every 30 days.

- **Step Therapy (ST):** In some cases, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) requires you to first try certain drugs to treat your medical condition before we will cover another drug for that condition. For example, if Drug A and Drug B both treat your medical condition, Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) may not cover Drug B unless you try Drug A first. If Drug A does not work for you, we will then cover Drug B.
- **Medicare Part B or Part D (B/D):** Depending on how this drug is used, it may be covered by either Medicare Part B (doctor and outpatient health care) or Medicare Part D (prescription drugs). Your doctor may need to provide the plan with more information about how this drug will be used to make sure it is correctly covered by Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP).
- **Morphine Milligram Equivalent (MME):** Additional quantity limits may apply across all drugs in the opioid class used for the treatment of pain. This additional limit is called a cumulative morphine milligram equivalent (MME) and is designed to monitor safe dosing levels of opioids for individuals who may be taking more than 1 opioid drug for pain management. If your prescriber prescribes more than this amount or thinks the limit is not right for your situation, you or your prescriber can ask the plan to cover the additional quantity.



Plans are offered through Troy Medicare, a Medicare Advantage HMO and HMO D-SNP organization with a Medicare contract. Enrollment in these plans depends on the plan's contract renewal with Medicare. Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP) also has a contract with state Medicaid.

The Formulary may change at any time. You will receive notice when necessary.

Benefits, formulary, pharmacy network, provider network, premium and/or copay/coinsurance may change on January 1 of each year. Member premiums, copays, coinsurance, and deductibles may vary based on the level of "Extra Help" you receive. Please contact the plan for further details.

This information is available for free in other languages and other formats, such as Braille and large print. Please call our Member Services number at 1-888-494-TROY (8769). TTY users should call 711. During the months of April through September, we are available from 8:00 am to 8:00 pm (ET), Monday through Friday. During the months of October through March, we are available from 8:00 am to 8:00 pm (ET), seven (7) days a week. Member Services also has free language interpreter services available for non-English speakers.

You must generally use network pharmacies to use your prescription drug benefit.

Troy does not discriminate or exclude people because of their race, color, national origin, ancestry, age, disability, ethnicity, sex, sexual orientation, gender, gender identity or expression, marital status, religion, or language.

2026 Troy Medicare

2026 Member Formulary

Formulary ID 26314

CURRENT AS OF 8/1/2026

Name of Drug	Drug Tier	Requirements/Limits
Analgesics - Treatment Of Pain		
Analgesics		
BAC (BUTALBITAL-ACETAMIN-CAFF) ORAL TABLET 50-325-40 MG	2	PA
<i>butalbital-acetaminophen oral tablet 50-325 mg</i>	2	PA
<i>butalbital-apap-caff-cod oral capsule 50-325-40-30 mg</i>	2	PA; MME
<i>butalbital-apap-caffeine oral capsule 50-325-40 mg</i>	2	PA
<i>butalbital-apap-caffeine oral solution 50-325-40 mg/15ml</i>	2	PA
<i>butalbital-apap-caffeine oral tablet 50-325-40 mg</i>	2	PA
<i>butalbital-asa-caff-codeine oral capsule 50-325-40-30 mg</i>	2	PA; MME
<i>butalbital-aspirin-caffeine oral capsule 50-325-40 mg</i>	2	PA
<i>nalbuphine hcl injection solution 10 mg/ml</i>	2	MME
Nonsteroidal Anti-Inflammatory Drugs		
<i>celecoxib oral capsule 100 mg, 200 mg, 400 mg, 50 mg</i>	1	
<i>diclofenac epolamine external patch 1.3 %</i>	2	
<i>diclofenac potassium oral tablet 50 mg</i>	2	
<i>diclofenac sodium er oral tablet extended release 24 hour 100 mg</i>	1	
<i>diclofenac sodium external gel 3 %</i>	2	
<i>diclofenac sodium external solution 1.5 %</i>	2	
<i>diclofenac sodium oral tablet delayed release 25 mg, 50 mg, 75 mg</i>	1	
<i>diflunisal oral tablet 500 mg</i>	2	
<i>etodolac er oral tablet extended release 24 hour 400 mg, 500 mg, 600 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>etodolac oral capsule 200 mg, 300 mg</i>	2	
<i>etodolac oral tablet 400 mg, 500 mg</i>	2	
<i>flurbiprofen oral tablet 100 mg</i>	2	
IBU ORAL TABLET 400 MG, 600 MG, 800 MG	1	
<i>ibuprofen oral suspension 100 mg/5ml</i>	1	
<i>ibuprofen oral tablet 400 mg, 600 mg, 800 mg</i>	1	
<i>indomethacin er oral capsule extended release 75 mg</i>	2	
<i>indomethacin oral capsule 25 mg, 50 mg</i>	2	PA
<i>ketorolac tromethamine oral tablet 10 mg</i>	2	PA; QL (20 EA per 30 days)
<i>meclofenamate sodium oral capsule 100 mg, 50 mg</i>	2	
<i>meloxicam oral tablet 15 mg, 7.5 mg</i>	1	
<i>nabumetone oral tablet 500 mg, 750 mg</i>	2	
<i>naproxen dr oral tablet delayed release 500 mg</i>	3	
<i>naproxen oral suspension 125 mg/5ml</i>	2	
<i>naproxen oral tablet 250 mg, 375 mg, 500 mg</i>	1	
<i>naproxen oral tablet delayed release 375 mg, 500 mg</i>	3	
<i>naproxen sodium oral tablet 275 mg, 550 mg</i>	2	
<i>piroxicam oral capsule 10 mg, 20 mg</i>	2	
<i>sulindac oral tablet 150 mg, 200 mg</i>	2	
Opioid Analgesics, Long-Acting		
<i>buprenorphine transdermal patch weekly 10 mcg/hr, 15 mcg/hr, 20 mcg/hr, 5 mcg/hr, 7.5 mcg/hr</i>	2	QL (4 EA per 28 days)
<i>fentanyl transdermal patch 72 hour 100 mcg/hr, 12 mcg/hr, 25 mcg/hr, 37.5 mcg/hr, 50 mcg/hr, 75 mcg/hr</i>	2	MME; QL (10 EA per 30 days)
<i>fentanyl transdermal patch 72 hour 62.5 mcg/hr, 87.5 mcg/hr</i>	4	MME; QL (10 EA per 30 days)
<i>methadone hcl oral solution 10 mg/5ml</i>	2	MME; QL (1200 ML per 30 days)
<i>methadone hcl oral solution 5 mg/5ml</i>	2	MME; QL (2400 ML per 30 days)
<i>methadone hcl oral tablet 10 mg</i>	2	MME; QL (240 EA per 30 days)
<i>methadone hcl oral tablet 5 mg</i>	2	MME; QL (180 EA per 30 days)
<i>morphine sulfate er oral tablet extended release 100 mg, 15 mg, 200 mg, 30 mg, 60 mg</i>	2	MME; QL (60 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
OXYCONTIN ORAL TABLET ER 12 HOUR ABUSE-DETERRENT 10 MG, 15 MG, 20 MG, 30 MG, 40 MG	4	PA; MME; QL (90 EA per 30 days)
OXYCONTIN ORAL TABLET ER 12 HOUR ABUSE-DETERRENT 60 MG, 80 MG	4	PA; MME; QL (60 EA per 30 days)
Opioid Analgesics, Short-Acting		
<i>acetaminophen-codeine oral solution 120-12 mg/5ml</i>	2	MME
<i>acetaminophen-codeine oral tablet 300-15 mg, 300-30 mg, 300-60 mg</i>	2	MME
<i>butorphanol tartrate nasal solution 10 mg/ml</i>	2	MME; QL (5 ML per 30 days)
ENDOCET ORAL TABLET 10-325 MG, 2.5-325 MG, 5-325 MG, 7.5-325 MG	2	MME
<i>hydrocodone-acetaminophen oral tablet 10-325 mg, 5-325 mg, 7.5-325 mg</i>	2	MME
<i>hydrocodone-ibuprofen oral tablet 10-200 mg, 5-200 mg, 7.5-200 mg</i>	2	MME
<i>hydromorphone hcl oral tablet 2 mg, 4 mg, 8 mg</i>	2	MME; QL (120 EA per 30 days)
<i>hydromorphone hcl pf injection solution 1 mg/ml, 10 mg/ml, 4 mg/ml, 50 mg/5ml, 500 mg/50ml</i>	2	MME
<i>morphine sulfate (concentrate) oral solution 100 mg/5ml</i>	2	MME
<i>morphine sulfate oral tablet 15 mg, 30 mg</i>	2	MME; QL (120 EA per 30 days)
<i>oxycodone hcl oral solution 5 mg/5ml</i>	2	MME; QL (5400 ML per 30 days)
<i>oxycodone hcl oral tablet 10 mg, 15 mg, 20 mg, 30 mg, 5 mg</i>	2	MME; QL (120 EA per 30 days)
<i>oxycodone hcl oral tablet abuse-deterrent 15 mg</i>	2	MME; QL (120 EA per 30 days)
<i>oxycodone-acetaminophen oral tablet 10-325 mg, 5-325 mg, 7.5-325 mg</i>	3	MME
<i>oxycodone-acetaminophen oral tablet 2.5-325 mg</i>	2	MME
<i>pentazocine-naloxone hcl oral tablet 50-0.5 mg</i>	2	MME
<i>tramadol hcl oral tablet 50 mg</i>	2	MME; QL (240 EA per 30 days)
<i>tramadol-acetaminophen oral tablet 37.5-325 mg</i>	2	MME
Anesthetics - Local Treatment Of Pain		
Local Anesthetics		
<i>lidocaine external ointment 5 %</i>	2	QL (50 GM per 30 days)
<i>lidocaine external patch 5 %</i>	2	PA; QL (90 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>lidocaine hcl external solution 4 %</i>	2	
<i>lidocaine viscous hcl mouth/throat solution 2 %</i>	2	
<i>lidocaine-prilocaine external cream 2.5-2.5 %</i>	2	
ZTLIDO EXTERNAL PATCH 1.8 %	4	PA; QL (90 EA per 30 days)
Anti-Addiction/Substance Abuse Treatment Agents - Treatment Of Substance Abuse Disorders		
Alcohol Deterrents/Anti-Craving		
<i>acamprosate calcium oral tablet delayed release 333 mg</i>	2	
<i>disulfiram oral tablet 250 mg, 500 mg</i>	2	
VIVITROL INTRAMUSCULAR SUSPENSION RECONSTITUTED 380 MG	5	QL (1 EA per 28 days)
Opioid Dependence		
<i>buprenorphine hcl sublingual tablet sublingual 2 mg, 8 mg</i>	2	
<i>buprenorphine hcl-naloxone hcl sublingual film 12-3 mg</i>	2	QL (90 EA per 30 days)
<i>buprenorphine hcl-naloxone hcl sublingual film 2-0.5 mg</i>	2	QL (150 EA per 30 days)
<i>buprenorphine hcl-naloxone hcl sublingual film 4-1 mg, 8-2 mg</i>	2	QL (120 EA per 30 days)
<i>buprenorphine hcl-naloxone hcl sublingual tablet sublingual 2-0.5 mg, 8-2 mg</i>	2	QL (120 EA per 30 days)
<i>lofexidine hcl oral tablet 0.18 mg</i>	5	PA; QL (224 EA per 14 days)
<i>naloxone hcl injection solution prefilled syringe 0.4 mg/ml</i>	1	
<i>naltrexone hcl oral tablet 50 mg</i>	1	
ZURNAI INJECTION SOLUTION AUTO-INJECTOR 1.5 MG/0.5ML	3	
Opioid Reversal Agents		
KLOXXADO NASAL LIQUID 8 MG/0.1ML	3	
<i>naloxone hcl injection solution 0.4 mg/ml, 4 mg/10ml</i>	1	
<i>naloxone hcl injection solution cartridge 0.4 mg/ml</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>naloxone hcl injection solution prefilled syringe 2 mg/2ml</i>	1	
OPVEE NASAL SOLUTION 2.7 MG/0.1ML	3	
REXTOVY NASAL LIQUID 4 MG/0.25ML	3	
REZENOPY NASAL LIQUID 10 MG/0.11ML	6	
Smoking Cessation Agents		
<i>bupropion hcl er (smoking det) oral tablet extended release 12 hour 150 mg</i>	1	
NICOTROL NS NASAL SOLUTION 10 MG/ML	4	
<i>varenicline tartrate (starter) oral tablet therapy pack 0.5 mg x 11 & 1 mg x 42</i>	2	QL (56 EA per 28 days)
<i>varenicline tartrate oral tablet 0.5 mg, 1 mg, 1 mg (56 pack)</i>	2	QL (56 EA per 28 days)
<i>varenicline tartrate(continue) oral tablet 1 mg</i>	2	QL (56 EA per 28 days)
Antibacterials - Treatment Of Bacterial Infections		
Aminoglycosides		
<i>amikacin sulfate injection solution 500 mg/2ml</i>	2	
ARIKAYCE INHALATION SUSPENSION 590 MG/8.4ML	5	PA
<i>gentamicin in saline intravenous solution 0.8-0.9 mg/ml-%, 1-0.9 mg/ml-%, 1.2-0.9 mg/ml-%, 1.6-0.9 mg/ml-%, 2-0.9 mg/ml-%</i>	2	
<i>gentamicin sulfate injection solution 40 mg/ml</i>	2	
<i>neomycin sulfate oral tablet 500 mg</i>	2	
<i>streptomycin sulfate intramuscular solution reconstituted 1 gm</i>	4	
<i>tobramycin sulfate injection solution 1.2 gm/30ml, 10 mg/ml, 2 gm/50ml, 80 mg/2ml</i>	2	
<i>tobramycin sulfate injection solution reconstituted 1.2 gm</i>	2	
Antibacterials, Other		
<i>aztreonam injection solution reconstituted 1 gm, 2 gm</i>	2	
<i>clindamycin hcl oral capsule 150 mg, 300 mg, 75 mg</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>clindamycin palmitate hcl oral solution reconstituted 75 mg/5ml</i>	2	
<i>clindamycin phosphate in d5w intravenous solution 300 mg/50ml, 600 mg/50ml, 900 mg/50ml</i>	2	
<i>clindamycin phosphate in nacl intravenous solution 300-0.9 mg/50ml-%, 600-0.9 mg/50ml-%, 900-0.9 mg/50ml-%</i>	2	
<i>clindamycin phosphate injection solution 300 mg/2ml, 900 mg/6ml</i>	2	
<i>clindamycin phosphate vaginal cream 2 %</i>	2	
<i>colistimethate sodium (cba) injection solution reconstituted 150 mg</i>	2	
<i>daptomycin intravenous solution reconstituted 350 mg</i>	2	
<i>daptomycin intravenous solution reconstituted 500 mg</i>	4	
<i>fosfomycin tromethamine oral packet 3 gm</i>	2	
<i>linezolid intravenous solution 600 mg/300ml</i>	2	
<i>linezolid oral suspension reconstituted 100 mg/5ml</i>	5	
<i>linezolid oral tablet 600 mg</i>	2	
<i>methenamine hippurate oral tablet 1 gm</i>	2	
<i>metronidazole intravenous solution 500 mg/100ml</i>	2	
<i>metronidazole oral capsule 375 mg</i>	2	
<i>metronidazole oral tablet 250 mg, 500 mg</i>	1	
<i>metronidazole vaginal gel 0.75 %</i>	2	
<i>nitrofurantoin macrocrystal oral capsule 100 mg, 25 mg, 50 mg</i>	2	
<i>nitrofurantoin monohyd macro oral capsule 100 mg</i>	2	
<i>polymyxin b sulfate injection solution reconstituted 500000 unit</i>	2	
PRIMAXIN IV INTRAVENOUS SOLUTION RECONSTITUTED 500-500 MG	4	
<i>sulfamethoxazole-trimethoprim intravenous solution 400-80 mg/5ml</i>	2	
<i>tigecycline intravenous solution reconstituted 50 mg</i>	4	PA

Name of Drug	Drug Tier	Requirements/Limits
<i>tinidazole oral tablet 250 mg, 500 mg</i>	2	
<i>trimethoprim oral tablet 100 mg</i>	1	
TYZAVAN INTRAVENOUS SOLUTION 1000 MG/200ML, 1250 MG/250ML, 1500 MG/300ML, 1750 MG/350ML, 2000 MG/400ML, 500 MG/100ML, 750 MG/150ML	4	
<i>vancomycin hcl in nacl intravenous solution 1-0.9 gm/200ml-%, 500-0.9 mg/100ml-%, 750-0.9 mg/150ml-%</i>	2	
<i>vancomycin hcl intravenous solution 1000 mg/200ml, 1250 mg/250ml, 1500 mg/300ml, 1750 mg/350ml, 2000 mg/400ml, 500 mg/100ml, 750 mg/150ml</i>	2	
<i>vancomycin hcl intravenous solution reconstituted 1 gm, 1.25 gm, 1.5 gm, 1.75 gm, 10 gm, 2 gm, 5 gm, 500 mg, 750 mg</i>	2	
<i>vancomycin hcl oral capsule 125 mg, 250 mg</i>	2	
ZOSYN INTRAVENOUS SOLUTION 2-0.25 GM/50ML	4	
Beta-Lactam, Cephalosporins		
<i>cefaclor er oral tablet extended release 12 hour 500 mg</i>	2	
<i>cefaclor oral capsule 250 mg, 500 mg</i>	2	
<i>cefadroxil oral capsule 500 mg</i>	1	
<i>cefadroxil oral suspension reconstituted 250 mg/5ml, 500 mg/5ml</i>	2	
<i>cefadroxil oral tablet 1 gm</i>	2	
<i>cefazolin sodium injection solution reconstituted 1 gm, 2 gm, 3 gm, 500 mg</i>	2	
<i>cefazolin sodium intravenous solution reconstituted 1 gm, 2 gm, 3 gm</i>	2	
<i>cefdinir oral capsule 300 mg</i>	1	
<i>cefdinir oral suspension reconstituted 125 mg/5ml, 250 mg/5ml</i>	2	
<i>cefepime hcl injection solution reconstituted 1 gm</i>	2	
<i>cefepime hcl intravenous solution 1 gm/50ml, 2 gm/100ml</i>	2	
<i>cefepime hcl intravenous solution reconstituted 2 gm</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>cefepime-dextrose intravenous solution reconstituted 1-5 gm-%(50ml), 2-5 gm-%(50ml)</i>	2	
<i>cefixime oral capsule 400 mg</i>	2	
<i>cefoxitin sodium intravenous solution reconstituted 1 gm, 10 gm, 2 gm</i>	2	
<i>cefpodoxime proxetil oral suspension reconstituted 100 mg/5ml, 50 mg/5ml</i>	2	
<i>cefpodoxime proxetil oral tablet 100 mg, 200 mg</i>	2	
<i>cefprozil oral suspension reconstituted 125 mg/5ml, 250 mg/5ml</i>	2	
<i>cefprozil oral tablet 250 mg, 500 mg</i>	2	
<i>ceftaroline fosamil intravenous solution reconstituted 400 mg, 600 mg</i>	5	
<i>ceftazidime injection solution reconstituted 1 gm, 6 gm</i>	2	
<i>ceftazidime intravenous solution reconstituted 2 gm</i>	2	
<i>ceftriaxone sodium in dextrose intravenous solution 20 mg/ml, 40 mg/ml</i>	2	
<i>ceftriaxone sodium injection solution reconstituted 1 gm, 2 gm, 250 mg, 500 mg</i>	2	
<i>ceftriaxone sodium intravenous solution reconstituted 1 gm, 10 gm, 2 gm</i>	2	
<i>ceftriaxone sodium-dextrose intravenous solution reconstituted 1-3.74 gm-%(50ml), 2-2.22 gm-%(50ml)</i>	2	
<i>cefuroxime axetil oral tablet 250 mg, 500 mg</i>	2	
<i>cefuroxime sodium injection solution reconstituted 750 mg</i>	2	
<i>cefuroxime sodium intravenous solution reconstituted 1.5 gm</i>	2	
<i>cephalexin oral capsule 250 mg, 500 mg</i>	1	
<i>cephalexin oral suspension reconstituted 125 mg/5ml, 250 mg/5ml</i>	1	
<i>cephalexin oral tablet 250 mg, 500 mg</i>	2	
TAZICEF INJECTION SOLUTION RECONSTITUTED 1 GM	4	
TAZICEF INTRAVENOUS SOLUTION RECONSTITUTED 1 GM, 2 GM, 6 GM	4	

Name of Drug	Drug Tier	Requirements/Limits
TEFLARO INTRAVENOUS SOLUTION RECONSTITUTED 400 MG, 600 MG	5	
Beta-Lactam, Penicillins		
<i>amoxicillin oral capsule 250 mg, 500 mg</i>	1	
<i>amoxicillin oral suspension reconstituted 125 mg/5ml, 200 mg/5ml, 250 mg/5ml, 400 mg/5ml</i>	1	
<i>amoxicillin oral tablet 500 mg, 875 mg</i>	1	
<i>amoxicillin oral tablet chewable 125 mg, 250 mg</i>	1	
<i>amoxicillin-pot clavulanate er oral tablet extended release 12 hour 1000-62.5 mg</i>	2	
<i>amoxicillin-pot clavulanate oral suspension reconstituted 200-28.5 mg/5ml, 250-62.5 mg/5ml, 400-57 mg/5ml, 600-42.9 mg/5ml</i>	2	
<i>amoxicillin-pot clavulanate oral tablet 250-125 mg, 500-125 mg, 875-125 mg</i>	2	
<i>ampicillin oral capsule 500 mg</i>	1	
<i>ampicillin sodium injection solution reconstituted 1 gm, 2 gm</i>	2	
<i>ampicillin sodium intravenous solution reconstituted 1 gm, 10 gm, 2 gm</i>	2	
<i>ampicillin-sulbactam sodium injection solution reconstituted 1.5 (1-0.5) gm, 3 (2-1) gm</i>	2	
<i>ampicillin-sulbactam sodium intravenous solution reconstituted 1.5 (1-0.5) gm, 15 (10-5) gm, 3 (2-1) gm</i>	2	
BICILLIN L-A INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 1200000 UNIT/2ML, 2400000 UNIT/4ML, 600000 UNIT/ML	4	
<i>dicloxacillin sodium oral capsule 250 mg, 500 mg</i>	2	
<i>nafcillin sodium injection solution reconstituted 1 gm</i>	2	
<i>nafcillin sodium injection solution reconstituted 2 gm</i>	4	
<i>oxacillin sodium in dextrose intravenous solution 2 gm/50ml</i>	2	
<i>penicillin g pot in dextrose intravenous solution 40000 unit/ml, 60000 unit/ml</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>penicillin g sodium injection solution reconstituted 5000000 unit</i>	2	
<i>penicillin v potassium oral solution reconstituted 125 mg/5ml, 250 mg/5ml</i>	1	
<i>penicillin v potassium oral tablet 250 mg, 500 mg</i>	1	
<i>piperacillin sod-tazobactam so intravenous solution reconstituted 13.5 (12-1.5) gm, 2.25 (2-0.25) gm, 3-0.375 gm, 3.375 (3-0.375) gm, 4.5 (4-0.5) gm, 40.5 (36-4.5) gm</i>	2	
<i>piperacillin-tazobactam-nacl intravenous solution reconstituted 2-0.25 gm/50ml, 3-0.375 gm/50ml, 4-0.5 gm/100ml</i>	2	
Carbapenems		
<i>ertapenem sodium injection solution reconstituted 1 gm</i>	4	
<i>imipenem-cilastatin intravenous solution reconstituted 250 mg, 500 mg</i>	2	
<i>meropenem intravenous solution reconstituted 1 gm, 500 mg</i>	2	
<i>meropenem-sodium chloride intravenous solution reconstituted 1 gm/50ml, 500 mg/50ml</i>	2	
Macrolides		
<i>azithromycin intravenous solution reconstituted 500 mg</i>	2	
<i>azithromycin oral suspension reconstituted 100 mg/5ml, 200 mg/5ml</i>	2	
<i>azithromycin oral tablet 250 mg, 250 mg (6 pack), 500 mg, 500 mg (3 pack), 600 mg</i>	1	
<i>clarithromycin er oral tablet extended release 24 hour 500 mg</i>	2	
<i>clarithromycin oral suspension reconstituted 125 mg/5ml, 250 mg/5ml</i>	2	
<i>clarithromycin oral tablet 250 mg, 500 mg</i>	2	
DIFICID ORAL SUSPENSION RECONSTITUTED 40 MG/ML	5	
ERYTHROCIN LACTOBIONATE INTRAVENOUS SOLUTION RECONSTITUTED 500 MG	4	
<i>erythromycin base oral tablet 250 mg, 500 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>erythromycin ethylsuccinate oral suspension reconstituted 200 mg/5ml</i>	2	
<i>erythromycin ethylsuccinate oral tablet 400 mg</i>	2	
<i>fidaxomicin oral tablet 200 mg</i>	5	
ZITHROMAX INTRAVENOUS SOLUTION RECONSTITUTED 500 MG	4	
Quinolones		
<i>ciprofloxacin hcl oral tablet 250 mg, 500 mg, 750 mg</i>	1	
<i>ciprofloxacin in d5w intravenous solution 200 mg/100ml</i>	2	
<i>levofloxacin in d5w intravenous solution 500 mg/100ml, 750 mg/150ml</i>	2	
<i>levofloxacin intravenous solution 25 mg/ml</i>	2	
<i>levofloxacin oral solution 25 mg/ml</i>	2	
<i>levofloxacin oral tablet 250 mg, 500 mg, 750 mg</i>	1	
<i>moxifloxacin hcl in nacl intravenous solution 400 mg/250ml</i>	2	
<i>moxifloxacin hcl intravenous solution 400 mg/250ml</i>	2	
<i>moxifloxacin hcl oral tablet 400 mg</i>	2	
<i>ofloxacin oral tablet 300 mg, 400 mg</i>	2	
Sulfonamides		
<i>sulfacetamide sodium (acne) external lotion 10 %</i>	2	
<i>sulfadiazine oral tablet 500 mg</i>	2	
<i>sulfamethoxazole-trimethoprim oral suspension 200-40 mg/5ml</i>	2	
<i>sulfamethoxazole-trimethoprim oral tablet 400-80 mg, 800-160 mg</i>	1	
Tetracyclines		
DOXY 100 INTRAVENOUS SOLUTION RECONSTITUTED 100 MG	2	
<i>doxycycline hyclate intravenous solution reconstituted 100 mg</i>	2	
<i>doxycycline hyclate oral capsule 100 mg, 50 mg</i>	2	
<i>doxycycline hyclate oral tablet 100 mg, 20 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>doxycycline monohydrate oral capsule 100 mg, 50 mg</i>	1	
<i>doxycycline monohydrate oral tablet 100 mg, 150 mg, 50 mg, 75 mg</i>	2	
<i>minocycline hcl oral capsule 100 mg, 50 mg, 75 mg</i>	1	
<i>minocycline hcl oral tablet 100 mg, 50 mg, 75 mg</i>	2	
<i>tetracycline hcl oral capsule 250 mg, 500 mg</i>	2	
Anticonvulsants - Treatment Of Seizures		
Anticonvulsants, Other		
<i>brivaracetam oral solution 10 mg/ml</i>	2	QL (600 ML per 30 days)
<i>brivaracetam oral tablet 10 mg, 100 mg, 25 mg, 50 mg, 75 mg</i>	2	QL (60 EA per 30 days)
DIACOMIT ORAL CAPSULE 250 MG, 500 MG	5	PA
DIACOMIT ORAL PACKET 250 MG, 500 MG	5	PA
<i>divalproex sodium er oral tablet extended release 24 hour 250 mg, 500 mg</i>	2	
<i>divalproex sodium oral capsule delayed release sprinkle 125 mg</i>	2	
<i>divalproex sodium oral tablet delayed release 125 mg, 250 mg, 500 mg</i>	2	
EPIDIOLEX ORAL SOLUTION 100 MG/ML	5	PA
<i>felbamate oral suspension 600 mg/5ml</i>	2	
<i>felbamate oral tablet 400 mg, 600 mg</i>	2	
FINTEPLA ORAL SOLUTION 2.2 MG/ML	5	PA
<i>lamotrigine er oral tablet extended release 24 hour 100 mg, 200 mg, 25 mg, 250 mg, 300 mg, 50 mg</i>	2	
<i>lamotrigine oral tablet 100 mg, 150 mg, 200 mg, 25 mg</i>	1	
<i>lamotrigine oral tablet chewable 25 mg, 5 mg</i>	2	
<i>lamotrigine starter kit-blue oral kit 35 x 25 mg</i>	2	
<i>lamotrigine starter kit-green oral kit 84 x 25 mg & 14x100 mg</i>	4	
<i>lamotrigine starter kit-orange oral kit 42 x 25 mg & 7 x 100 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>levetiracetam er oral tablet extended release 24 hour 500 mg, 750 mg</i>	1	
<i>levetiracetam oral solution 100 mg/ml, 500 mg/5ml</i>	1	
<i>levetiracetam oral tablet 1000 mg, 250 mg, 500 mg, 750 mg</i>	1	
<i>levetiracetam oral tablet disintegrating soluble 250 mg, 500 mg</i>	2	ST; QL (60 EA per 30 days)
<i>perampanel oral suspension 0.5 mg/ml</i>	5	ST; QL (720 ML per 30 days)
<i>perampanel oral tablet 10 mg, 12 mg, 4 mg, 6 mg, 8 mg</i>	5	ST; QL (30 EA per 30 days)
<i>perampanel oral tablet 2 mg</i>	4	ST; QL (30 EA per 30 days)
SPRITAM ORAL TABLET DISINTEGRATING SOLUBLE 250 MG, 500 MG	4	ST; QL (60 EA per 30 days)
SUBVENITE ORAL SUSPENSION 10 MG/ML	4	
<i>topiramate oral capsule sprinkle 15 mg, 25 mg, 50 mg</i>	2	
<i>topiramate oral solution 25 mg/ml</i>	2	PA
<i>topiramate oral tablet 100 mg, 200 mg, 25 mg, 50 mg</i>	1	
<i>valproic acid oral capsule 250 mg</i>	2	
<i>valproic acid oral solution 250 mg/5ml</i>	2	
XCOPRI (250 MG DAILY DOSE) ORAL TABLET THERAPY PACK 100 & 150 MG	4	ST
XCOPRI (350 MG DAILY DOSE) ORAL TABLET THERAPY PACK 150 & 200 MG	4	ST
XCOPRI ORAL TABLET 100 MG, 150 MG, 200 MG, 25 MG, 50 MG	4	ST
XCOPRI ORAL TABLET THERAPY PACK 14 X 12.5 MG & 14 X 25 MG, 14 X 150 MG & 14 X 200 MG, 14 X 50 MG & 14 X 100 MG	4	ST
Calcium Channel Modifying Agents		
<i>ethosuximide oral capsule 250 mg</i>	2	
<i>ethosuximide oral solution 250 mg/5ml</i>	2	
<i>methsuximide oral capsule 300 mg</i>	2	
Gamma-Aminobutyric Acid (Gaba) Augmenting Agents		
<i>clobazam oral suspension 2.5 mg/ml</i>	2	QL (480 ML per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>clobazam oral tablet 10 mg, 20 mg</i>	2	QL (60 EA per 30 days)
<i>diazepam rectal gel 10 mg, 2.5 mg, 20 mg</i>	2	
<i>gabapentin oral capsule 100 mg, 400 mg</i>	1	QL (270 EA per 30 days)
<i>gabapentin oral capsule 200 mg</i>	5	QL (120 EA per 30 days)
<i>gabapentin oral capsule 300 mg</i>	1	QL (360 EA per 30 days)
<i>gabapentin oral solution 250 mg/5ml, 300 mg/6ml</i>	2	QL (2160 ML per 30 days)
<i>gabapentin oral tablet 600 mg</i>	1	QL (180 EA per 30 days)
<i>gabapentin oral tablet 800 mg</i>	1	QL (120 EA per 30 days)
<i>midazolam intramuscular solution auto-injector 10 mg/0.7ml</i>	2	QL (2.8 ML per 30 days)
NAYZILAM NASAL SOLUTION 5 MG/0.1ML	4	PA; QL (10 EA per 30 days)
<i>phenobarbital oral elixir 20 mg/5ml, 30 mg/7.5ml, 60 mg/15ml</i>	2	PA
<i>phenobarbital oral tablet 100 mg, 15 mg, 16.2 mg, 30 mg, 32.4 mg, 60 mg, 64.8 mg, 97.2 mg</i>	2	PA
<i>pregabalin oral capsule 100 mg, 150 mg, 200 mg, 25 mg, 50 mg, 75 mg</i>	1	QL (90 EA per 30 days)
<i>pregabalin oral capsule 225 mg, 300 mg</i>	1	QL (60 EA per 30 days)
<i>pregabalin oral solution 20 mg/ml</i>	1	QL (900 ML per 30 days)
<i>primidone oral tablet 250 mg, 50 mg</i>	2	
RELGAABI ORAL CAPSULE 200 MG	5	ST; QL (120 EA per 30 days)
RELGAABI ORAL CAPSULE 300 MG	5	ST; QL (360 EA per 30 days)
RELGAABI ORAL CAPSULE 400 MG	5	ST; QL (270 EA per 30 days)
SYMPAZAN ORAL FILM 10 MG, 20 MG	5	ST; QL (60 EA per 30 days)
SYMPAZAN ORAL FILM 5 MG	4	ST; QL (60 EA per 30 days)
<i>tiagabine hcl oral tablet 12 mg, 16 mg, 2 mg, 4 mg</i>	2	
VALTOCO 10 MG DOSE NASAL LIQUID 10 MG/0.1ML	4	PA; QL (10 EA per 30 days)
VALTOCO 15 MG DOSE NASAL LIQUID THERAPY PACK 2 X 7.5 MG/0.1ML	4	PA; QL (10 EA per 30 days)
VALTOCO 20 MG DOSE NASAL LIQUID THERAPY PACK 2 X 10 MG/0.1ML	4	PA; QL (10 EA per 30 days)
VALTOCO 5 MG DOSE NASAL LIQUID 5 MG/0.1ML	4	PA; QL (10 EA per 30 days)
<i>vigabatrin oral packet 500 mg</i>	5	PA; QL (180 EA per 30 days)
<i>vigabatrin oral tablet 500 mg</i>	5	PA; QL (180 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
VIGAFYDE ORAL SOLUTION 100 MG/ML	5	PA
ZTALMY ORAL SUSPENSION 50 MG/ML	5	PA
Sodium Channel Agents		
<i>carbamazepine er oral capsule extended release 12 hour 100 mg, 200 mg, 300 mg</i>	2	
<i>carbamazepine er oral tablet extended release 12 hour 100 mg, 200 mg, 400 mg</i>	2	
<i>carbamazepine oral suspension 100 mg/5ml</i>	2	
<i>carbamazepine oral tablet 200 mg</i>	1	
<i>carbamazepine oral tablet chewable 100 mg, 200 mg</i>	2	
DILANTIN ORAL CAPSULE 100 MG, 30 MG	4	
<i>eslicarbazepine acetate oral tablet 200 mg, 400 mg</i>	2	QL (30 EA per 30 days)
<i>eslicarbazepine acetate oral tablet 600 mg, 800 mg</i>	2	QL (60 EA per 30 days)
<i>lacosamide oral solution 10 mg/ml, 100 mg/10ml, 50 mg/5ml</i>	2	QL (1200 ML per 30 days)
<i>lacosamide oral tablet 100 mg, 150 mg, 200 mg, 50 mg</i>	2	QL (60 EA per 30 days)
<i>oxcarbazepine er oral tablet extended release 24 hour 150 mg, 300 mg</i>	2	
<i>oxcarbazepine er oral tablet extended release 24 hour 600 mg</i>	5	
<i>oxcarbazepine oral suspension 300 mg/5ml</i>	2	
<i>oxcarbazepine oral tablet 150 mg, 300 mg, 600 mg</i>	1	
PHENYTEK ORAL CAPSULE 200 MG, 300 MG	4	
PHENYTOIN INFATABS ORAL TABLET CHEWABLE 50 MG	2	
<i>phenytoin oral suspension 125 mg/5ml</i>	2	
<i>phenytoin oral tablet chewable 50 mg</i>	2	
<i>phenytoin sodium extended oral capsule 100 mg, 200 mg, 300 mg</i>	2	
<i>rufinamide oral suspension 40 mg/ml</i>	2	PA; QL (2400 ML per 30 days)
<i>rufinamide oral tablet 200 mg, 400 mg</i>	2	PA; QL (240 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
ZONISADE ORAL SUSPENSION 100 MG/5ML	4	ST
<i>zonisamide oral capsule 100 mg, 25 mg, 50 mg</i>	2	
Antidementia Agents - Management Of Dementia		
Antidementia Agents, Other		
<i>memantine hcl-donepezil hcl er oral capsule extended release 24 hour 14-10 mg, 21-10 mg, 28-10 mg</i>	2	
NAMZARIC ORAL CAPSULE EXTENDED RELEASE 24 HOUR 7-10 MG	4	
Cholinesterase Inhibitors		
<i>donepezil hcl oral tablet 10 mg, 5 mg</i>	1	
<i>donepezil hcl oral tablet 23 mg</i>	2	
<i>donepezil hcl oral tablet dispersible 10 mg, 5 mg</i>	1	
<i>galantamine hydrobromide er oral capsule extended release 24 hour 16 mg, 24 mg, 8 mg</i>	1	
<i>galantamine hydrobromide oral tablet 12 mg, 4 mg, 8 mg</i>	1	
<i>rivastigmine tartrate oral capsule 1.5 mg, 3 mg, 4.5 mg, 6 mg</i>	1	QL (60 EA per 30 days)
<i>rivastigmine transdermal patch 24 hour 13.3 mg/24hr, 4.6 mg/24hr, 9.5 mg/24hr</i>	2	QL (30 EA per 30 days)
N-Methyl-D-Aspartate (Nmda) Receptor Antagonist		
<i>memantine hcl er oral capsule extended release 24 hour 14 mg, 21 mg, 28 mg, 7 mg</i>	2	QL (30 EA per 30 days)
<i>memantine hcl oral tablet 10 mg, 28 x 5 mg & 21 x 10 mg, 5 mg</i>	1	
Antidepressants - Treatment Of Depression		
Antidepressants, Other		
AUVELITY ORAL TABLET EXTENDED RELEASE 45-105 MG	5	PA
AUVELITY TITRATION PACK ORAL TABLET EXTENDED RELEASE THERAPY PACK 30-105 MG & 45-105 MG	5	PA

Name of Drug	Drug Tier	Requirements/Limits
<i>bupropion hcl er (sr) oral tablet extended release 12 hour 100 mg, 150 mg, 200 mg</i>	1	
<i>bupropion hcl er (xl) oral tablet extended release 24 hour 150 mg, 300 mg, 450 mg</i>	1	
<i>bupropion hcl oral tablet 100 mg, 75 mg</i>	1	
<i>duloxetine hcl oral capsule delayed release particles 120 mg, 80 mg, 90 mg</i>	2	QL (30 EA per 30 days)
EXXUA ORAL TABLET EXTENDED RELEASE 24 HOUR 18.2 MG, 36.3 MG, 54.5 MG, 72.6 MG	5	ST
EXXUA TITRATION PACK ORAL TABLET EXTENDED RELEASE 24 HOUR 18.2 MG	5	ST
<i>mirtazapine oral tablet 15 mg, 30 mg, 45 mg, 7.5 mg</i>	1	
<i>mirtazapine oral tablet dispersible 15 mg, 30 mg, 45 mg</i>	2	
<i>perphenazine-amitriptyline oral tablet 2-10 mg, 2-25 mg, 4-10 mg, 4-25 mg, 4-50 mg</i>	2	PA
ZURZUVAE ORAL CAPSULE 20 MG, 25 MG, 30 MG	5	PA
Monoamine Oxidase Inhibitors		
EMSAM TRANSDERMAL PATCH 24 HOUR 12 MG/24HR, 6 MG/24HR, 9 MG/24HR	5	PA
MARPLAN ORAL TABLET 10 MG	4	
<i>phenelzine sulfate oral tablet 15 mg</i>	2	
<i>tranylcypromine sulfate oral tablet 10 mg</i>	2	
Ssri/Snri (Selective Serotonin Reuptake Inhibitor/Serotonin And Norepinephrine Reuptake Inhibitor)		
<i>citalopram hydrobromide oral solution 10 mg/5ml</i>	2	
<i>citalopram hydrobromide oral tablet 10 mg, 20 mg, 40 mg</i>	1	
<i>desvenlafaxine succinate er oral tablet extended release 24 hour 100 mg</i>	2	QL (60 EA per 30 days)
<i>desvenlafaxine succinate er oral tablet extended release 24 hour 25 mg, 50 mg</i>	2	QL (30 EA per 30 days)
<i>escitalopram oxalate oral solution 10 mg/10ml, 5 mg/5ml</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>escitalopram oxalate oral tablet 10 mg, 20 mg, 5 mg</i>	1	
FETZIMA ORAL CAPSULE EXTENDED RELEASE 24 HOUR 120 MG, 20 MG, 40 MG, 80 MG	4	ST; QL (30 EA per 30 days)
FETZIMA TITRATION ORAL CAPSULE ER 24 HOUR THERAPY PACK 20 & 40 MG	4	ST; QL (28 EA per 180 days)
<i>fluoxetine hcl oral capsule 10 mg, 20 mg, 40 mg</i>	1	
<i>fluoxetine hcl oral capsule delayed release 90 mg</i>	2	
<i>fluoxetine hcl oral solution 20 mg/5ml</i>	1	
<i>fluoxetine hcl oral tablet 10 mg, 20 mg, 60 mg</i>	2	
<i>fluvoxamine maleate oral tablet 100 mg, 25 mg, 50 mg</i>	2	
<i>nefazodone hcl oral tablet 100 mg, 150 mg, 200 mg, 250 mg, 50 mg</i>	2	
<i>paroxetine hcl er oral tablet extended release 24 hour 12.5 mg, 25 mg, 37.5 mg</i>	2	
<i>paroxetine hcl oral suspension 10 mg/5ml</i>	2	
<i>paroxetine hcl oral tablet 10 mg, 20 mg, 30 mg, 40 mg</i>	1	
RALDESY ORAL SOLUTION 10 MG/ML	4	
<i>sertraline hcl oral concentrate 20 mg/ml</i>	2	
<i>sertraline hcl oral tablet 100 mg, 25 mg, 50 mg</i>	1	
<i>trazodone hcl oral tablet 100 mg, 150 mg, 50 mg</i>	1	
<i>trazodone hcl oral tablet 300 mg</i>	2	
TRINTELLIX ORAL TABLET 10 MG, 20 MG, 5 MG	3	QL (30 EA per 30 days)
<i>venlafaxine hcl er oral capsule extended release 24 hour 150 mg, 37.5 mg, 75 mg</i>	2	
<i>venlafaxine hcl er oral tablet extended release 24 hour 225 mg</i>	2	
<i>venlafaxine hcl oral tablet 100 mg, 25 mg, 37.5 mg, 50 mg, 75 mg</i>	2	
<i>vilazodone hcl oral tablet 10 mg, 20 mg, 40 mg</i>	2	
Tricyclics		
<i>amitriptyline hcl oral tablet 10 mg, 100 mg, 150 mg, 25 mg, 50 mg, 75 mg</i>	2	PA

Name of Drug	Drug Tier	Requirements/Limits
<i>amoxapine oral tablet 100 mg, 150 mg, 25 mg, 50 mg</i>	2	PA
<i>clomipramine hcl oral capsule 25 mg, 50 mg, 75 mg</i>	2	PA
<i>desipramine hcl oral tablet 10 mg, 100 mg, 150 mg, 25 mg, 50 mg, 75 mg</i>	2	
<i>doxepin hcl oral capsule 10 mg, 100 mg, 150 mg, 25 mg, 50 mg, 75 mg</i>	2	
<i>doxepin hcl oral concentrate 10 mg/ml</i>	2	
<i>imipramine hcl oral tablet 10 mg, 25 mg, 50 mg</i>	2	
<i>imipramine pamoate oral capsule 100 mg, 125 mg, 150 mg, 75 mg</i>	2	
<i>nortriptyline hcl oral capsule 10 mg, 25 mg, 50 mg, 75 mg</i>	1	PA
<i>nortriptyline hcl oral solution 10 mg/5ml</i>	2	PA
<i>protriptyline hcl oral tablet 10 mg, 5 mg</i>	2	
<i>trimipramine maleate oral capsule 100 mg, 25 mg, 50 mg</i>	2	
Antiemetics - Treatment Of Vomiting Or Nausea		
Antiemetics, Other		
<i>chlorpromazine hcl oral concentrate 100 mg/ml, 30 mg/ml</i>	2	
<i>chlorpromazine hcl oral tablet 10 mg, 100 mg, 200 mg, 25 mg, 50 mg</i>	2	
<i>meclizine hcl oral tablet 12.5 mg, 25 mg</i>	2	
<i>metoclopramide hcl oral solution 5 mg/5ml</i>	2	
<i>metoclopramide hcl oral tablet 10 mg, 5 mg</i>	1	
<i>perphenazine oral tablet 16 mg, 2 mg, 4 mg, 8 mg</i>	2	
<i>prochlorperazine maleate oral tablet 10 mg, 5 mg</i>	1	
<i>prochlorperazine rectal suppository 25 mg</i>	2	
<i>promethazine hcl oral tablet 12.5 mg, 25 mg, 50 mg</i>	2	PA
<i>promethazine hcl rectal suppository 12.5 mg, 25 mg</i>	2	PA
PROMETHEGAN RECTAL SUPPOSITORY 50 MG	4	PA

Name of Drug	Drug Tier	Requirements/Limits
<i>scopolamine transdermal patch 72 hour 1 mg/3days</i>	2	
<i>trimethobenzamide hcl oral capsule 300 mg</i>	2	
Emetogenic Therapy Adjuncts		
<i>aprepitant oral capsule 125 mg, 40 mg, 80 mg</i>	2	B/D
<i>aprepitant oral capsule therapy pack 80 & 125 mg</i>	2	B/D
<i>dronabinol oral capsule 10 mg, 2.5 mg, 5 mg</i>	2	B/D
EMEND ORAL SUSPENSION RECONSTITUTED 125 MG/5ML	4	B/D
<i>granisetron hcl oral tablet 1 mg</i>	2	B/D
<i>ondansetron hcl oral solution 4 mg/5ml</i>	2	B/D
<i>ondansetron hcl oral tablet 24 mg</i>	2	B/D
<i>ondansetron hcl oral tablet 4 mg, 8 mg</i>	1	B/D
<i>ondansetron oral tablet dispersible 4 mg, 8 mg</i>	2	B/D
Antifungals - Treatment Of Fungal Or Yeast Infections		
Antifungals		
<i>amphotericin b intravenous solution reconstituted 50 mg</i>	2	B/D
<i>amphotericin b liposome intravenous suspension reconstituted 50 mg</i>	5	B/D
<i>caspofungin acetate intravenous solution reconstituted 50 mg, 70 mg</i>	4	PA
<i>clotrimazole external cream 1 %</i>	2	QL (45 GM per 28 days)
<i>clotrimazole external solution 1 %</i>	2	QL (30 ML per 28 days)
<i>clotrimazole mouth/throat troche 10 mg</i>	2	
CRESEMBA ORAL CAPSULE 186 MG, 74.5 MG	5	PA
<i>econazole nitrate external cream 1 %</i>	2	
<i>fluconazole in sodium chloride intravenous solution 200-0.9 mg/100ml-%, 400-0.9 mg/200ml-%</i>	2	
<i>fluconazole oral suspension reconstituted 10 mg/ml, 40 mg/ml</i>	2	
<i>fluconazole oral tablet 100 mg, 150 mg, 200 mg, 50 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>flucytosine oral capsule 250 mg, 500 mg</i>	5	PA
<i>griseofulvin microsize oral suspension 125 mg/5ml</i>	2	
<i>itraconazole oral capsule 100 mg</i>	2	
<i>itraconazole oral solution 10 mg/ml</i>	2	
<i>ketoconazole external cream 2 %</i>	2	
<i>ketoconazole external shampoo 2 %</i>	1	
<i>ketoconazole oral tablet 200 mg</i>	2	
KLAYESTA EXTERNAL POWDER 100000 UNIT/GM	1	
<i>micafungin sodium intravenous solution reconstituted 100 mg</i>	4	
<i>micafungin sodium intravenous solution reconstituted 50 mg</i>	2	
NYAMYC EXTERNAL POWDER 100000 UNIT/GM	1	
<i>nystatin external cream 100000 unit/gm</i>	1	
<i>nystatin external ointment 100000 unit/gm</i>	1	
<i>nystatin external powder 100000 unit/gm</i>	1	
<i>nystatin mouth/throat suspension 100000 unit/ml</i>	2	
<i>nystatin oral tablet 500000 unit</i>	2	
NYSTOP EXTERNAL POWDER 100000 UNIT/GM	1	
<i>posaconazole intravenous solution 300 mg/16.7ml</i>	2	
<i>posaconazole oral suspension 40 mg/ml</i>	5	PA
<i>posaconazole oral tablet delayed release 100 mg</i>	2	PA
<i>terbinafine hcl oral tablet 250 mg</i>	1	
<i>terconazole vaginal cream 0.4 %, 0.8 %</i>	2	
<i>terconazole vaginal suppository 80 mg</i>	2	
<i>voriconazole intravenous solution 200 mg/20ml</i>	2	PA
<i>voriconazole intravenous solution reconstituted 200 mg</i>	4	PA
<i>voriconazole oral suspension reconstituted 40 mg/ml</i>	5	
<i>voriconazole oral tablet 200 mg, 50 mg</i>	2	

Antigout Agents - Treatment Or Prevention Of Gouty Arthritis

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Formulary ID 26314, Version 14
Page 30

Last Updated 7/2026

Please refer to our Evidence of Coverage for more information about this coverage. You can find information on what the symbols and abbreviations on this table mean by going to Pages 7-8.

Name of Drug	Drug Tier	Requirements/Limits
Antigout Agents		
<i>allopurinol oral tablet 100 mg, 300 mg</i>	1	
<i>colchicine oral capsule 0.6 mg</i>	2	
<i>colchicine oral tablet 0.6 mg</i>	2	
<i>colchicine-probenecid oral tablet 0.5-500 mg</i>	2	
<i>febuxostat oral tablet 40 mg, 80 mg</i>	2	ST
<i>probenecid oral tablet 500 mg</i>	2	
Antimigraine Agents - Treatment Of Migraine Headaches		
Antimigraine Agents		
NURTEC ORAL TABLET DISPERSIBLE 75 MG	3	PA; QL (18 EA per 30 days)
UBRELVY ORAL TABLET 100 MG, 50 MG	3	PA; QL (16 EA per 30 days)
ZAVZPRET NASAL SOLUTION 10 MG/ACT	5	PA; QL (8 EA per 30 days)
Ergot Alkaloids		
<i>dihydroergotamine mesylate nasal solution 4 mg/ml</i>	5	PA; QL (8 ML per 30 days)
<i>ergotamine-caffeine oral tablet 1-100 mg</i>	2	PA
Prophylactic		
AIMOVIG SUBCUTANEOUS SOLUTION AUTO-INJECTOR 140 MG/ML, 70 MG/ML	3	PA; QL (1 ML per 30 days)
EMGALITY (300 MG DOSE) SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 100 MG/ML	3	PA; QL (3 ML per 30 days)
EMGALITY SUBCUTANEOUS SOLUTION AUTO-INJECTOR 120 MG/ML	3	PA; QL (2 ML per 30 days)
EMGALITY SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 120 MG/ML	3	PA; QL (2 ML per 30 days)
QULIPTA ORAL TABLET 10 MG, 30 MG, 60 MG	3	PA; QL (30 EA per 30 days)
Serotonin (5-Ht) Receptor Agonist		
<i>naratriptan hcl oral tablet 1 mg, 2.5 mg</i>	2	QL (9 EA per 28 days)
<i>rizatriptan benzoate oral tablet 10 mg, 5 mg</i>	2	QL (12 EA per 30 days)
<i>rizatriptan benzoate oral tablet dispersible 10 mg, 5 mg</i>	2	QL (12 EA per 30 days)
<i>sumatriptan nasal solution 20 mg/act, 5 mg/act</i>	2	QL (12 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>sumatriptan succinate oral tablet 100 mg, 25 mg, 50 mg</i>	2	QL (9 EA per 30 days)
<i>sumatriptan succinate subcutaneous solution 6 mg/0.5ml</i>	2	QL (4 ML per 30 days)
<i>sumatriptan succinate subcutaneous solution auto-injector 6 mg/0.5ml</i>	2	QL (4 ML per 30 days)
<i>zolmitriptan oral tablet 2.5 mg, 5 mg</i>	2	QL (9 EA per 28 days)
<i>zolmitriptan oral tablet dispersible 2.5 mg, 5 mg</i>	2	QL (9 EA per 28 days)
Antimyasthenic Agents - Treatment Of Myasthenia		
Parasympathomimetics		
<i>pyridostigmine bromide er oral tablet extended release 180 mg</i>	2	
<i>pyridostigmine bromide er oral tablet extended release 24 hour 105 mg</i>	2	
<i>pyridostigmine bromide oral tablet 60 mg</i>	1	
Antimycobacterials - Treatment For Infections By Tuberculosis-Type Organisms		
Antimycobacterials, Other		
<i>dapsone oral tablet 100 mg, 25 mg</i>	2	
<i>rifabutin oral capsule 150 mg</i>	4	
Antituberculars		
<i>ethambutol hcl oral tablet 100 mg, 400 mg</i>	2	
<i>isoniazid oral tablet 100 mg, 300 mg</i>	1	
<i>pretomanid oral tablet 200 mg</i>	4	PA
PRIFTIN ORAL TABLET 150 MG	4	
<i>pyrazinamide oral tablet 500 mg</i>	2	
<i>rifampin intravenous solution reconstituted 600 mg</i>	4	
<i>rifampin oral capsule 150 mg, 300 mg</i>	2	
SIRTURO ORAL TABLET 100 MG, 20 MG	5	PA
Antineoplastics - Treatment Of Cancer		
Alkylating Agents		
<i>cyclophosphamide oral capsule 25 mg, 50 mg</i>	2	B/D
<i>cyclophosphamide oral tablet 25 mg, 50 mg</i>	2	B/D

Name of Drug	Drug Tier	Requirements/Limits
LEUKERAN ORAL TABLET 2 MG	5	PA
<i>lomustine oral capsule 10 mg</i>	4	
<i>lomustine oral capsule 100 mg, 40 mg</i>	5	
MATULANE ORAL CAPSULE 50 MG	5	
VALCHLOR EXTERNAL GEL 0.016 %	5	PA
Antiandrogens		
<i>abiraterone acetate oral tablet 250 mg</i>	2	PA
<i>abiraterone acetate oral tablet 500 mg</i>	5	PA
ABIRTEGA ORAL TABLET 250 MG	4	PA; QL (120 EA per 30 days)
<i>bicalutamide oral tablet 50 mg</i>	1	
ERLEADA ORAL TABLET 240 MG, 60 MG	5	PA
EULEXIN ORAL CAPSULE 125 MG	5	PA
<i>nilutamide oral tablet 150 mg</i>	5	PA
NUBEQA ORAL TABLET 300 MG	5	PA
XTANDI ORAL CAPSULE 40 MG	5	PA
XTANDI ORAL TABLET 40 MG, 80 MG	5	PA
YONSA ORAL TABLET 125 MG	5	PA
Antiangiogenic Agents		
<i>lenalidomide oral capsule 10 mg, 15 mg, 2.5 mg, 20 mg, 25 mg, 5 mg</i>	5	PA
<i>pomalidomide oral capsule 1 mg, 2 mg, 3 mg, 4 mg</i>	5	PA
REVLIMID ORAL CAPSULE 10 MG, 15 MG, 2.5 MG, 20 MG, 25 MG, 5 MG	5	PA
THALOMID ORAL CAPSULE 100 MG, 50 MG	5	PA
Antiestrogens/Modifiers		
SOLTAMOX ORAL SOLUTION 10 MG/5ML	5	PA
<i>tamoxifen citrate oral tablet 10 mg, 20 mg</i>	1	
<i>toremifene citrate oral tablet 60 mg</i>	5	PA
Antimetabolites		
DROXIA ORAL CAPSULE 200 MG, 300 MG, 400 MG	4	
<i>hydroxyurea oral capsule 500 mg</i>	2	
INQOVI ORAL TABLET 35-100 MG	5	PA
<i>mercaptopurine oral suspension 2000 mg/100ml</i>	5	PA

Name of Drug	Drug Tier	Requirements/Limits
<i>mercaptopurine oral tablet 50 mg</i>	2	
ONUREG ORAL TABLET 200 MG, 300 MG	5	PA
SIKLOS ORAL TABLET 100 MG, 1000 MG	4	
TABLOID ORAL TABLET 40 MG	4	PA
XROMI ORAL SOLUTION 100 MG/ML	4	
Antineoplastics, Other		
AKEEGA ORAL TABLET 100-500 MG, 50-500 MG	5	PA
AVMAPKI FAKZYNJA CO-PACK ORAL THERAPY PACK 0.8 & 200 MG	5	PA; QL (66 EA per 28 days)
BESREMI SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 500 MCG/ML	5	PA
DANZITEN ORAL TABLET 71 MG, 95 MG	5	PA
GOMEKLI ORAL CAPSULE 1 MG, 2 MG	5	PA
GOMEKLI ORAL TABLET SOLUBLE 1 MG	5	PA
IDHIFA ORAL TABLET 100 MG, 50 MG	5	PA
INLURIYO ORAL TABLET 200 MG	5	PA
IWILFIN ORAL TABLET 192 MG	5	PA
JYLAMVO ORAL SOLUTION 2 MG/ML	4	PA
KOMZIFTI ORAL CAPSULE 200 MG	5	PA
KRAZATI ORAL TABLET 200 MG	5	PA
LAZCLUZE ORAL TABLET 240 MG, 80 MG	5	PA
LIFYORLI (125 MG DOSE) ORAL CAPSULE THERAPY PACK 1 X 25 MG & 1 X 100 MG	5	PA
LIFYORLI (150 MG DOSE) ORAL CAPSULE THERAPY PACK 2 X 25 MG & 1 X 100 MG	5	PA
LONSURF ORAL TABLET 15-6.14 MG, 20-8.19 MG	5	PA
LUMAKRAS ORAL TABLET 120 MG, 240 MG, 320 MG	5	PA
LYSODREN ORAL TABLET 500 MG	5	
MODEYSO ORAL CAPSULE 125 MG	5	PA; QL (20 EA per 28 days)
NINLARO ORAL CAPSULE 2.3 MG, 3 MG, 4 MG	5	PA
OJJAARA ORAL TABLET 100 MG, 150 MG, 200 MG	5	PA
ORSERDU ORAL TABLET 345 MG, 86 MG	5	PA

Name of Drug	Drug Tier	Requirements/Limits
REVUFORJ ORAL TABLET 110 MG, 160 MG, 25 MG	5	PA
REZLIDHIA ORAL CAPSULE 150 MG	5	PA
ROMVIMZA ORAL CAPSULE 14 MG, 20 MG, 30 MG	5	PA; QL (8 EA per 28 days)
RYLAZE INTRAMUSCULAR SOLUTION 10 MG/0.5ML	5	PA
TIBSOVO ORAL TABLET 250 MG	5	PA
VORANIGO ORAL TABLET 10 MG, 40 MG	5	PA
WELIREG ORAL TABLET 40 MG	5	PA
XATMEP ORAL SOLUTION 2.5 MG/ML	4	PA
XPOVIO (100 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 50 MG	5	PA
XPOVIO (40 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 10 MG, 40 MG	5	PA
XPOVIO (40 MG TWICE WEEKLY) ORAL TABLET THERAPY PACK 40 MG	5	PA
XPOVIO (60 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 60 MG	5	PA
XPOVIO (60 MG TWICE WEEKLY) ORAL TABLET THERAPY PACK 20 MG	5	PA
XPOVIO (80 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 40 MG, 80 MG	5	PA
XPOVIO (80 MG TWICE WEEKLY) ORAL TABLET THERAPY PACK 20 MG	5	PA
ZOLINZA ORAL CAPSULE 100 MG	5	PA
Aromatase Inhibitors, 3Rd Generation		
<i>anastrozole oral tablet 1 mg</i>	1	
<i>exemestane oral tablet 25 mg</i>	2	
<i>letrozole oral tablet 2.5 mg</i>	1	
Molecular Target Inhibitors		
ALECENSA ORAL CAPSULE 150 MG	5	PA
ALUNBRIG ORAL TABLET 180 MG, 30 MG, 90 MG	5	PA
ALUNBRIG ORAL TABLET THERAPY PACK 90 & 180 MG	5	PA
AUGTYRO ORAL CAPSULE 160 MG, 40 MG	5	PA

Name of Drug	Drug Tier	Requirements/Limits
AYVAKIT ORAL TABLET 100 MG, 200 MG, 25 MG, 300 MG, 50 MG	5	PA
BALVERSA ORAL TABLET 3 MG, 4 MG, 5 MG	5	PA
BOSULIF ORAL CAPSULE 100 MG, 50 MG	5	PA
BOSULIF ORAL TABLET 100 MG, 400 MG, 500 MG	5	PA
<i>bosutinib oral tablet 100 mg, 400 mg, 500 mg</i>	5	PA
BRAFTOVI ORAL CAPSULE 75 MG	5	PA
BRUKINSA ORAL CAPSULE 80 MG	5	PA
BRUKINSA ORAL TABLET 160 MG	5	PA
CABOMETYX ORAL TABLET 20 MG, 40 MG, 60 MG	5	PA
CALQUENCE ORAL TABLET 100 MG	5	PA
CAPRELSA ORAL TABLET 100 MG, 300 MG	5	PA
COMETRIQ (100 MG DAILY DOSE) ORAL KIT 80 & 20 MG	5	PA
COMETRIQ (140 MG DAILY DOSE) ORAL KIT 3 X 20 MG & 80 MG	5	PA
COMETRIQ (60 MG DAILY DOSE) ORAL KIT 20 MG	5	PA
COPIKTRA ORAL CAPSULE 15 MG, 25 MG	5	PA
COTELLIC ORAL TABLET 20 MG	5	PA
<i>dasatinib oral tablet 100 mg, 140 mg, 20 mg, 50 mg, 70 mg, 80 mg</i>	5	PA
DAURISMO ORAL TABLET 100 MG, 25 MG	5	PA
ENSACOVE ORAL CAPSULE 100 MG, 25 MG	5	PA; QL (30 EA per 30 days)
ERIVEDGE ORAL CAPSULE 150 MG	5	PA
<i>erlotinib hcl oral tablet 100 mg, 150 mg, 25 mg</i>	5	PA
<i>everolimus oral tablet 10 mg, 2.5 mg, 5 mg, 7.5 mg</i>	5	PA
<i>everolimus oral tablet soluble 2 mg, 3 mg, 5 mg</i>	5	PA
FOTIVDA ORAL CAPSULE 0.89 MG, 1.34 MG	5	PA
FRUZAQLA ORAL CAPSULE 1 MG, 5 MG	5	PA
GAVRETO ORAL CAPSULE 100 MG	5	PA
<i>gefitinib oral tablet 250 mg</i>	5	PA

Name of Drug	Drug Tier	Requirements/Limits
GILOTRIF ORAL TABLET 20 MG, 30 MG, 40 MG	5	PA
HERNEXEOS ORAL TABLET 60 MG	5	PA; QL (90 EA per 30 days)
HYRNUO ORAL TABLET 10 MG	5	PA; QL (120 EA per 30 days)
IBRANCE ORAL CAPSULE 100 MG, 125 MG, 75 MG	5	PA
IBRANCE ORAL TABLET 100 MG, 125 MG, 75 MG	5	PA
IBTROZI ORAL CAPSULE 200 MG	5	PA
ICLUSIG ORAL TABLET 10 MG, 15 MG, 30 MG, 45 MG	5	PA
<i>imatinib mesylate oral tablet 100 mg, 400 mg</i>	2	PA
IMBRUVICA ORAL CAPSULE 140 MG, 70 MG	5	PA
IMBRUVICA ORAL SUSPENSION 70 MG/ML	5	PA
IMBRUVICA ORAL TABLET 140 MG, 280 MG, 420 MG	5	PA
<i>imkeldi oral solution 80 mg/ml</i>	5	PA
INLYTA ORAL TABLET 1 MG, 5 MG	5	PA
INREBIC ORAL CAPSULE 100 MG	5	PA
ITOVEBI ORAL TABLET 3 MG, 9 MG	5	PA
JAKAFI ORAL TABLET 10 MG, 15 MG, 20 MG, 25 MG, 5 MG	5	PA
JAKAFI XR ORAL TABLET EXTENDED RELEASE 24 HOUR 11 MG, 22 MG, 33 MG, 44 MG, 55 MG	5	PA
JAYPIRCA ORAL TABLET 100 MG, 50 MG	5	PA
KISQALI (200 MG DOSE) ORAL TABLET THERAPY PACK 200 MG	5	PA
KISQALI (400 MG DOSE) ORAL TABLET THERAPY PACK 200 MG	5	PA
KISQALI (600 MG DOSE) ORAL TABLET THERAPY PACK 200 MG	5	PA
KOSELUGO ORAL CAPSULE 10 MG, 25 MG	5	PA
KOSELUGO ORAL CAPSULE SPRINKLE 5 MG, 7.5 MG	5	PA
<i>lapatinib ditosylate oral tablet 250 mg</i>	5	PA

Name of Drug	Drug Tier	Requirements/Limits
LENVIMA (10 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 10 MG	5	PA
LENVIMA (12 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 3 X 4 MG	5	PA
LENVIMA (14 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 10 & 4 MG	5	PA
LENVIMA (18 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 10 MG & 2 X 4 MG	5	PA
LENVIMA (20 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 2 X 10 MG	5	PA
LENVIMA (24 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 2 X 10 MG & 4 MG	5	PA
LENVIMA (4 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 4 MG	5	PA
LENVIMA (8 MG DAILY DOSE) ORAL CAPSULE THERAPY PACK 2 X 4 MG	5	PA
LORBRENA ORAL TABLET 100 MG, 25 MG	5	PA
LYNPARZA ORAL TABLET 100 MG, 150 MG	5	PA
LYTGOBI (12 MG DAILY DOSE) ORAL TABLET THERAPY PACK 4 MG	5	PA
LYTGOBI (16 MG DAILY DOSE) ORAL TABLET THERAPY PACK 4 MG	5	PA
LYTGOBI (20 MG DAILY DOSE) ORAL TABLET THERAPY PACK 4 MG	5	PA
MEKINIST ORAL SOLUTION RECONSTITUTED 0.05 MG/ML	5	PA
MEKINIST ORAL TABLET 0.5 MG, 2 MG	5	PA
MEKTOVI ORAL TABLET 15 MG	5	PA
NERLYNX ORAL TABLET 40 MG	5	PA
<i>nilotinib d-tartrate oral capsule 150 mg, 200 mg, 50 mg</i>	5	PA
<i>nilotinib hcl oral capsule 150 mg, 200 mg, 50 mg</i>	5	PA
ODOMZO ORAL CAPSULE 200 MG	5	PA
OGSIVEO ORAL TABLET 100 MG, 150 MG	5	PA
OJEMDA ORAL SUSPENSION RECONSTITUTED 25 MG/ML	5	PA

Name of Drug	Drug Tier	Requirements/Limits
OJEMDA ORAL TABLET 100 MG, 100 MG (16 PACK), 100 MG (24 PACK)	5	PA
<i>pazopanib hcl oral tablet 200 mg, 400 mg</i>	5	PA
PEMAZYRE ORAL TABLET 13.5 MG, 4.5 MG, 9 MG	5	PA
PIQRAY (200 MG DAILY DOSE) ORAL TABLET THERAPY PACK 200 MG	5	PA
PIQRAY (250 MG DAILY DOSE) ORAL TABLET THERAPY PACK 200 & 50 MG	5	PA
PIQRAY (300 MG DAILY DOSE) ORAL TABLET THERAPY PACK 2 X 150 MG	5	PA
QINLOCK ORAL TABLET 50 MG	5	PA
RETEVMO ORAL TABLET 120 MG, 160 MG, 40 MG, 80 MG	5	PA
ROZLYTREK ORAL CAPSULE 100 MG, 200 MG	5	PA
ROZLYTREK ORAL PACKET 50 MG	5	PA
RUBRACA ORAL TABLET 200 MG, 250 MG, 300 MG	5	PA
RYDAPT ORAL CAPSULE 25 MG	5	PA
SCSEMBLIX ORAL TABLET 100 MG, 20 MG, 40 MG	5	PA
<i>sorafenib tosylate oral tablet 200 mg</i>	5	PA
STIVARGA ORAL TABLET 40 MG	5	PA
<i>sunitinib malate oral capsule 12.5 mg, 25 mg, 37.5 mg, 50 mg</i>	5	PA
TABRECTA ORAL TABLET 150 MG, 200 MG	5	PA
TAFINLAR ORAL CAPSULE 50 MG, 75 MG	5	PA
TAFINLAR ORAL TABLET SOLUBLE 10 MG	5	PA
TAGRISSO ORAL TABLET 40 MG, 80 MG	5	PA
TALZENNA ORAL CAPSULE 0.1 MG, 0.25 MG, 0.35 MG, 0.5 MG, 0.75 MG, 1 MG	5	PA
TAZVERIK ORAL TABLET 200 MG	5	PA
TEPMETKO ORAL TABLET 225 MG	5	PA
TRUQAP ORAL TABLET 200 MG	5	PA
TRUQAP ORAL TABLET THERAPY PACK 160 MG, 200 MG	5	PA
TUKYSA ORAL TABLET 150 MG, 50 MG	5	PA

Name of Drug	Drug Tier	Requirements/Limits
TURALIO ORAL CAPSULE 125 MG	5	PA
VANFLYTA ORAL TABLET 17.7 MG, 26.5 MG	5	PA
VENCLEXTA ORAL TABLET 10 MG	4	PA
VENCLEXTA ORAL TABLET 100 MG, 50 MG	5	PA
VENCLEXTA STARTING PACK ORAL TABLET THERAPY PACK 10 & 50 & 100 MG	5	PA
VERZENIO ORAL TABLET 100 MG, 150 MG, 200 MG, 50 MG	5	PA
VIJOICE ORAL PACKET 50 MG	5	PA
VIJOICE ORAL TABLET THERAPY PACK 125 MG, 200 & 50 MG, 50 MG	5	PA
VITRAKVI ORAL CAPSULE 100 MG, 25 MG	5	PA
VITRAKVI ORAL SOLUTION 20 MG/ML	5	PA
VIZIMPRO ORAL TABLET 15 MG, 30 MG, 45 MG	5	PA
VONJO ORAL CAPSULE 100 MG	5	PA
XALKORI ORAL CAPSULE 200 MG, 250 MG	5	PA
XALKORI ORAL CAPSULE SPRINKLE 150 MG, 20 MG, 50 MG	5	PA
XOSPATA ORAL TABLET 40 MG	5	PA
ZEJULA ORAL TABLET 100 MG, 200 MG, 300 MG	5	PA
ZELBORAF ORAL TABLET 240 MG	5	PA
ZYDELIG ORAL TABLET 100 MG, 150 MG	5	PA
ZYKADIA ORAL TABLET 150 MG	5	PA
Retinoids		
<i>bexarotene external gel 1 %</i>	5	PA
<i>bexarotene oral capsule 75 mg</i>	5	PA
PANRETIN EXTERNAL GEL 0.1 %	5	PA
<i>tretinoin oral capsule 10 mg</i>	5	PA
Treatment Adjuncts		
LEDERLE LEUCOVORIN ORAL TABLET 5 MG	2	
<i>leucovorin calcium oral tablet 10 mg, 15 mg, 25 mg, 5 mg</i>	2	
<i>mesna oral tablet 400 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
Antiparasitics - Treatment Of Infections From Parasites		
Anthelmintics		
<i>albendazole oral tablet 200 mg</i>	2	
<i>ivermectin oral tablet 3 mg</i>	1	
<i>praziquantel oral tablet 600 mg</i>	2	
Antiprotozoals		
<i>atovaquone oral suspension 750 mg/5ml</i>	2	
<i>atovaquone-proguanil hcl oral tablet 250-100 mg, 62.5-25 mg</i>	2	
<i>chloroquine phosphate oral tablet 250 mg, 500 mg</i>	2	
COARTEM ORAL TABLET 20-120 MG	4	
<i>hydroxychloroquine sulfate oral tablet 100 mg, 200 mg, 300 mg, 400 mg</i>	1	
IMPAVIDO ORAL CAPSULE 50 MG	5	PA; QL (84 EA per 28 days)
<i>mefloquine hcl oral tablet 250 mg</i>	2	
<i>nitazoxanide oral tablet 500 mg</i>	4	
<i>pentamidine isethionate inhalation solution reconstituted 300 mg</i>	2	B/D
<i>pentamidine isethionate injection solution reconstituted 300 mg</i>	4	
<i>primaquine phosphate oral tablet 26.3 (15 base) mg</i>	2	
<i>pyrimethamine oral tablet 25 mg</i>	5	QL (90 EA per 30 days)
<i>quinine sulfate oral capsule 324 mg</i>	2	
Antiparkinson Agents - Treatment Of Parkinson's Disease		
Anticholinergics		
<i>benztropine mesylate oral tablet 0.5 mg, 1 mg, 2 mg</i>	1	PA
<i>trihexyphenidyl hcl oral solution 0.4 mg/ml</i>	2	
<i>trihexyphenidyl hcl oral tablet 2 mg, 5 mg</i>	1	
Antiparkinson Agents, Other		
<i>amantadine hcl oral capsule 100 mg</i>	1	
<i>amantadine hcl oral solution 50 mg/5ml</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>amantadine hcl oral tablet 100 mg</i>	1	
<i>carbidopa-levodopa-entacapone oral tablet 12.5-50-200 mg, 18.75-75-200 mg, 25-100-200 mg, 31.25-125-200 mg, 37.5-150-200 mg, 50-200-200 mg</i>	2	
<i>entacapone oral tablet 200 mg</i>	2	
GOCOVRI ORAL CAPSULE EXTENDED RELEASE 24 HOUR 137 MG, 68.5 MG	5	PA
ONGENTYS ORAL CAPSULE 25 MG, 50 MG	4	ST
Dopamine Agonists		
<i>apomorphine hcl subcutaneous solution cartridge 30 mg/3ml</i>	5	PA
<i>bromocriptine mesylate oral capsule 5 mg</i>	2	
<i>bromocriptine mesylate oral tablet 2.5 mg</i>	2	
NEUPRO TRANSDERMAL PATCH 24 HOUR 1 MG/24HR, 2 MG/24HR, 3 MG/24HR, 4 MG/24HR, 6 MG/24HR, 8 MG/24HR	4	
<i>pramipexole dihydrochloride er oral tablet extended release 24 hour 0.375 mg, 0.75 mg, 1.5 mg, 2.25 mg, 3 mg, 3.75 mg, 4.5 mg</i>	2	
<i>pramipexole dihydrochloride oral tablet 0.125 mg, 0.25 mg, 0.5 mg, 0.75 mg, 1 mg, 1.5 mg</i>	1	
<i>ropinirole hcl er oral tablet extended release 24 hour 12 mg, 2 mg, 4 mg, 6 mg, 8 mg</i>	2	
<i>ropinirole hcl oral tablet 0.25 mg, 0.5 mg, 1 mg, 2 mg, 3 mg, 4 mg, 5 mg</i>	1	
Dopamine Precursors And/Or L-Amino Acid Decarboxylase Inhibitors		
<i>carbidopa oral tablet 25 mg</i>	2	
<i>carbidopa-levodopa er oral tablet extended release 25-100 mg, 50-200 mg</i>	1	
<i>carbidopa-levodopa oral tablet 10-100 mg, 25-100 mg, 25-250 mg</i>	1	
<i>carbidopa-levodopa oral tablet dispersible 10-100 mg, 25-100 mg, 25-250 mg</i>	1	
Monoamine Oxidase B (Mao-B) Inhibitors		
<i>rasagiline mesylate oral tablet 0.5 mg, 1 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>selegiline hcl oral capsule 5 mg</i>	2	
<i>selegiline hcl oral tablet 5 mg</i>	2	
Antipsychotics - Treatment Of Behavioral And Emotional Disorders		
1St Generation/Typical		
<i>fluphenazine decanoate injection solution 25 mg/ml</i>	2	
<i>fluphenazine hcl injection solution 2.5 mg/ml</i>	4	
<i>fluphenazine hcl oral concentrate 5 mg/ml</i>	2	
<i>fluphenazine hcl oral elixir 2.5 mg/5ml</i>	2	
<i>fluphenazine hcl oral tablet 1 mg, 10 mg, 2.5 mg, 5 mg</i>	2	
<i>haloperidol decanoate intramuscular solution 100 mg/ml, 100 mg/ml 1 ml, 50 mg/ml, 50 mg/ml(1ml)</i>	2	
<i>haloperidol lactate injection solution 5 mg/ml</i>	2	
<i>haloperidol lactate oral concentrate 2 mg/ml</i>	1	
<i>haloperidol oral tablet 0.5 mg, 1 mg, 10 mg, 2 mg, 20 mg, 5 mg</i>	1	
<i>loxapine succinate oral capsule 10 mg, 25 mg, 5 mg, 50 mg</i>	2	
<i>molindone hcl oral tablet 10 mg, 25 mg, 5 mg</i>	4	
<i>pimozide oral tablet 1 mg, 2 mg</i>	2	
<i>thioridazine hcl oral tablet 10 mg, 100 mg, 25 mg, 50 mg</i>	2	
<i>thiothixene oral capsule 1 mg, 10 mg, 2 mg, 5 mg</i>	2	
<i>trifluoperazine hcl oral tablet 1 mg, 10 mg, 2 mg, 5 mg</i>	2	
2Nd Generation/Atypical		
ABILIFY ASIMTUFII INTRAMUSCULAR PREFILLED SYRINGE 720 MG/2.4ML	3	QL (2.4 ML per 56 days)
ABILIFY ASIMTUFII INTRAMUSCULAR PREFILLED SYRINGE 960 MG/3.2ML	3	QL (3.2 ML per 56 days)
ABILIFY MAINTENA INTRAMUSCULAR PREFILLED SYRINGE 300 MG, 400 MG	3	QL (1 EA per 28 days)
ABILIFY MAINTENA INTRAMUSCULAR SUSPENSION RECONSTITUTED ER 300 MG, 400 MG	3	QL (1 EA per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>aripiprazole oral solution 1 mg/ml</i>	2	QL (900 ML per 30 days)
<i>aripiprazole oral tablet 10 mg, 15 mg, 2 mg, 20 mg, 30 mg, 5 mg</i>	1	QL (30 EA per 30 days)
<i>aripiprazole oral tablet dispersible 10 mg, 15 mg</i>	4	QL (60 EA per 30 days)
ARISTADA INITIO INTRAMUSCULAR PREFILLED SYRINGE 675 MG/2.4ML	5	PA; QL (4.8 ML per 365 days)
ARISTADA INTRAMUSCULAR PREFILLED SYRINGE 1064 MG/3.9ML	5	PA; QL (3.9 ML per 56 days)
ARISTADA INTRAMUSCULAR PREFILLED SYRINGE 441 MG/1.6ML	5	PA; QL (1.6 ML per 28 days)
ARISTADA INTRAMUSCULAR PREFILLED SYRINGE 662 MG/2.4ML	5	PA; QL (2.4 ML per 28 days)
ARISTADA INTRAMUSCULAR PREFILLED SYRINGE 882 MG/3.2ML	5	PA; QL (3.2 ML per 28 days)
<i>asenapine maleate sublingual tablet sublingual 10 mg, 2.5 mg, 5 mg</i>	2	QL (60 EA per 30 days)
CAPLYTA ORAL CAPSULE 10.5 MG, 21 MG, 42 MG	5	PA
ERZOFRI INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 117 MG/0.75ML	5	PA; QL (0.75 ML per 28 days)
ERZOFRI INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 156 MG/ML	5	PA; QL (1 ML per 28 days)
ERZOFRI INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 234 MG/1.5ML	5	PA; QL (1.5 ML per 28 days)
ERZOFRI INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 351 MG/2.25ML	5	PA; QL (2.25 ML per 28 days)
ERZOFRI INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 39 MG/0.25ML	4	PA; QL (0.25 ML per 28 days)
ERZOFRI INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 78 MG/0.5ML	5	PA; QL (0.5 ML per 28 days)
FANAPT ORAL TABLET 1 MG, 10 MG, 12 MG, 2 MG, 4 MG, 6 MG, 8 MG	5	PA; QL (60 EA per 30 days)
FANAPT TITRATION PACK A ORAL TABLET 1 & 2 & 4 & 6 MG	4	PA; QL (8 EA per 180 days)
FANAPT TITRATION PACK B ORAL TABLET 1 & 2 & 6 & 8 MG	4	PA; QL (12 EA per 180 days)
FANAPT TITRATION PACK C ORAL TABLET 1 & 2 & 6 MG	4	PA; QL (8 EA per 180 days)

Name of Drug	Drug Tier	Requirements/Limits
INVEGA HAFYERA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 1092 MG/3.5ML	5	PA; QL (3.5 ML per 180 days)
INVEGA HAFYERA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 1560 MG/5ML	5	PA; QL (5 ML per 180 days)
INVEGA SUSTENNA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 117 MG/0.75ML	5	PA; QL (0.75 ML per 28 days)
INVEGA SUSTENNA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 156 MG/ML	5	PA; QL (1 ML per 28 days)
INVEGA SUSTENNA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 234 MG/1.5ML	5	PA; QL (1.5 ML per 28 days)
INVEGA SUSTENNA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 39 MG/0.25ML	4	PA; QL (0.25 ML per 28 days)
INVEGA SUSTENNA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 78 MG/0.5ML	5	PA; QL (0.5 ML per 28 days)
INVEGA TRINZA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 273 MG/0.88ML	5	PA; QL (0.88 ML per 84 days)
INVEGA TRINZA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 410 MG/1.32ML	5	PA; QL (1.32 ML per 84 days)
INVEGA TRINZA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 546 MG/1.75ML	5	PA; QL (1.75 ML per 84 days)
INVEGA TRINZA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 819 MG/2.63ML	5	PA; QL (2.63 ML per 84 days)
<i>lurasidone hcl oral tablet 120 mg, 20 mg, 40 mg, 60 mg</i>	2	QL (30 EA per 30 days)
<i>lurasidone hcl oral tablet 80 mg</i>	2	QL (60 EA per 30 days)
LYBALVI ORAL TABLET 10-10 MG, 15-10 MG, 20-10 MG, 5-10 MG	5	PA
NUPLAZID ORAL CAPSULE 34 MG	5	PA; QL (30 EA per 30 days)
NUPLAZID ORAL TABLET 10 MG	5	PA; QL (30 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>olanzapine intramuscular solution reconstituted 10 mg</i>	4	QL (90 EA per 30 days)
<i>olanzapine oral tablet 10 mg, 15 mg, 2.5 mg, 20 mg, 5 mg, 7.5 mg</i>	1	QL (30 EA per 30 days)
<i>olanzapine oral tablet dispersible 10 mg, 15 mg, 20 mg, 5 mg</i>	2	QL (30 EA per 30 days)
OPIPZA ORAL FILM 10 MG, 5 MG	5	PA; QL (90 EA per 30 days)
OPIPZA ORAL FILM 2 MG	5	PA; QL (30 EA per 30 days)
<i>paliperidone er oral tablet extended release 24 hour 1.5 mg, 3 mg, 9 mg</i>	2	QL (30 EA per 30 days)
<i>paliperidone er oral tablet extended release 24 hour 6 mg</i>	2	QL (60 EA per 30 days)
PERSERIS SUBCUTANEOUS PREFILLED SYRINGE 120 MG, 90 MG	5	PA; QL (1 EA per 28 days)
<i>quetiapine fumarate er oral tablet extended release 24 hour 150 mg, 200 mg</i>	2	QL (30 EA per 30 days)
<i>quetiapine fumarate er oral tablet extended release 24 hour 300 mg, 400 mg, 50 mg</i>	2	QL (60 EA per 30 days)
<i>quetiapine fumarate oral tablet 100 mg, 150 mg, 200 mg, 300 mg, 400 mg</i>	1	QL (60 EA per 30 days)
<i>quetiapine fumarate oral tablet 25 mg, 50 mg</i>	1	QL (90 EA per 30 days)
REXULTI ORAL TABLET 0.25 MG, 0.5 MG, 1 MG, 2 MG, 3 MG, 4 MG	5	PA; QL (30 EA per 30 days)
<i>risperidone microspheres er intramuscular suspension reconstituted er 12.5 mg, 25 mg, 37.5 mg, 50 mg</i>	2	QL (2 EA per 28 days)
<i>risperidone oral solution 1 mg/ml</i>	2	QL (360 ML per 30 days)
<i>risperidone oral tablet 0.25 mg, 0.5 mg, 1 mg, 2 mg</i>	1	QL (60 EA per 30 days)
<i>risperidone oral tablet 3 mg</i>	1	QL (120 EA per 30 days)
<i>risperidone oral tablet 4 mg</i>	1	QL (90 EA per 30 days)
<i>risperidone oral tablet dispersible 0.25 mg, 0.5 mg, 1 mg, 2 mg</i>	2	QL (60 EA per 30 days)
<i>risperidone oral tablet dispersible 3 mg</i>	2	QL (120 EA per 30 days)
<i>risperidone oral tablet dispersible 4 mg</i>	2	QL (90 EA per 30 days)
RYKINDO INTRAMUSCULAR SUSPENSION RECONSTITUTED ER 25 MG, 37.5 MG, 50 MG	5	PA; QL (2 EA per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
SECUADO TRANSDERMAL PATCH 24 HOUR 3.8 MG/24HR, 5.7 MG/24HR, 7.6 MG/24HR	5	PA; QL (30 EA per 30 days)
UZEDY SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 100 MG/0.28ML	5	PA; QL (0.28 ML per 28 days)
UZEDY SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 125 MG/0.35ML	5	PA; QL (0.35 ML per 28 days)
UZEDY SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 150 MG/0.42ML	5	PA; QL (0.42 ML per 56 days)
UZEDY SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 200 MG/0.56ML	5	PA; QL (0.56 ML per 56 days)
UZEDY SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 250 MG/0.7ML	5	PA; QL (0.7 ML per 56 days)
UZEDY SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 50 MG/0.14ML	5	PA; QL (0.14 ML per 28 days)
UZEDY SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 75 MG/0.21ML	5	PA; QL (0.21 ML per 28 days)
VRAYLAR ORAL CAPSULE 0.5 MG, 0.75 MG, 1.5 MG, 3 MG, 4.5 MG, 6 MG	5	PA; QL (30 EA per 30 days)
<i>ziprasidone hcl oral capsule 20 mg, 40 mg, 60 mg, 80 mg</i>	2	QL (60 EA per 30 days)
<i>ziprasidone mesylate intramuscular solution reconstituted 20 mg</i>	2	QL (6 EA per 3 days)
Treatment-Resistant		
<i>clozapine oral tablet 100 mg</i>	2	QL (270 EA per 30 days)
<i>clozapine oral tablet 200 mg</i>	2	QL (120 EA per 30 days)
<i>clozapine oral tablet 25 mg, 50 mg</i>	2	QL (90 EA per 30 days)
<i>clozapine oral tablet dispersible 100 mg</i>	2	QL (270 EA per 30 days)
<i>clozapine oral tablet dispersible 12.5 mg</i>	2	
<i>clozapine oral tablet dispersible 150 mg</i>	2	QL (180 EA per 30 days)
<i>clozapine oral tablet dispersible 200 mg</i>	2	QL (120 EA per 30 days)
<i>clozapine oral tablet dispersible 25 mg</i>	2	QL (90 EA per 30 days)
VERSACLOZ ORAL SUSPENSION 50 MG/ML	4	QL (600 ML per 30 days)
Antispasticity Agents - Treatment Of Muscle Spasms		
Antispasticity Agents		
<i>baclofen oral tablet 10 mg, 20 mg, 5 mg</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>dantrolene sodium oral capsule 100 mg, 25 mg, 50 mg</i>	2	
<i>tizanidine hcl oral tablet 2 mg, 4 mg</i>	1	
Antivirals - Treatment Of Infections By Viruses		
Anti-Cytomegalovirus (Cmv) Agents		
LIVTENCITY ORAL TABLET 200 MG	5	PA
PREVYMIS ORAL PACKET 120 MG, 20 MG	5	PA
PREVYMIS ORAL TABLET 240 MG, 480 MG	5	PA
<i>valganciclovir hcl oral solution reconstituted 50 mg/ml</i>	5	
<i>valganciclovir hcl oral tablet 450 mg</i>	2	
Anti-Hepatitis B (Hbv) Agents		
<i>adefovir dipivoxil oral tablet 10 mg</i>	4	QL (30 EA per 30 days)
BARACLUDE ORAL SOLUTION 0.05 MG/ML	4	
<i>entecavir oral tablet 0.5 mg, 1 mg</i>	2	QL (30 EA per 30 days)
<i>lamivudine oral solution 10 mg/ml, 300 mg/30ml</i>	2	QL (960 ML per 30 days)
<i>lamivudine oral tablet 100 mg, 300 mg</i>	2	QL (30 EA per 30 days)
<i>lamivudine oral tablet 150 mg</i>	2	QL (60 EA per 30 days)
<i>tenofovir disoproxil fumarate oral tablet 300 mg</i>	2	QL (30 EA per 30 days)
VEMLIDY ORAL TABLET 25 MG	5	PA; QL (30 EA per 30 days)
VIREAD ORAL POWDER 40 MG/GM	5	QL (240 GM per 30 days)
VIREAD ORAL TABLET 150 MG, 200 MG, 250 MG	5	QL (30 EA per 30 days)
Anti-Hepatitis C (Hcv) Agents		
MAVYRET ORAL PACKET 50-20 MG	5	PA
MAVYRET ORAL TABLET 100-40 MG	5	PA
<i>ribavirin oral capsule 200 mg</i>	2	
<i>ribavirin oral tablet 200 mg</i>	2	
<i>sofosbuvir-velpatasvir oral tablet 400-100 mg</i>	5	PA
VOSEVI ORAL TABLET 400-100-100 MG	5	PA
Antitherpetic Agents		
<i>acyclovir oral capsule 200 mg</i>	1	
<i>acyclovir oral suspension 200 mg/5ml</i>	1	
<i>acyclovir oral tablet 400 mg, 800 mg</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>acyclovir sodium intravenous solution 50 mg/ml</i>	2	B/D
<i>famciclovir oral tablet 125 mg, 250 mg, 500 mg</i>	2	
<i>trifluridine ophthalmic solution 1 %</i>	2	
<i>valacyclovir hcl oral tablet 1 gm, 500 mg</i>	2	
Anti-Hiv Agents, Integrase Inhibitors (Insti)		
ISENTRESS HD ORAL TABLET 600 MG	5	QL (60 EA per 30 days)
ISENTRESS ORAL PACKET 100 MG	4	QL (60 EA per 30 days)
ISENTRESS ORAL TABLET 400 MG	5	QL (120 EA per 30 days)
ISENTRESS ORAL TABLET CHEWABLE 100 MG, 25 MG	4	QL (180 EA per 30 days)
TIVICAY ORAL TABLET 50 MG	5	QL (60 EA per 30 days)
TIVICAY PD ORAL TABLET SOLUBLE 5 MG	4	QL (180 EA per 30 days)
Anti-Hiv Agents, Non-Nucleoside Reverse Transcriptase Inhibitors (Nnrti)		
EDURANT PED ORAL TABLET SOLUBLE 2.5 MG	4	QL (180 EA per 30 days)
<i>efavirenz oral tablet 600 mg</i>	2	QL (30 EA per 30 days)
<i>etravirine oral tablet 100 mg</i>	2	QL (120 EA per 30 days)
<i>etravirine oral tablet 200 mg</i>	5	QL (60 EA per 30 days)
IDVYNZO ORAL TABLET 100-0.25 MG	5	QL (30 EA per 30 days)
INTELENCE ORAL TABLET 25 MG	4	QL (120 EA per 30 days)
<i>nevirapine er oral tablet extended release 24 hour 400 mg</i>	2	QL (30 EA per 30 days)
<i>nevirapine oral suspension 50 mg/5ml</i>	2	QL (1200 ML per 30 days)
<i>nevirapine oral tablet 200 mg</i>	1	QL (60 EA per 30 days)
PIFELTRO ORAL TABLET 100 MG	5	QL (30 EA per 30 days)
<i>rilpivirine hcl oral tablet 25 mg</i>	5	QL (30 EA per 30 days)
Anti-Hiv Agents, Nucleoside And Nucleotide Reverse Transcriptase Inhibitors (Nrti)		
<i>abacavir sulfate oral solution 20 mg/ml</i>	2	QL (960 ML per 30 days)
<i>abacavir sulfate oral tablet 300 mg</i>	2	QL (60 EA per 30 days)
<i>abacavir sulfate-lamivudine oral tablet 600-300 mg</i>	2	QL (30 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
CIMDUO ORAL TABLET 300-300 MG	5	QL (30 EA per 30 days)
DESCOVY ORAL TABLET 120-15 MG, 200-25 MG	5	QL (30 EA per 30 days)
<i>emtricitabine oral capsule 200 mg</i>	2	QL (30 EA per 30 days)
<i>emtricitabine-tenofovir df oral tablet 100-150 mg, 133-200 mg, 167-250 mg, 200-300 mg</i>	2	QL (30 EA per 30 days)
EMTRIVA ORAL SOLUTION 10 MG/ML	4	
<i>lamivudine-zidovudine oral tablet 150-300 mg</i>	2	QL (60 EA per 30 days)
<i>zidovudine oral capsule 100 mg</i>	2	QL (180 EA per 30 days)
<i>zidovudine oral syrup 50 mg/5ml</i>	2	QL (1920 ML per 30 days)
<i>zidovudine oral tablet 300 mg</i>	2	QL (90 EA per 30 days)
Anti-Hiv Agents, Other		
BIKTARVY ORAL TABLET 30-120-15 MG, 50-200-25 MG	5	QL (30 EA per 30 days)
DELSTRIGO ORAL TABLET 100-300-300 MG	5	QL (30 EA per 30 days)
DOVATO ORAL TABLET 50-300 MG	5	QL (30 EA per 30 days)
<i>efavirenz-emtricitab-tenofo df oral tablet 600-200-300 mg</i>	2	QL (30 EA per 30 days)
<i>efavirenz-lamivudine-tenofovir oral tablet 400-300-300 mg, 600-300-300 mg</i>	5	QL (30 EA per 30 days)
<i>emtricitab-rilpivir-tenofovf df oral tablet 200-25-300 mg</i>	5	QL (30 EA per 30 days)
EVOTAZ ORAL TABLET 300-150 MG	5	QL (30 EA per 30 days)
GENVOYA ORAL TABLET 150-150-200-10 MG	5	QL (30 EA per 30 days)
JULUCA ORAL TABLET 50-25 MG	5	QL (30 EA per 30 days)
<i>maraviroc oral tablet 150 mg</i>	5	QL (60 EA per 30 days)
<i>maraviroc oral tablet 300 mg</i>	5	QL (120 EA per 30 days)
ODEFSEY ORAL TABLET 200-25-25 MG	5	QL (30 EA per 30 days)
PREZCOBIX ORAL TABLET 675-150 MG, 800-150 MG	5	QL (30 EA per 30 days)
RUKOBIA ORAL TABLET EXTENDED RELEASE 12 HOUR 600 MG	5	QL (60 EA per 30 days)
SELZENTRY ORAL SOLUTION 20 MG/ML	3	QL (1840 ML per 30 days)
STRIBILD ORAL TABLET 150-150-200-300 MG	5	QL (30 EA per 30 days)
SUNLENCA ORAL TABLET 300 MG	5	QL (10 EA per 365 days)

Name of Drug	Drug Tier	Requirements/Limits
SUNLENCA ORAL TABLET THERAPY PACK 4 X 300 MG	5	QL (8 EA per 365 days)
SUNLENCA ORAL TABLET THERAPY PACK 5 X 300 MG	5	QL (10 EA per 365 days)
SUNLENCA SUBCUTANEOUS SOLUTION 463.5 MG/1.5ML	5	QL (6 ML per 365 days)
SYMTUZA ORAL TABLET 800-150-200-10 MG	5	QL (30 EA per 30 days)
TRIUMEQ ORAL TABLET 600-50-300 MG	5	QL (30 EA per 30 days)
<i>trimeq pd oral tablet soluble 60-5-30 mg</i>	2	QL (180 EA per 30 days)
TYBOST ORAL TABLET 150 MG	3	QL (30 EA per 30 days)
Anti-Hiv Agents, Protease Inhibitors (Pi)		
APTIVUS ORAL CAPSULE 250 MG	5	QL (120 EA per 30 days)
<i>atazanavir sulfate oral capsule 150 mg, 300 mg</i>	2	QL (30 EA per 30 days)
<i>atazanavir sulfate oral capsule 200 mg</i>	2	QL (60 EA per 30 days)
<i>darunavir oral tablet 600 mg</i>	2	QL (60 EA per 30 days)
<i>darunavir oral tablet 800 mg</i>	2	QL (30 EA per 30 days)
<i>fosamprenavir calcium oral tablet 700 mg</i>	5	QL (120 EA per 30 days)
KALETRA ORAL SOLUTION 400-100 MG/5ML	4	QL (390 ML per 30 days)
<i>lopinavir-ritonavir oral tablet 100-25 mg</i>	2	QL (300 EA per 30 days)
<i>lopinavir-ritonavir oral tablet 200-50 mg</i>	2	QL (120 EA per 30 days)
NORVIR ORAL PACKET 100 MG	4	QL (360 EA per 30 days)
PREZISTA ORAL SUSPENSION 100 MG/ML	5	QL (400 ML per 30 days)
PREZISTA ORAL TABLET 150 MG	4	QL (180 EA per 30 days)
PREZISTA ORAL TABLET 75 MG	4	QL (300 EA per 30 days)
REYATAZ ORAL PACKET 50 MG	4	
<i>ritonavir oral tablet 100 mg</i>	2	QL (360 EA per 30 days)
VIRACEPT ORAL TABLET 250 MG	5	QL (300 EA per 30 days)
VIRACEPT ORAL TABLET 625 MG	5	QL (120 EA per 30 days)
Anti-Influenza Agents		
<i>oseltamivir phosphate oral capsule 30 mg</i>	2	QL (84 EA per 180 days)
<i>oseltamivir phosphate oral capsule 45 mg, 75 mg</i>	2	QL (42 EA per 180 days)
<i>oseltamivir phosphate oral suspension reconstituted 6 mg/ml</i>	2	QL (540 ML per 180 days)

Name of Drug	Drug Tier	Requirements/Limits
RELENZA DISKHALER INHALATION AEROSOL POWDER BREATH ACTIVATED 5 MG/ACT	4	QL (60 EA per 180 days)
<i>rimantadine hcl oral tablet 100 mg</i>	2	
Antiviral, Coronavirus Agents		
LAGEVRIO ORAL CAPSULE 200 MG	3	QL (40 EA per 5 days)
PAXLOVID (150/100) ORAL TABLET THERAPY PACK 10 X 150 MG & 10 X 100MG	3	QL (20 EA per 5 days)
PAXLOVID (300/100 & 150/100) ORAL TABLET THERAPY PACK 6 X 150 MG & 5 X 100MG	3	QL (11 EA per 5 days)
PAXLOVID (300/100) ORAL TABLET THERAPY PACK 20 X 150 MG & 10 X 100MG	3	QL (30 EA per 5 days)
Anxiolytics - Treatment Of Anxiety Or Nervousness		
Anxiolytics, Other		
<i>buspirone hcl oral tablet 10 mg, 15 mg, 30 mg, 5 mg, 7.5 mg</i>	1	
<i>hydroxyzine pamoate oral capsule 100 mg, 25 mg, 50 mg</i>	2	
Benzodiazepines		
ALPRAZOLAM INTENSOL ORAL CONCENTRATE 1 MG/ML	2	QL (300 ML per 30 days)
<i>alprazolam oral tablet 0.25 mg, 0.5 mg, 1 mg</i>	2	QL (120 EA per 30 days)
<i>alprazolam oral tablet 2 mg</i>	2	QL (150 EA per 30 days)
<i>clonazepam oral tablet 0.5 mg, 1 mg</i>	2	QL (90 EA per 30 days)
<i>clonazepam oral tablet 2 mg</i>	2	QL (300 EA per 30 days)
<i>clonazepam oral tablet dispersible 0.125 mg, 0.25 mg, 0.5 mg, 1 mg</i>	2	QL (90 EA per 30 days)
<i>clonazepam oral tablet dispersible 2 mg</i>	2	QL (300 EA per 30 days)
<i>clorazepate dipotassium oral tablet 15 mg</i>	2	QL (180 EA per 30 days)
<i>clorazepate dipotassium oral tablet 3.75 mg, 7.5 mg</i>	2	QL (90 EA per 30 days)
DIAZEPAM INTENSOL ORAL CONCENTRATE 5 MG/ML	2	QL (240 ML per 30 days)
<i>diazepam oral concentrate 5 mg/ml</i>	2	QL (240 ML per 30 days)
<i>diazepam oral solution 5 mg/5ml</i>	2	QL (1200 ML per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>diazepam oral tablet 10 mg, 2 mg, 5 mg</i>	2	QL (120 EA per 30 days)
LORAZEPAM INTENSOL ORAL CONCENTRATE 2 MG/ML	2	QL (150 ML per 30 days)
<i>lorazepam oral concentrate 2 mg/ml</i>	2	QL (150 ML per 30 days)
<i>lorazepam oral tablet 0.5 mg, 1 mg</i>	2	QL (90 EA per 30 days)
<i>lorazepam oral tablet 2 mg</i>	2	QL (150 EA per 30 days)
Bipolar Agents - Treatment For Bipolar Illnesses		
Mood Stabilizers		
EQUETRO ORAL CAPSULE EXTENDED RELEASE 12 HOUR 100 MG, 200 MG, 300 MG	4	
<i>lithium carbonate er oral tablet extended release 300 mg, 450 mg</i>	2	
<i>lithium carbonate oral capsule 150 mg, 300 mg, 600 mg</i>	2	
<i>lithium carbonate oral tablet 300 mg</i>	2	
<i>lithium oral solution 8 meq/5ml</i>	2	
Blood Glucose Regulators - Control Of Diabetes		
Antidiabetic Agents		
<i>acarbose oral tablet 100 mg, 25 mg, 50 mg</i>	2	QL (90 EA per 30 days)
<i>dapagliflozin oral tablet 10 mg, 5 mg</i>	2	QL (30 EA per 30 days)
<i>glimepiride oral tablet 1 mg</i>	1	QL (240 EA per 30 days)
<i>glimepiride oral tablet 2 mg</i>	1	QL (120 EA per 30 days)
<i>glimepiride oral tablet 4 mg</i>	1	QL (60 EA per 30 days)
<i>glipizide er oral tablet extended release 24 hour 10 mg</i>	1	QL (60 EA per 30 days)
<i>glipizide er oral tablet extended release 24 hour 2.5 mg</i>	1	QL (240 EA per 30 days)
<i>glipizide er oral tablet extended release 24 hour 5 mg</i>	1	QL (120 EA per 30 days)
<i>glipizide oral tablet 10 mg</i>	1	QL (120 EA per 30 days)
<i>glipizide oral tablet 15 mg, 2.5 mg</i>	1	QL (60 EA per 30 days)
<i>glipizide oral tablet 5 mg</i>	1	QL (240 EA per 30 days)
<i>glipizide-metformin hcl oral tablet 2.5-250 mg</i>	1	QL (240 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>glipizide-metformin hcl oral tablet 2.5-500 mg, 5-500 mg</i>	1	QL (120 EA per 30 days)
<i>glyburide micronized oral tablet 3 mg</i>	1	PA; QL (90 EA per 30 days)
<i>glyburide micronized oral tablet 6 mg</i>	1	PA; QL (60 EA per 30 days)
<i>glyburide oral tablet 1.25 mg, 2.5 mg</i>	1	PA; QL (60 EA per 30 days)
<i>glyburide oral tablet 5 mg</i>	1	PA; QL (120 EA per 30 days)
<i>glyburide-metformin oral tablet 1.25-250 mg</i>	2	PA; QL (240 EA per 30 days)
<i>glyburide-metformin oral tablet 2.5-500 mg, 5-500 mg</i>	2	PA; QL (120 EA per 30 days)
GLYXAMBI ORAL TABLET 10-5 MG, 25-5 MG	3	QL (30 EA per 30 days)
JANUMET ORAL TABLET 50-1000 MG, 50-500 MG	3	QL (60 EA per 30 days)
JANUMET XR ORAL TABLET EXTENDED RELEASE 24 HOUR 100-1000 MG	3	QL (30 EA per 30 days)
JANUMET XR ORAL TABLET EXTENDED RELEASE 24 HOUR 50-1000 MG, 50-500 MG	3	QL (60 EA per 30 days)
JANUVIA ORAL TABLET 100 MG, 25 MG, 50 MG	3	QL (30 EA per 30 days)
JARDIANCE ORAL TABLET 10 MG, 25 MG	3	QL (30 EA per 30 days)
JENTADUETO ORAL TABLET 2.5-1000 MG, 2.5-500 MG, 2.5-850 MG	3	QL (60 EA per 30 days)
JENTADUETO XR ORAL TABLET EXTENDED RELEASE 24 HOUR 2.5-1000 MG	3	QL (60 EA per 30 days)
JENTADUETO XR ORAL TABLET EXTENDED RELEASE 24 HOUR 5-1000 MG	3	QL (30 EA per 30 days)
<i>liraglutide subcutaneous solution pen-injector 18 mg/3ml</i>	2	PA; QL (9 ML per 30 days)
<i>metformin hcl er oral tablet extended release 24 hour 500 mg</i>	1	QL (120 EA per 30 days)
<i>metformin hcl er oral tablet extended release 24 hour 750 mg</i>	1	QL (60 EA per 30 days)
<i>metformin hcl oral tablet 1000 mg</i>	1	QL (75 EA per 30 days)
<i>metformin hcl oral tablet 500 mg</i>	1	QL (150 EA per 30 days)
<i>metformin hcl oral tablet 850 mg</i>	1	QL (90 EA per 30 days)
MOUNJARO SUBCUTANEOUS SOLUTION AUTO-INJECTOR 10 MG/0.5ML, 12.5 MG/0.5ML, 15 MG/0.5ML, 2.5 MG/0.5ML, 5 MG/0.5ML, 7.5 MG/0.5ML	3	PA; QL (2 ML per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>nateglinide oral tablet 120 mg, 60 mg</i>	2	QL (90 EA per 30 days)
OZEMPIC (0.25 OR 0.5 MG/DOSE) SUBCUTANEOUS SOLUTION PEN-INJECTOR 2 MG/3ML	3	PA; QL (3 ML per 28 days)
OZEMPIC (1 MG/DOSE) SUBCUTANEOUS SOLUTION PEN-INJECTOR 4 MG/3ML	3	PA; QL (3 ML per 28 days)
OZEMPIC (2 MG/DOSE) SUBCUTANEOUS SOLUTION PEN-INJECTOR 8 MG/3ML	3	PA; QL (3 ML per 28 days)
<i>pioglitazone hcl oral tablet 15 mg, 30 mg, 45 mg</i>	2	QL (30 EA per 30 days)
<i>pioglitazone hcl-metformin hcl oral tablet 15-500 mg, 15-850 mg</i>	2	QL (90 EA per 30 days)
<i>repaglinide oral tablet 0.5 mg, 1 mg</i>	2	QL (120 EA per 30 days)
<i>repaglinide oral tablet 2 mg</i>	2	QL (240 EA per 30 days)
RYBELSUS ORAL TABLET 14 MG, 3 MG, 7 MG	3	PA; QL (30 EA per 30 days)
SYMLINPEN 120 SUBCUTANEOUS SOLUTION PEN-INJECTOR 2700 MCG/2.7ML	5	
SYMLINPEN 60 SUBCUTANEOUS SOLUTION PEN-INJECTOR 1500 MCG/1.5ML	5	
SYNJARDY ORAL TABLET 12.5-1000 MG, 12.5-500 MG, 5-1000 MG, 5-500 MG	3	QL (60 EA per 30 days)
SYNJARDY XR ORAL TABLET EXTENDED RELEASE 24 HOUR 10-1000 MG, 12.5-1000 MG, 5-1000 MG	3	QL (60 EA per 30 days)
SYNJARDY XR ORAL TABLET EXTENDED RELEASE 24 HOUR 25-1000 MG	3	QL (30 EA per 30 days)
TRADJENTA ORAL TABLET 5 MG	3	QL (30 EA per 30 days)
TRIJARDY XR ORAL TABLET EXTENDED RELEASE 24 HOUR 10-5-1000 MG, 25-5-1000 MG	3	QL (30 EA per 30 days)
TRIJARDY XR ORAL TABLET EXTENDED RELEASE 24 HOUR 12.5-2.5-1000 MG, 5-2.5-1000 MG	3	QL (60 EA per 30 days)
TRULICITY SUBCUTANEOUS SOLUTION AUTO-INJECTOR 0.75 MG/0.5ML, 1.5 MG/0.5ML, 3 MG/0.5ML, 4.5 MG/0.5ML	3	PA; QL (2 ML per 28 days)
XIGDUO XR ORAL TABLET EXTENDED RELEASE 24 HOUR 10-1000 MG, 10-500 MG, 5-500 MG	3	QL (30 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
XIGDUO XR ORAL TABLET EXTENDED RELEASE 24 HOUR 2.5-1000 MG, 5-1000 MG	3	QL (60 EA per 30 days)
Glycemic Agents		
BAQSIMI ONE PACK NASAL POWDER 3 MG/DOSE	3	QL (4 EA per 30 days)
BAQSIMI TWO PACK NASAL POWDER 3 MG/DOSE	3	QL (4 EA per 30 days)
<i>diazoxide oral suspension 50 mg/ml</i>	2	
<i>glucagon emergency injection solution reconstituted 1 mg, 1 mg/ml</i>	3	QL (4 EA per 30 days)
<i>mifepristone oral tablet 300 mg</i>	5	PA
Insulins		
FIASP FLEXTOUCH SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 UNIT/ML	3	
FIASP INJECTION SOLUTION 100 UNIT/ML	3	
FIASP PENFILL SUBCUTANEOUS SOLUTION CARTRIDGE 100 UNIT/ML	3	
<i>gauze pad 2"x2"</i>	1	
GAUZE PAD 2"X2"	1	
HUMALOG INJECTION SOLUTION 100 UNIT/ML	3	
HUMALOG JUNIOR KWIKPEN SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 UNIT/ML	3	
HUMALOG KWIKPEN SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 UNIT/ML, 200 UNIT/ML	3	
HUMALOG MIX 50/50 KWIKPEN SUBCUTANEOUS SUSPENSION PEN-INJECTOR (50-50) 100 UNIT/ML	3	
HUMALOG MIX 75/25 KWIKPEN SUBCUTANEOUS SUSPENSION PEN-INJECTOR (75-25) 100 UNIT/ML	3	
HUMALOG MIX 75/25 SUBCUTANEOUS SUSPENSION (75-25) 100 UNIT/ML	3	
HUMALOG SUBCUTANEOUS SOLUTION CARTRIDGE 100 UNIT/ML	3	
HUMALOG TEMPO PEN SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 UNIT/ML	3	

Name of Drug	Drug Tier	Requirements/Limits
HUMULIN 70/30 KWIKPEN SUBCUTANEOUS SUSPENSION PEN- INJECTOR (70-30) 100 UNIT/ML	3	
HUMULIN 70/30 SUBCUTANEOUS SUSPENSION (70-30) 100 UNIT/ML	3	
HUMULIN N KWIKPEN SUBCUTANEOUS SUSPENSION PEN-INJECTOR 100 UNIT/ML	3	
HUMULIN N SUBCUTANEOUS SUSPENSION 100 UNIT/ML	3	
HUMULIN R INJECTION SOLUTION 100 UNIT/ML	3	
HUMULIN R U-500 (CONCENTRATED) SUBCUTANEOUS SOLUTION 500 UNIT/ML	3	
HUMULIN R U-500 KWIKPEN SUBCUTANEOUS SOLUTION PEN- INJECTOR 500 UNIT/ML	3	
<i>insulin asp prot & asp flexpen subcutaneous suspension pen-injector (70-30) 100 unit/ml</i>	2	
<i>insulin aspart flexpen subcutaneous solution pen- injector 100 unit/ml</i>	2	
<i>insulin aspart injection solution 100 unit/ml</i>	2	
<i>insulin aspart prot & aspart subcutaneous suspension (70-30) 100 unit/ml</i>	2	
<i>insulin lispro (1 unit dial) subcutaneous solution pen-injector 100 unit/ml</i>	2	
<i>insulin lispro injection solution 100 unit/ml</i>	2	
<i>insulin lispro junior kwikpen subcutaneous solution pen-injector 100 unit/ml</i>	2	
<i>insulin lispro prot & lispro subcutaneous suspension pen-injector (75-25) 100 unit/ml</i>	2	
<i>insulin syringe 27g x 1/2" 0.5 ml, 27g x 1/2" 1 ml, 28g x 1/2" 0.5 ml, 28g x 1/2" 1 ml, 29g x 1/2" 0.3 ml, 29g x 1/2" 0.5 ml, 29g x 1/2" 1 ml, 29g x 5/16" 1 ml, 30g x 1/2" 0.3 ml, 30g x 1/2" 0.5 ml, 30g x 1/2" 1 ml, 30g x 5/16" 0.3 ml, 30g x 5/16" 0.5 ml, 30g x 5/16" 1 ml, 31g x 1/2" 0.3 ml, 31g x 1/4" 0.3 ml, 31g x 1/4" 0.5 ml, 31g x 1/4" 1 ml, 31g x 5/16" 0.3 ml, 31g x 5/16" 0.5 ml, 31g x 5/16" 1 ml</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
INSULIN SYRINGE 27G X 1/2" 1 ML, 27G X 5/8" 1 ML, 28G X 1/2" 0.5 ML, 28G X 1/2" 1 ML, 29G 0.3 ML, 29G X 1/2" 0.3 ML, 29G X 1/2" 0.5 ML, 29G X 1/2" 1 ML, 30G X 1/2" 0.3 ML, 30G X 1/2" 0.5 ML, 30G X 1/2" 1 ML, 30G X 5/16" 0.3 ML, 30G X 5/16" 0.5 ML, 30G X 5/16" 1 ML, 31G X 15/64" 0.3 ML, 31G X 15/64" 0.5 ML, 31G X 15/64" 1 ML, 31G X 5/16" 0.3 ML, 31G X 5/16" 0.5 ML, 31G X 5/16" 1 ML, 31G X 6MM 0.5 ML, U-100 1 ML	1	
LANTUS SOLOSTAR SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 UNIT/ML	3	
LANTUS SUBCUTANEOUS SOLUTION 100 UNIT/ML	3	
NOVOLIN 70/30 FLEXPEN RELION SUBCUTANEOUS SUSPENSION PEN-INJECTOR (70-30) 100 UNIT/ML	3	
NOVOLIN 70/30 FLEXPEN SUBCUTANEOUS SUSPENSION PEN-INJECTOR (70-30) 100 UNIT/ML	3	
NOVOLIN 70/30 RELION SUBCUTANEOUS SUSPENSION (70-30) 100 UNIT/ML	3	
NOVOLIN 70/30 SUBCUTANEOUS SUSPENSION (70-30) 100 UNIT/ML	3	
NOVOLIN N FLEXPEN RELION SUBCUTANEOUS SUSPENSION PEN-INJECTOR 100 UNIT/ML	3	
NOVOLIN N FLEXPEN SUBCUTANEOUS SUSPENSION PEN-INJECTOR 100 UNIT/ML	3	
NOVOLIN N RELION SUBCUTANEOUS SUSPENSION 100 UNIT/ML	3	
NOVOLIN N SUBCUTANEOUS SUSPENSION 100 UNIT/ML	3	
NOVOLIN R FLEXPEN INJECTION SOLUTION PEN-INJECTOR 100 UNIT/ML	3	
NOVOLIN R FLEXPEN RELION INJECTION SOLUTION PEN-INJECTOR 100 UNIT/ML	3	
NOVOLIN R INJECTION SOLUTION 100 UNIT/ML	3	
NOVOLIN R RELION INJECTION SOLUTION 100 UNIT/ML	3	

Name of Drug	Drug Tier	Requirements/Limits
NOVOLOG 70/30 FLEXPEN RELION SUBCUTANEOUS SUSPENSION PEN-INJECTOR (70-30) 100 UNIT/ML	3	
NOVOLOG FLEXPEN RELION SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 UNIT/ML	3	
NOVOLOG FLEXPEN SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 UNIT/ML	3	
NOVOLOG INJECTION SOLUTION 100 UNIT/ML	3	
NOVOLOG MIX 70/30 FLEXPEN SUBCUTANEOUS SUSPENSION PEN-INJECTOR (70-30) 100 UNIT/ML	3	
NOVOLOG MIX 70/30 RELION SUBCUTANEOUS SUSPENSION (70-30) 100 UNIT/ML	3	
NOVOLOG MIX 70/30 SUBCUTANEOUS SUSPENSION (70-30) 100 UNIT/ML	3	
NOVOLOG PENFILL SUBCUTANEOUS SOLUTION CARTRIDGE 100 UNIT/ML	3	
NOVOLOG RELION INJECTION SOLUTION 100 UNIT/ML	3	
OMNIPOD 5 DEXG7G6 INTRO GEN 5 KIT	3	
OMNIPOD 5 DEXG7G6 PODS GEN 5	3	
OMNIPOD 5 G7 INTRO (GEN 5) KIT	3	
OMNIPOD 5 G7 PODS (GEN 5)	3	
OMNIPOD 5 LIBRE INTRO KIT	3	
OMNIPOD 5 LIBRE PODS	3	
OMNIPOD DASH INTRO (GEN 4) KIT	3	
OMNIPOD DASH PODS (GEN 4)	3	
PEN NEEDLES 29G X 12.7MM , 29G X 12MM , 30G X 5 MM , 30G X 8 MM , 31G X 4 MM , 31G X 5 MM , 31G X 6 MM , 31G X 8 MM , 32G X 4 MM , 32G X 6 MM	1	
<i>pen needles 29g x 4mm , 30g x 5 mm , 31g x 5 mm , 31g x 6 mm , 31g x 8 mm , 32g x 4 mm , 32g x 5 mm , 32g x 6 mm , 32g x 8 mm</i>	1	
SOLIQUA SUBCUTANEOUS SOLUTION PEN-INJECTOR 100-33 UNT-MCG/ML	3	QL (15 ML per 25 days)

Name of Drug	Drug Tier	Requirements/Limits
TOUJEO MAX SOLOSTAR SUBCUTANEOUS SOLUTION PEN-INJECTOR 300 UNIT/ML	3	
TOUJEO SOLOSTAR SUBCUTANEOUS SOLUTION PEN-INJECTOR 300 UNIT/ML	3	
Blood Products And Modifiers - Prevention Of Clotting And Increasing Blood Cell Production		
Anticoagulants		
<i>dabigatran etexilate mesylate oral capsule 110 mg, 150 mg, 75 mg</i>	2	QL (60 EA per 30 days)
ELIQUIS (1.5 MG PACK) ORAL TABLET SOLUBLE 3 X 0.5 MG	3	
ELIQUIS (2 MG PACK) ORAL TABLET SOLUBLE 4 X 0.5 MG	3	
ELIQUIS DVT/PE STARTER PACK ORAL TABLET THERAPY PACK 5 MG	3	QL (148 EA per 365 days)
ELIQUIS ORAL CAPSULE SPRINKLE 0.15 MG	3	QL (74 EA per 30 days)
ELIQUIS ORAL TABLET 2.5 MG	3	QL (60 EA per 30 days)
ELIQUIS ORAL TABLET 5 MG	3	QL (74 EA per 30 days)
ELIQUIS ORAL TABLET SOLUBLE 0.5 MG	3	
<i>enoxaparin sodium injection solution 300 mg/3ml</i>	2	
<i>enoxaparin sodium injection solution prefilled syringe 100 mg/ml, 120 mg/0.8ml, 150 mg/ml, 30 mg/0.3ml, 40 mg/0.4ml, 60 mg/0.6ml, 80 mg/0.8ml</i>	2	
<i>fondaparinux sodium subcutaneous solution 10 mg/0.8ml, 5 mg/0.4ml, 7.5 mg/0.6ml</i>	5	
<i>fondaparinux sodium subcutaneous solution 2.5 mg/0.5ml</i>	2	
<i>heparin sodium (porcine) injection solution 10000 unit/ml, 5000 unit/ml</i>	2	
<i>heparin sodium (porcine) pf injection solution 1000 unit/ml</i>	2	
JANTOVEN ORAL TABLET 1 MG, 10 MG, 2 MG, 2.5 MG, 3 MG, 4 MG, 5 MG, 6 MG, 7.5 MG	1	
<i>warfarin sodium oral tablet 1 mg, 10 mg, 2 mg, 2.5 mg, 3 mg, 4 mg, 5 mg, 6 mg, 7.5 mg</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
XARELTO ORAL SUSPENSION RECONSTITUTED 1 MG/ML	3	QL (900 ML per 30 days)
XARELTO ORAL TABLET 10 MG, 20 MG	3	QL (30 EA per 30 days)
XARELTO ORAL TABLET 15 MG, 2.5 MG	3	QL (60 EA per 30 days)
XARELTO STARTER PACK ORAL TABLET THERAPY PACK 15 & 20 MG	3	QL (102 EA per 365 days)
Blood Products And Modifiers, Other		
<i>anagrelide hcl oral capsule 0.5 mg, 1 mg</i>	2	
ARANESP (ALBUMIN FREE) INJECTION SOLUTION 100 MCG/ML, 200 MCG/ML	5	PA
ARANESP (ALBUMIN FREE) INJECTION SOLUTION 25 MCG/ML, 40 MCG/ML, 60 MCG/ML	4	PA
ARANESP (ALBUMIN FREE) INJECTION SOLUTION PREFILLED SYRINGE 10 MCG/0.4ML, 25 MCG/0.42ML, 40 MCG/0.4ML	4	PA
ARANESP (ALBUMIN FREE) INJECTION SOLUTION PREFILLED SYRINGE 100 MCG/0.5ML, 150 MCG/0.3ML, 200 MCG/0.4ML, 300 MCG/0.6ML, 500 MCG/ML, 60 MCG/0.3ML	5	PA
<i>eltrombopag olamine oral packet 12.5 mg</i>	5	PA; QL (360 EA per 30 days)
<i>eltrombopag olamine oral packet 25 mg</i>	5	PA; QL (180 EA per 30 days)
<i>eltrombopag olamine oral tablet 12.5 mg, 25 mg</i>	5	PA; QL (30 EA per 30 days)
<i>eltrombopag olamine oral tablet 50 mg, 75 mg</i>	5	PA; QL (60 EA per 30 days)
EPOGEN INJECTION SOLUTION 10000 UNIT/ML, 2000 UNIT/ML, 20000 UNIT/ML, 3000 UNIT/ML, 4000 UNIT/ML	4	PA
FULPHILA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 6 MG/0.6ML	5	PA
FYLNETRA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 6 MG/0.6ML	5	PA
LEUKINE INJECTION SOLUTION RECONSTITUTED 250 MCG	4	PA
NEULASTA ONPRO SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 6 MG/0.6ML	5	PA
NEULASTA SUBCUTANEOUS SOLUTION 4 MG/0.4ML	5	PA

Name of Drug	Drug Tier	Requirements/Limits
NEULASTA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 6 MG/0.6ML	5	PA
PROCRIT INJECTION SOLUTION 10000 UNIT/ML, 2000 UNIT/ML, 3000 UNIT/ML, 4000 UNIT/ML	4	PA
PROCRIT INJECTION SOLUTION 20000 UNIT/ML, 40000 UNIT/ML	5	PA
PYRUKYND ORAL TABLET 20 MG, 5 MG, 50 MG	5	PA
PYRUKYND TAPER PACK ORAL TABLET THERAPY PACK 5 MG, 7 X 20 MG & 7 X 5 MG, 7 X 50 MG & 7 X 20 MG	5	PA
RETACRIT INJECTION SOLUTION 10000 UNIT/ML, 10000 UNIT/ML(1ML), 2000 UNIT/ML, 20000 UNIT/ML, 3000 UNIT/ML, 4000 UNIT/ML, 40000 UNIT/ML	4	PA
TAVNEOS ORAL CAPSULE 10 MG	5	PA
<i>tranexamic acid oral tablet 650 mg</i>	2	
XOLREMDI ORAL CAPSULE 100 MG	5	PA
ZARXIO INJECTION SOLUTION PREFILLED SYRINGE 300 MCG/0.5ML, 480 MCG/0.8ML	5	PA
Platelet Modifying Agents		
<i>aspirin-dipyridamole er oral capsule extended release 12 hour 25-200 mg</i>	2	
BRILINTA ORAL TABLET 90 MG	4	
<i>cilostazol oral tablet 100 mg, 50 mg</i>	1	
<i>clopidogrel bisulfate oral tablet 75 mg</i>	1	
<i>dipyridamole oral tablet 25 mg, 50 mg, 75 mg</i>	2	PA
DOPTELET ORAL TABLET 20 MG, 20 MG (10 PACK), 20 MG(15 PACK)	5	PA
DOPTELET SPRINKLE ORAL CAPSULE SPRINKLE 10 MG	5	PA
<i>prasugrel hcl oral tablet 10 mg, 5 mg</i>	2	
<i>ticagrelor oral tablet 60 mg, 90 mg</i>	2	
Cardiovascular Agents - Treatment Of Conditions Affecting The Heart And Blood Vessels		
Alpha-Adrenergic Agonists		

Name of Drug	Drug Tier	Requirements/Limits
<i>clonidine hcl oral tablet 0.1 mg, 0.2 mg, 0.3 mg</i>	1	
<i>clonidine transdermal patch weekly 0.1 mg/24hr, 0.2 mg/24hr, 0.3 mg/24hr</i>	2	
<i>droxidopa oral capsule 100 mg, 200 mg, 300 mg</i>	2	
<i>guanfacine hcl oral tablet 1 mg, 2 mg</i>	1	PA
<i>midodrine hcl oral tablet 10 mg, 2.5 mg, 5 mg</i>	2	
Alpha-Adrenergic Blocking Agents		
<i>doxazosin mesylate oral tablet 1 mg, 2 mg, 4 mg, 8 mg</i>	1	
<i>phenoxybenzamine hcl oral capsule 10 mg</i>	5	PA
<i>prazosin hcl oral capsule 1 mg, 2 mg, 5 mg</i>	1	
<i>terazosin hcl oral capsule 1 mg, 10 mg, 2 mg, 5 mg</i>	1	
Angiotensin II Receptor Antagonists		
<i>candesartan cilexetil oral tablet 16 mg, 32 mg, 4 mg, 8 mg</i>	2	
<i>irbesartan oral tablet 150 mg, 300 mg, 75 mg</i>	1	
<i>losartan potassium oral tablet 100 mg, 25 mg, 50 mg</i>	1	
<i>olmesartan medoxomil oral tablet 20 mg, 40 mg, 5 mg</i>	1	
<i>telmisartan oral tablet 20 mg, 40 mg, 80 mg</i>	1	
<i>valsartan oral tablet 160 mg, 320 mg, 40 mg, 80 mg</i>	1	
Angiotensin-Converting Enzyme (Ace) Inhibitors		
<i>benazepril hcl oral tablet 10 mg, 20 mg, 40 mg, 5 mg</i>	1	
<i>captopril oral tablet 100 mg, 12.5 mg, 25 mg, 50 mg</i>	2	
<i>enalapril maleate oral tablet 10 mg, 2.5 mg, 20 mg, 5 mg</i>	1	
<i>fosinopril sodium oral tablet 10 mg, 20 mg, 40 mg</i>	1	
<i>lisinopril oral tablet 10 mg, 2.5 mg, 20 mg, 30 mg, 40 mg, 5 mg</i>	1	
<i>moexipril hcl oral tablet 15 mg, 7.5 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>perindopril erbumine oral tablet 2 mg, 4 mg, 8 mg</i>	2	
<i>quinapril hcl oral tablet 10 mg, 20 mg, 40 mg, 5 mg</i>	1	
<i>ramipril oral capsule 1.25 mg, 10 mg, 2.5 mg, 5 mg</i>	1	
<i>trandolapril oral tablet 1 mg, 2 mg, 4 mg</i>	1	
Antiarrhythmics		
<i>amiodarone hcl oral tablet 100 mg, 200 mg, 400 mg</i>	2	
<i>disopyramide phosphate oral capsule 100 mg, 150 mg</i>	2	
<i>dofetilide oral capsule 125 mcg, 250 mcg, 500 mcg</i>	2	
<i>flecainide acetate oral tablet 100 mg, 150 mg, 50 mg</i>	2	
<i>mexiletine hcl oral capsule 150 mg, 200 mg, 250 mg</i>	2	
NORPACE CR ORAL CAPSULE EXTENDED RELEASE 12 HOUR 100 MG, 150 MG	4	
<i>propafenone hcl oral tablet 150 mg, 225 mg, 300 mg</i>	2	
<i>quinidine gluconate er oral tablet extended release 324 mg</i>	2	
<i>quinidine sulfate oral tablet 200 mg, 300 mg</i>	2	
<i>sotalol hcl (af) oral tablet 120 mg, 160 mg, 80 mg</i>	1	
<i>sotalol hcl oral tablet 120 mg, 160 mg, 240 mg, 80 mg</i>	1	
Beta-Adrenergic Blocking Agents		
<i>acebutolol hcl oral capsule 200 mg, 400 mg</i>	1	
<i>atenolol oral tablet 100 mg, 25 mg, 50 mg</i>	1	
<i>betaxolol hcl oral tablet 10 mg, 20 mg</i>	2	
<i>bisoprolol fumarate oral tablet 10 mg, 5 mg</i>	1	
<i>carvedilol oral tablet 12.5 mg, 25 mg, 3.125 mg, 6.25 mg</i>	1	
<i>labetalol hcl oral tablet 100 mg, 200 mg, 300 mg</i>	2	
<i>metoprolol succinate er oral tablet extended release 24 hour 100 mg, 200 mg, 25 mg, 50 mg</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>metoprolol tartrate oral tablet 100 mg, 25 mg, 50 mg</i>	1	
<i>nadolol oral tablet 20 mg, 40 mg, 80 mg</i>	2	
<i>nebivolol hcl oral tablet 10 mg, 2.5 mg, 5 mg</i>	1	QL (30 EA per 30 days)
<i>nebivolol hcl oral tablet 20 mg</i>	1	QL (60 EA per 30 days)
<i>pindolol oral tablet 10 mg, 5 mg</i>	2	
<i>propranolol hcl er oral capsule extended release 24 hour 120 mg, 160 mg, 60 mg, 80 mg</i>	1	
<i>propranolol hcl oral solution 20 mg/5ml, 40 mg/5ml</i>	2	
<i>propranolol hcl oral tablet 10 mg, 20 mg, 40 mg, 60 mg, 80 mg</i>	1	
<i>timolol maleate oral tablet 10 mg, 20 mg, 5 mg</i>	2	
Calcium Channel Blocking Agents, Dihydropyridines		
<i>amlodipine besylate oral tablet 10 mg, 2.5 mg, 5 mg</i>	1	
<i>felodipine er oral tablet extended release 24 hour 10 mg, 2.5 mg, 5 mg</i>	2	
<i>isradipine oral capsule 2.5 mg, 5 mg</i>	2	
<i>nifedipine er oral tablet extended release 24 hour 30 mg, 60 mg, 90 mg</i>	1	
<i>nifedipine er osmotic release oral tablet extended release 24 hour 30 mg, 60 mg, 90 mg</i>	1	
<i>nifedipine oral capsule 10 mg, 20 mg</i>	2	PA
<i>nimodipine oral capsule 30 mg</i>	2	
Calcium Channel Blocking Agents, Nondihydropyridines		
CARDAMYST NASAL SOLUTION 2 X 70 MG/DOSE	5	PA; QL (4 EA per 30 days)
CARTIA XT ORAL CAPSULE EXTENDED RELEASE 24 HOUR 120 MG, 180 MG, 240 MG, 300 MG	2	
<i>diltiazem hcl er beads oral capsule extended release 24 hour 120 mg, 180 mg, 240 mg, 300 mg, 360 mg, 420 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>diltiazem hcl er coated beads oral capsule extended release 24 hour 120 mg, 180 mg, 240 mg, 300 mg, 360 mg</i>	2	
<i>diltiazem hcl er oral capsule extended release 12 hour 120 mg, 60 mg, 90 mg</i>	2	
<i>diltiazem hcl er oral capsule extended release 24 hour 120 mg, 180 mg, 240 mg</i>	2	
<i>diltiazem hcl oral tablet 120 mg, 30 mg, 60 mg, 90 mg</i>	1	
<i>dilt-xr oral capsule extended release 24 hour 120 mg, 180 mg, 240 mg</i>	2	
<i>verapamil hcl er oral capsule extended release 24 hour 100 mg, 120 mg, 180 mg, 200 mg, 240 mg, 300 mg, 360 mg</i>	2	
<i>verapamil hcl er oral tablet extended release 120 mg, 180 mg, 240 mg</i>	1	
<i>verapamil hcl oral tablet 120 mg, 40 mg, 80 mg</i>	1	
Cardiovascular Agents, Other		
<i>acetazolamide oral tablet 125 mg, 250 mg</i>	2	
<i>aliskiren fumarate oral tablet 150 mg, 300 mg</i>	2	
<i>amiloride-hydrochlorothiazide oral tablet 5-50 mg</i>	1	
<i>amlodipine besy-benazepril hcl oral capsule 10-20 mg, 10-40 mg, 2.5-10 mg, 5-10 mg, 5-20 mg, 5-40 mg</i>	1	
<i>amlodipine besylate-valsartan oral tablet 10-160 mg, 10-320 mg, 5-160 mg, 5-320 mg</i>	1	
<i>amlodipine-atorvastatin oral tablet 10-10 mg, 10-20 mg, 10-40 mg, 10-80 mg, 2.5-10 mg, 2.5-20 mg, 2.5-40 mg, 5-10 mg, 5-20 mg, 5-40 mg, 5-80 mg</i>	2	
<i>amlodipine-olmesartan oral tablet 10-20 mg, 10-40 mg, 5-20 mg, 5-40 mg</i>	1	
<i>amlodipine-valsartan-hctz oral tablet 10-160-12.5 mg, 10-160-25 mg, 10-320-25 mg, 5-160-12.5 mg, 5-160-25 mg</i>	2	
<i>atenolol-chlorthalidone oral tablet 100-25 mg, 50-25 mg</i>	1	
<i>benazepril-hydrochlorothiazide oral tablet 10-12.5 mg, 20-12.5 mg, 20-25 mg, 5-6.25 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>bisoprolol-hydrochlorothiazide oral tablet 10-6.25 mg, 2.5-6.25 mg, 5-6.25 mg</i>	1	
CAMZYOS ORAL CAPSULE 10 MG, 15 MG, 2.5 MG, 5 MG	5	PA; QL (30 EA per 30 days)
<i>candesartan cilexetil-hctz oral tablet 16-12.5 mg, 32-12.5 mg, 32-25 mg</i>	2	
CORLANOR ORAL SOLUTION 5 MG/5ML	4	PA; QL (450 ML per 30 days)
<i>digoxin oral solution 0.05 mg/ml</i>	2	QL (150 ML per 30 days)
<i>digoxin oral tablet 125 mcg, 250 mcg</i>	1	QL (30 EA per 30 days)
<i>enalapril-hydrochlorothiazide oral tablet 10-25 mg, 5-12.5 mg</i>	1	
ENTRESTO ORAL CAPSULE SPRINKLE 15-16 MG, 6-6 MG	3	QL (240 EA per 30 days)
ENTRESTO ORAL TABLET 24-26 MG, 49-51 MG, 97-103 MG	3	QL (60 EA per 30 days)
<i>fosinopril sodium-hctz oral tablet 10-12.5 mg, 20-12.5 mg</i>	2	
<i>irbesartan-hydrochlorothiazide oral tablet 150-12.5 mg, 300-12.5 mg</i>	1	
<i>ivabradine hcl oral tablet 5 mg, 7.5 mg</i>	2	PA
KERENDIA ORAL TABLET 10 MG, 20 MG, 40 MG	4	PA; QL (30 EA per 30 days)
<i>lisinopril-hydrochlorothiazide oral tablet 10-12.5 mg, 20-12.5 mg, 20-25 mg</i>	1	
LODOCO ORAL TABLET 0.5 MG	4	PA
<i>losartan potassium-hctz oral tablet 100-12.5 mg, 100-25 mg, 50-12.5 mg</i>	1	
<i>metoprolol-hydrochlorothiazide oral tablet 100-25 mg, 100-50 mg, 50-25 mg</i>	2	
<i>metyrosine oral capsule 250 mg</i>	5	PA
MYQORZO ORAL TABLET 10 MG, 15 MG, 20 MG, 5 MG	5	PA; QL (30 EA per 30 days)
NEXLETOL ORAL TABLET 180 MG	3	PA; QL (30 EA per 30 days)
NEXLIZET ORAL TABLET 180-10 MG	3	PA; QL (30 EA per 30 days)
<i>olmesartan medoxomil-hctz oral tablet 20-12.5 mg, 40-12.5 mg, 40-25 mg</i>	1	
<i>olmesartan-amlodipine-hctz oral tablet 20-5-12.5 mg, 40-10-12.5 mg, 40-10-25 mg, 40-5-12.5 mg, 40-5-25 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>pentoxifylline er oral tablet extended release 400 mg</i>	1	
<i>quinapril-hydrochlorothiazide oral tablet 10-12.5 mg, 20-12.5 mg, 20-25 mg</i>	1	
<i>ranolazine er oral tablet extended release 12 hour 1000 mg, 500 mg</i>	2	
<i>spironolactone-hctz oral tablet 25-25 mg</i>	2	
<i>telmisartan-amlodipine oral tablet 40-10 mg, 40-5 mg, 80-10 mg, 80-5 mg</i>	2	
<i>telmisartan-hctz oral tablet 40-12.5 mg, 80-12.5 mg, 80-25 mg</i>	2	
<i>triamterene-hctz oral capsule 37.5-25 mg</i>	1	
<i>triamterene-hctz oral tablet 37.5-25 mg, 75-50 mg</i>	1	
<i>valsartan-hydrochlorothiazide oral tablet 160-12.5 mg, 160-25 mg, 320-12.5 mg, 320-25 mg, 80-12.5 mg</i>	1	
VERQUVO ORAL TABLET 10 MG, 2.5 MG, 5 MG	3	QL (30 EA per 30 days)
WEGOVY ORAL TABLET 1.5 MG, 25 MG, 4 MG, 9 MG	5	PA; QL (30 EA per 30 days)
WEGOVY SUBCUTANEOUS SOLUTION AUTO-INJECTOR 0.25 MG/0.5ML, 0.5 MG/0.5ML, 1 MG/0.5ML	5	PA; QL (2 ML per 28 days)
WEGOVY SUBCUTANEOUS SOLUTION AUTO-INJECTOR 1.7 MG/0.75ML, 2.4 MG/0.75ML	5	PA; QL (3 ML per 28 days)
Diuretics, Loop		
<i>bumetanide oral tablet 0.5 mg, 1 mg, 2 mg</i>	2	
<i>furosemide injection solution 10 mg/ml</i>	2	
<i>furosemide oral solution 10 mg/ml, 8 mg/ml</i>	1	
<i>furosemide oral tablet 20 mg, 40 mg, 80 mg</i>	1	
<i>torseamide oral tablet 10 mg, 100 mg, 20 mg, 5 mg</i>	2	
Diuretics, Potassium-Sparing		
<i>amiloride hcl oral tablet 5 mg</i>	1	
<i>eplerenone oral tablet 25 mg, 50 mg</i>	2	
<i>spironolactone oral tablet 100 mg, 25 mg, 50 mg</i>	1	
Diuretics, Thiazide		

Name of Drug	Drug Tier	Requirements/Limits
<i>chlorthalidone oral tablet 25 mg, 50 mg</i>	1	
<i>hydrochlorothiazide oral capsule 12.5 mg</i>	1	
<i>hydrochlorothiazide oral tablet 12.5 mg, 25 mg, 50 mg</i>	1	
<i>indapamide oral tablet 1.25 mg, 2.5 mg</i>	1	
<i>metolazone oral tablet 10 mg, 2.5 mg, 5 mg</i>	2	
Dyslipidemics, Fibric Acid Derivatives		
<i>fenofibrate micronized oral capsule 134 mg, 200 mg, 43 mg, 67 mg</i>	2	
<i>fenofibrate oral capsule 134 mg, 200 mg, 67 mg</i>	2	
<i>fenofibrate oral tablet 145 mg, 160 mg, 48 mg, 54 mg</i>	2	
<i>fenofibric acid oral capsule delayed release 135 mg, 45 mg</i>	2	
<i>gemfibrozil oral tablet 600 mg</i>	1	
Dyslipidemics, Hmg Coa Reductase Inhibitors		
<i>atorvastatin calcium oral tablet 10 mg, 20 mg, 40 mg, 80 mg</i>	1	
<i>lovastatin oral tablet 10 mg, 20 mg, 40 mg</i>	1	
<i>pravastatin sodium oral tablet 10 mg, 20 mg, 40 mg, 80 mg</i>	1	
<i>rosuvastatin calcium oral tablet 10 mg, 20 mg, 40 mg, 5 mg</i>	1	
<i>simvastatin oral tablet 10 mg, 20 mg, 40 mg, 5 mg, 80 mg</i>	1	
Dyslipidemics, Other		
<i>cholestyramine light oral packet 4 gm</i>	2	
<i>cholestyramine light oral powder 4 gm/dose</i>	2	
<i>cholestyramine oral packet 4 gm</i>	2	
<i>cholestyramine oral powder 4 gm/dose</i>	2	
<i>colesevelam hcl oral packet 3.75 gm</i>	2	
<i>colesevelam hcl oral tablet 625 mg</i>	2	
<i>colestipol hcl oral granules 5 gm</i>	2	
<i>colestipol hcl oral packet 5 gm</i>	2	
<i>colestipol hcl oral tablet 1 gm</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>ezetimibe oral tablet 10 mg</i>	1	
<i>ezetimibe-simvastatin oral tablet 10-10 mg, 10-20 mg, 10-40 mg, 10-80 mg</i>	2	
<i>icosapent ethyl oral capsule 0.5 gm, 1 gm</i>	2	
<i>niacin er (antihyperlipidemic) oral tablet extended release 1000 mg, 500 mg, 750 mg</i>	2	
<i>omega-3-acid ethyl esters oral capsule 1 gm</i>	2	
PREVALITE ORAL PACKET 4 GM	2	
PREVALITE ORAL POWDER 4 GM/DOSE	2	
REPATHA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 140 MG/ML	3	PA; QL (3 ML per 28 days)
REPATHA SURECLICK SUBCUTANEOUS SOLUTION AUTO-INJECTOR 140 MG/ML	3	PA; QL (3 ML per 28 days)
Vasodilators, Direct-Acting Arterial		
<i>hydralazine hcl oral tablet 10 mg, 100 mg, 25 mg, 50 mg</i>	2	
<i>isosorb dinitrate-hydralazine oral tablet 20-37.5 mg</i>	2	
<i>minoxidil oral tablet 10 mg, 2.5 mg</i>	2	
Vasodilators, Direct-Acting Arterial/ Venous		
<i>isosorbide dinitrate oral tablet 10 mg, 20 mg, 30 mg, 5 mg</i>	1	
<i>isosorbide mononitrate er oral tablet extended release 24 hour 120 mg, 30 mg, 60 mg</i>	1	
<i>isosorbide mononitrate oral tablet 10 mg, 20 mg</i>	1	
NITRO-BID TRANSDERMAL OINTMENT 2 %	4	
NITRO-DUR TRANSDERMAL PATCH 24 HOUR 0.3 MG/HR, 0.8 MG/HR	4	
<i>nitroglycerin rectal ointment 0.4 %</i>	2	
<i>nitroglycerin sublingual tablet sublingual 0.3 mg, 0.4 mg, 0.6 mg</i>	2	
<i>nitroglycerin transdermal patch 24 hour 0.1 mg/hr, 0.2 mg/hr, 0.4 mg/hr, 0.6 mg/hr</i>	2	
<i>nitroglycerin translingual solution 0.4 mg/spray</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
Central Nervous System Agents - Treatment Of Disorders Of The Brain And Spinal Column		
Attention Deficit Hyperactivity Disorder Agents, Amphetamines		
<i>amphetamine-dextroamphetamine oral capsule extended release 24 hour 10 mg, 15 mg, 20 mg, 25 mg, 30 mg, 5 mg</i>	2	QL (30 EA per 30 days)
<i>amphetamine-dextroamphetamine oral tablet 10 mg, 20 mg, 30 mg, 5 mg, 7.5 mg</i>	2	QL (60 EA per 30 days)
<i>amphetamine-dextroamphetamine oral tablet 12.5 mg</i>	2	QL (120 EA per 30 days)
<i>amphetamine-dextroamphetamine oral tablet 15 mg</i>	2	QL (90 EA per 30 days)
<i>dextroamphetamine sulfate er oral capsule extended release 24 hour 10 mg</i>	2	QL (150 EA per 30 days)
<i>dextroamphetamine sulfate er oral capsule extended release 24 hour 15 mg</i>	2	QL (120 EA per 30 days)
<i>dextroamphetamine sulfate er oral capsule extended release 24 hour 5 mg</i>	2	QL (90 EA per 30 days)
<i>dextroamphetamine sulfate oral tablet 10 mg, 5 mg</i>	2	QL (180 EA per 30 days)
Attention Deficit Hyperactivity Disorder Agents, Non-Amphetamines		
<i>atomoxetine hcl oral capsule 10 mg, 100 mg, 18 mg, 25 mg, 40 mg, 60 mg, 80 mg</i>	2	
<i>clonidine hcl er oral tablet extended release 12 hour 0.1 mg</i>	2	QL (120 EA per 30 days)
<i>dexmethylphenidate hcl er oral capsule extended release 24 hour 10 mg, 15 mg, 20 mg, 25 mg, 30 mg, 35 mg, 40 mg, 5 mg</i>	2	QL (30 EA per 30 days)
<i>dexmethylphenidate hcl oral tablet 10 mg</i>	2	QL (60 EA per 30 days)
<i>dexmethylphenidate hcl oral tablet 2.5 mg, 5 mg</i>	2	QL (90 EA per 30 days)
<i>guanfacine hcl er oral tablet extended release 24 hour 1 mg, 2 mg, 4 mg</i>	1	PA; QL (30 EA per 30 days)
<i>guanfacine hcl er oral tablet extended release 24 hour 3 mg</i>	1	PA; QL (60 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>methylphenidate hcl er (cd) oral capsule extended release 10 mg, 20 mg, 30 mg, 40 mg, 50 mg, 60 mg</i>	2	QL (30 EA per 30 days)
<i>methylphenidate hcl er (la) oral capsule extended release 24 hour 10 mg, 20 mg, 30 mg, 40 mg, 60 mg</i>	2	QL (30 EA per 30 days)
<i>methylphenidate hcl er (osm) oral tablet extended release 18 mg</i>	2	QL (120 EA per 30 days)
<i>methylphenidate hcl er (osm) oral tablet extended release 27 mg, 54 mg, 72 mg</i>	2	QL (30 EA per 30 days)
<i>methylphenidate hcl er (osm) oral tablet extended release 36 mg</i>	2	QL (60 EA per 30 days)
<i>methylphenidate hcl er (xr) oral capsule extended release 24 hour 10 mg, 15 mg, 20 mg, 30 mg, 40 mg, 50 mg, 60 mg</i>	2	QL (30 EA per 30 days)
<i>methylphenidate hcl er oral tablet extended release 10 mg</i>	2	QL (30 EA per 30 days)
<i>methylphenidate hcl er oral tablet extended release 20 mg</i>	2	QL (90 EA per 30 days)
<i>methylphenidate hcl er oral tablet extended release 24 hour 18 mg</i>	2	QL (120 EA per 30 days)
<i>methylphenidate hcl er(diffus) oral tablet extended release 27 mg, 54 mg</i>	2	QL (30 EA per 30 days)
<i>methylphenidate hcl er(diffus) oral tablet extended release 36 mg</i>	2	QL (60 EA per 30 days)
<i>methylphenidate hcl oral solution 10 mg/5ml</i>	2	QL (900 ML per 30 days)
<i>methylphenidate hcl oral solution 5 mg/5ml</i>	2	QL (1800 ML per 30 days)
<i>methylphenidate hcl oral tablet 10 mg, 20 mg, 5 mg</i>	2	QL (90 EA per 30 days)
<i>methylphenidate hcl oral tablet chewable 10 mg</i>	2	QL (180 EA per 30 days)
<i>methylphenidate hcl oral tablet chewable 2.5 mg, 5 mg</i>	2	QL (90 EA per 30 days)
Central Nervous System, Other		
AQNEURSA ORAL PACKET 1 GM	5	PA; QL (120 EA per 30 days)
AUSTEDO ORAL TABLET 12 MG, 6 MG, 9 MG	5	PA
AUSTEDO XR ORAL TABLET EXTENDED RELEASE 24 HOUR 12 MG, 18 MG, 24 MG, 30 MG, 36 MG, 42 MG, 48 MG, 6 MG	5	PA

Name of Drug	Drug Tier	Requirements/Limits
AUSTEDO XR PATIENT TITRATION ORAL TABLET EXTENDED RELEASE THERAPY PACK 12 & 18 & 24 & 30 MG	5	PA
BYSANTI ORAL TABLET 1 MG, 10 MG, 12 MG, 2 MG, 4 MG, 6 MG, 8 MG	5	PA; QL (60 EA per 30 days)
BYSANTI TITRATION PACK A ORAL TABLET THERAPY PACK 1 & 2 & 4 & 6 MG	4	PA; QL (8 EA per 180 days)
BYSANTI TITRATION PACK B ORAL TABLET THERAPY PACK 1 & 2 & 6 & 8 MG	4	PA; QL (12 EA per 180 days)
BYSANTI TITRATION PACK C ORAL TABLET THERAPY PACK 1 & 2 & 6 MG	4	PA; QL (8 EA per 180 days)
COBENFY ORAL CAPSULE 100-20 MG, 125-30 MG, 50-20 MG	5	PA; QL (56 EA per 28 days)
COBENFY STARTER PACK ORAL CAPSULE THERAPY PACK 50-20 & 100-20 MG	5	PA; QL (56 EA per 180 days)
EVRYSDI ORAL SOLUTION RECONSTITUTED 0.75 MG/ML	5	PA
EVRYSDI ORAL TABLET 5 MG	5	PA
FIRDAPSE ORAL TABLET 10 MG	5	PA
INGREZZA ORAL CAPSULE 40 MG, 60 MG, 80 MG	5	PA; QL (30 EA per 30 days)
INGREZZA ORAL CAPSULE SPRINKLE 40 MG, 60 MG, 80 MG	5	PA; QL (30 EA per 30 days)
INGREZZA ORAL CAPSULE THERAPY PACK 40 & 80 MG	5	PA; QL (56 EA per 365 days)
LEQEMBI IQLIK SUBCUTANEOUS SOLUTION AUTO-INJECTOR 360 MG/1.8ML	5	PA
NUEDEXTA ORAL CAPSULE 20-10 MG	5	PA
RADICAVA ORS ORAL SUSPENSION 105 MG/5ML	5	PA
RADICAVA ORS STARTER KIT ORAL SUSPENSION 105 MG/5ML	5	PA
<i>riluzole oral tablet 50 mg</i>	2	
<i>tetrabenazine oral tablet 12.5 mg</i>	4	PA
<i>tetrabenazine oral tablet 25 mg</i>	5	PA
VEOZAH ORAL TABLET 45 MG	4	PA
Fibromyalgia Agents		

Name of Drug	Drug Tier	Requirements/Limits
DRIZALMA SPRINKLE ORAL CAPSULE DELAYED RELEASE SPRINKLE 20 MG, 30 MG, 40 MG, 60 MG	4	ST
<i>duloxetine hcl oral capsule delayed release particles 20 mg, 30 mg, 60 mg</i>	2	QL (60 EA per 30 days)
<i>milnacipran hcl oral 12.5 & 25 & 50 mg</i>	2	
<i>milnacipran hcl oral tablet 100 mg, 12.5 mg, 25 mg, 50 mg</i>	2	
SAVELLA TITRATION PACK ORAL 12.5 & 25 & 50 MG	4	ST
Multiple Sclerosis Agents		
BAFIERTAM ORAL CAPSULE DELAYED RELEASE 95 MG	5	PA
BETASERON SUBCUTANEOUS KIT 0.3 MG	5	PA
<i>cladribine (10 tabs) oral tablet therapy pack 10 mg</i>	5	PA
<i>cladribine (4 tabs) oral tablet therapy pack 10 mg</i>	5	PA
<i>cladribine (5 tabs) oral tablet therapy pack 10 mg</i>	5	PA
<i>cladribine (6 tabs) oral tablet therapy pack 10 mg</i>	5	PA
<i>cladribine (7 tabs) oral tablet therapy pack 10 mg</i>	5	PA
<i>cladribine (8 tabs) oral tablet therapy pack 10 mg</i>	5	PA
<i>cladribine (9 tabs) oral tablet therapy pack 10 mg</i>	5	PA
<i>cladribine 10 mg tablet therapy pack oral oral tablet therapy pack 10 mg</i>	5	PA
<i>dalfampridine er oral tablet extended release 12 hour 10 mg</i>	2	PA
<i>dimethyl fumarate oral capsule delayed release 120 mg</i>	2	PA; QL (56 EA per 28 days)
<i>dimethyl fumarate oral capsule delayed release 240 mg</i>	2	PA; QL (60 EA per 30 days)
<i>dimethyl fumarate starter pack oral capsule delayed release therapy pack 120 & 240 mg</i>	2	PA
<i>fingolimod hcl oral capsule 0.5 mg</i>	4	PA; QL (30 EA per 30 days)
<i>glatiramer acetate subcutaneous solution prefilled syringe 20 mg/ml</i>	5	PA; QL (30 ML per 30 days)
<i>glatiramer acetate subcutaneous solution prefilled syringe 40 mg/ml</i>	5	PA; QL (12 ML per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
GLATOPA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 20 MG/ML	5	PA; QL (30 ML per 30 days)
GLATOPA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 40 MG/ML	5	PA; QL (12 ML per 28 days)
KESIMPTA SUBCUTANEOUS SOLUTION AUTO-INJECTOR 20 MG/0.4ML	5	PA
MAYZENT ORAL TABLET 0.25 MG, 1 MG, 2 MG	5	PA
MAYZENT STARTER PACK ORAL TABLET THERAPY PACK 12 X 0.25 MG	5	PA
MAYZENT STARTER PACK ORAL TABLET THERAPY PACK 7 X 0.25 MG	4	PA
PONVORY ORAL TABLET 20 MG	5	PA
PONVORY STARTER PACK ORAL TABLET THERAPY PACK 2-3-4-5-6-7-8-9 & 10 MG	5	PA
REBIF REBIDOSE SUBCUTANEOUS SOLUTION AUTO-INJECTOR 22 MCG/0.5ML, 44 MCG/0.5ML	5	PA
REBIF REBIDOSE TITRATION PACK SUBCUTANEOUS SOLUTION AUTO- INJECTOR 6X8.8 & 6X22 MCG	5	PA
REBIF SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 22 MCG/0.5ML, 44 MCG/0.5ML	5	PA
REBIF TITRATION PACK SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 6X8.8 & 6X22 MCG	5	PA
TASCENSO ODT ORAL TABLET DISPERSIBLE 0.25 MG, 0.5 MG	5	PA
<i>teriflunomide oral tablet 14 mg, 7 mg</i>	2	PA; QL (30 EA per 30 days)
ZEPOSIA 7-DAY STARTER PACK ORAL CAPSULE THERAPY PACK 4 X 0.23MG & 3 X 0.46MG	5	PA
ZEPOSIA ORAL CAPSULE 0.92 MG	5	PA
ZEPOSIA STARTER KIT ORAL CAPSULE THERAPY PACK 0.23MG & 0.46MG 0.92MG(21)	5	PA

Dental And Oral Agents - Treatment Of Mouth And Gum Disorders

Name of Drug	Drug Tier	Requirements/Limits
Dental And Oral Agents		
<i>cevimeline hcl oral capsule 30 mg</i>	2	
<i>chlorhexidine gluconate mouth/throat solution 0.12 %</i>	1	
<i>pilocarpine hcl oral tablet 5 mg, 7.5 mg</i>	2	
<i>triamcinolone acetonide mouth/throat paste 0.1 %</i>	2	
Dermatological Agents - Treatment Of Skin Conditions		
Acne And Rosacea Agents		
<i>acitretin oral capsule 10 mg, 17.5 mg, 25 mg</i>	2	PA
<i>adapalene external gel 0.3 %</i>	2	
<i>adapalene-benzoyl peroxide external gel 0.1-2.5 %</i>	2	
AMNESTEEM ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG	2	
<i>benzoyl peroxide-erythromycin external gel 5-3 %</i>	2	
CLARAVIS ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG	2	
<i>clindamycin phos-benzoyl perox external gel 1-5 %, 1.2-2.5 %, 1.2-5 %</i>	2	
<i>isotretinoin oral capsule 10 mg, 20 mg, 30 mg, 40 mg</i>	2	
<i>tazarotene external cream 0.05 %, 0.1 %</i>	2	
<i>tazarotene external gel 0.05 %, 0.1 %</i>	2	
<i>tretinoin external cream 0.025 %, 0.05 %, 0.1 %</i>	2	
<i>tretinoin external gel 0.01 %, 0.025 %</i>	2	
ZENATANE ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG	2	
Dermatitis And Pruritus Agents		
<i>alclometasone dipropionate external cream 0.05 %</i>	2	
<i>alclometasone dipropionate external ointment 0.05 %</i>	2	
<i>ammonium lactate external cream 12 %</i>	2	
<i>ammonium lactate external lotion 12 %</i>	2	
<i>betamethasone dipropionate aug external cream 0.05 %</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>betamethasone dipropionate aug external gel 0.05 %</i>	2	
<i>betamethasone dipropionate aug external lotion 0.05 %</i>	2	
<i>betamethasone dipropionate aug external ointment 0.05 %</i>	2	
<i>betamethasone dipropionate external cream 0.05 %</i>	2	
<i>betamethasone dipropionate external lotion 0.05 %</i>	2	
<i>betamethasone dipropionate external ointment 0.05 %</i>	2	
<i>betamethasone valerate external cream 0.1 %</i>	2	
<i>betamethasone valerate external lotion 0.1 %</i>	2	
<i>betamethasone valerate external ointment 0.1 %</i>	2	
<i>clobetasol prop emollient base external cream 0.05 %</i>	2	QL (60 GM per 30 days)
<i>clobetasol propionate e external cream 0.05 %</i>	2	
<i>clobetasol propionate external cream 0.05 %</i>	2	QL (60 GM per 30 days)
<i>clobetasol propionate external gel 0.05 %</i>	2	QL (60 GM per 30 days)
<i>clobetasol propionate external ointment 0.05 %</i>	2	QL (60 GM per 30 days)
<i>clobetasol propionate external solution 0.05 %</i>	2	QL (50 ML per 30 days)
<i>desonide external cream 0.05 %</i>	2	
<i>desonide external lotion 0.05 %</i>	2	
<i>desonide external ointment 0.05 %</i>	2	
<i>desoximetasone external cream 0.05 %, 0.25 %</i>	2	
<i>desoximetasone external gel 0.05 %</i>	2	
<i>desoximetasone external ointment 0.05 %, 0.25 %</i>	2	
<i>doxepin hcl external cream 5 %</i>	2	PA; QL (45 GM per 30 days)
EUCRISA EXTERNAL OINTMENT 2 %	4	PA
<i>fluocinolone acetonide external cream 0.01 %, 0.025 %</i>	2	
<i>fluocinolone acetonide external ointment 0.025 %</i>	2	
<i>fluocinolone acetonide external solution 0.01 %</i>	2	
<i>fluocinonide emulsified base external cream 0.05 %</i>	2	
<i>fluocinonide external cream 0.05 %</i>	2	QL (120 GM per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>fluocinonide external gel 0.05 %</i>	2	QL (120 GM per 30 days)
<i>fluocinonide external ointment 0.05 %</i>	2	QL (60 GM per 30 days)
<i>fluocinonide external solution 0.05 %</i>	2	QL (60 ML per 30 days)
<i>fluticasone propionate external cream 0.05 %</i>	2	
<i>fluticasone propionate external lotion 0.05 %</i>	2	
<i>fluticasone propionate external ointment 0.005 %</i>	2	
<i>halobetasol propionate external cream 0.05 %</i>	2	
<i>halobetasol propionate external ointment 0.05 %</i>	2	
<i>hydrocortisone (perianal) external cream 1 %, 2.5 %</i>	1	
<i>hydrocortisone butyrate external cream 0.1 %</i>	2	
<i>hydrocortisone butyrate external ointment 0.1 %</i>	2	
<i>hydrocortisone butyrate external solution 0.1 %</i>	2	
<i>hydrocortisone external cream 1 %, 2.5 %</i>	1	
<i>hydrocortisone external lotion 2.5 %</i>	1	
<i>hydrocortisone external ointment 1 %, 2.5 %</i>	1	
<i>hydrocortisone valerate external cream 0.2 %</i>	2	
<i>hydrocortisone valerate external ointment 0.2 %</i>	2	
HYFTOR EXTERNAL GEL 0.2 %	5	PA
<i>mometasone furoate external cream 0.1 %</i>	2	
<i>mometasone furoate external ointment 0.1 %</i>	2	
<i>mometasone furoate external solution 0.1 %</i>	2	
<i>pimecrolimus external cream 1 %</i>	2	ST
<i>selenium sulfide external lotion 2.5 %</i>	2	
<i>tacrolimus external ointment 0.03 %, 0.1 %</i>	2	ST
<i>triamcinolone acetonide external cream 0.025 %, 0.1 %, 0.5 %</i>	2	
<i>triamcinolone acetonide external lotion 0.025 %, 0.1 %</i>	2	
<i>triamcinolone acetonide external ointment 0.025 %, 0.05 %, 0.1 %, 0.5 %</i>	2	
<i>triamcinolone in absorbase external ointment 0.05 %</i>	2	
Dermatological Agents, Other		
<i>alcohol pad , 70 %</i>	1	
ALCOHOL PAD 70 %	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>alcohol sheet , 70 %</i>	1	
<i>calcipotriene external cream 0.005 %</i>	2	QL (120 GM per 30 days)
<i>calcipotriene external ointment 0.005 %</i>	2	QL (120 GM per 30 days)
<i>calcipotriene external solution 0.005 %</i>	2	QL (120 ML per 30 days)
<i>calcitriol external ointment 3 mcg/gm</i>	2	
<i>clotrimazole-betamethasone external cream 1-0.05 %</i>	2	QL (45 GM per 28 days)
<i>clotrimazole-betamethasone external lotion 1-0.05 %</i>	2	QL (60 ML per 28 days)
<i>fluorouracil external cream 5 %</i>	2	QL (40 GM per 30 days)
<i>fluorouracil external solution 2 %, 5 %</i>	2	
<i>imiquimod external cream 5 %</i>	2	
<i>methoxsalen rapid oral capsule 10 mg</i>	2	
<i>nystatin-triamcinolone external cream 100000-0.1 unit/gm-%</i>	2	
<i>nystatin-triamcinolone external ointment 100000-0.1 unit/gm-%</i>	2	
OTEZLA ORAL TABLET 20 MG, 30 MG	5	PA
OTEZLA ORAL TABLET THERAPY PACK 10 & 20 & 30 MG, 4 X 10 & 51 X20 MG	5	PA
OTEZLA XR ORAL TABLET EXTENDED RELEASE 24 HOUR 75 MG	5	PA
OTEZLA/OTEZLA XR INITIATION PK ORAL TABLET THERAPY PACK 10&20&30&(ER)75 MG	5	PA
<i>podofilox external solution 0.5 %</i>	2	
SANTYL EXTERNAL OINTMENT 250 UNIT/GM	3	QL (90 GM per 30 days)
<i>silver sulfadiazine external cream 1 %</i>	2	
<i>sodium chloride irrigation solution 0.9 %</i>	2	
Pediculicides/Scabicides		
<i>malathion external lotion 0.5 %</i>	2	
<i>permethrin external cream 5 %</i>	2	QL (60 GM per 30 days)
Topical Anti-Infectives		
<i>acyclovir external cream 5 %</i>	2	
<i>acyclovir external ointment 5 %</i>	2	
<i>ciclopirox external solution 8 %</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>ciclopirox olamine external cream 0.77 %</i>	2	
<i>ciclopirox olamine external suspension 0.77 %</i>	2	
<i>clindamycin phos (once-daily) external gel 1 %</i>	2	
<i>clindamycin phos (twice-daily) external gel 1 %</i>	2	
<i>clindamycin phosphate external lotion 1 %</i>	2	
<i>clindamycin phosphate external solution 1 %</i>	2	
<i>clindamycin phosphate external swab 1 %</i>	2	
<i>ery external pad 2 %</i>	2	
<i>erythromycin external gel 2 %</i>	2	
<i>erythromycin external solution 2 %</i>	2	
<i>gentamicin sulfate external cream 0.1 %</i>	1	
<i>gentamicin sulfate external ointment 0.1 %</i>	1	
<i>metronidazole external cream 0.75 %</i>	2	
<i>metronidazole external gel 0.75 %, 1 %</i>	2	
<i>metronidazole external lotion 0.75 %</i>	2	
<i>mupirocin external ointment 2 %</i>	2	QL (88 GM per 30 days)
<i>penciclovir external cream 1 %</i>	2	

**Electrolytes/Minerals/ Metals/ Vitamins
- Products That Supplement Or Replace
Electrolytes, Minerals, Metals Or
Vitamins**

Electrolyte/ Mineral Replacement

<i>carglumic acid oral tablet soluble 200 mg</i>	5	PA
ISOLYTE-S INTRAVENOUS SOLUTION	4	
ISOLYTE-S PH 7.4 INTRAVENOUS SOLUTION	4	
<i>kcl in dextrose-nacl intravenous solution 20-5-0.45 meq/l-%-%</i>	2	
KLOR-CON 10 ORAL TABLET EXTENDED RELEASE 10 MEQ	3	
KLOR-CON M10 ORAL TABLET EXTENDED RELEASE 10 MEQ	3	
KLOR-CON M15 ORAL TABLET EXTENDED RELEASE 15 MEQ	3	
KLOR-CON M20 ORAL TABLET EXTENDED RELEASE 20 MEQ	3	

Name of Drug	Drug Tier	Requirements/Limits
KLOR-CON ORAL TABLET EXTENDED RELEASE 8 MEQ	3	
<i>magnesium sulfate injection solution 50 %, 50 % (10ml syringe)</i>	2	
<i>potassium chloride crys er oral tablet extended release 10 meq, 15 meq, 20 meq</i>	1	
<i>potassium chloride er oral capsule extended release 10 meq, 8 meq</i>	1	
<i>potassium chloride er oral tablet extended release 10 meq, 15 meq, 20 meq, 8 meq</i>	1	
<i>potassium chloride intravenous solution 2 meq/ml, 2 meq/ml (20 ml), 40 meq/100ml</i>	2	
<i>potassium chloride oral solution 10 %, 20 meq/15ml (10%), 40 meq/15ml (20%)</i>	2	
<i>potassium citrate er oral tablet extended release 10 meq (1080 mg), 15 meq (1620 mg), 5 meq (540 mg)</i>	2	
<i>sodium chloride (pf) injection solution 0.9 %</i>	2	
<i>sodium chloride intravenous solution 0.45 %, 0.9 %, 3 %</i>	2	
<i>sodium fluoride oral tablet 2.2 (1 f) mg</i>	2	
Electrolyte/Mineral/Metal Modifiers		
CUVRIOR ORAL TABLET 300 MG	5	PA
<i>deferasirox granules oral packet 180 mg, 360 mg, 90 mg</i>	5	PA
<i>deferasirox oral packet 180 mg, 360 mg, 90 mg</i>	5	PA
<i>deferasirox oral tablet 180 mg, 360 mg, 90 mg</i>	2	PA
<i>deferasirox oral tablet soluble 125 mg</i>	2	PA
<i>deferasirox oral tablet soluble 250 mg, 500 mg</i>	5	PA
<i>deferiprone oral tablet 1000 mg, 500 mg</i>	5	PA
<i>penicillamine oral tablet 250 mg</i>	5	PA
<i>tolvaptan (hyponatremia) oral tablet 15 mg, 30 mg</i>	5	PA
<i>tolvaptan oral tablet 15 mg, 30 mg</i>	5	PA
<i>tolvaptan oral tablet therapy pack 15 mg, 30 & 15 mg, 45 & 15 mg, 60 & 30 mg, 90 & 30 mg</i>	5	PA
<i>trientine hcl oral capsule 250 mg</i>	5	PA
Electrolytes/Minerals/Metals/Vitamins		

Name of Drug	Drug Tier	Requirements/Limits
CLINISOL SF INTRAVENOUS SOLUTION 15 %	4	B/D
<i>dextrose intravenous solution 10 %, 5 %</i>	2	
<i>dextrose-sodium chloride intravenous solution 10-0.2 %, 10-0.45 %, 2.5-0.45 %, 5-0.2 %, 5-0.45 %, 5-0.9 %</i>	2	
INTRALIPID INTRAVENOUS EMULSION 20 %, 30 %	4	B/D
ISOLYTE-P IN D5W INTRAVENOUS SOLUTION	4	
<i>levocarnitine oral solution 1 gm/10ml</i>	2	
<i>levocarnitine oral tablet 330 mg</i>	2	
<i>levocarnitine sf oral solution 1 gm/10ml</i>	2	
NUTRILIPID INTRAVENOUS EMULSION 20 %	4	B/D
PLENAMINE INTRAVENOUS SOLUTION 15 %	4	B/D
<i>pnv 27-ca/fe/fa oral tablet 60-1 mg</i>	2	
<i>prenatal oral tablet 27-1 mg</i>	2	
Phosphate Binders		
<i>calcium acetate (phos binder) oral capsule 667 mg</i>	2	
<i>lanthanum carbonate oral tablet chewable 1000 mg, 500 mg, 750 mg</i>	2	
<i>sevelamer carbonate oral packet 0.8 gm, 2.4 gm</i>	2	
<i>sevelamer carbonate oral tablet 800 mg</i>	2	
Potassium Binders		
LOKELMA ORAL PACKET 10 GM, 5 GM	3	
<i>sodium polystyrene sulfonate combination suspension 15 gm/60ml</i>	2	
<i>sodium polystyrene sulfonate oral powder</i>	2	
SPS (SODIUM POLYSTYRENE SULF) COMBINATION SUSPENSION 15 GM/60ML	2	
SPS (SODIUM POLYSTYRENE SULF) RECTAL SUSPENSION 30 GM/120ML	2	
VELTASSA ORAL PACKET 16.8 GM, 25.2 GM	5	QL (30 EA per 30 days)
VELTASSA ORAL PACKET 8.4 GM	5	QL (90 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
Vitamins		
<i>trinatal rx 1 oral tablet 60-1 mg</i>	2	
Gastrointestinal Agents - Treatment Of Stomach And Intestinal Conditions		
Anti-Constipation Agents		
<i>constulose oral solution 10 gm/15ml</i>	2	
<i>enulose oral solution 10 gm/15ml</i>	2	
GAVILYTE-C ORAL SOLUTION RECONSTITUTED 240 GM	2	
GAVILYTE-G ORAL SOLUTION RECONSTITUTED 236 GM	2	
GAVILYTE-N WITH FLAVOR PACK ORAL SOLUTION RECONSTITUTED 420 GM	2	
<i>generlac oral solution 10 gm/15ml</i>	2	
<i>lactulose encephalopathy oral solution 10 gm/15ml</i>	2	
<i>lactulose oral solution 10 gm/15ml, 20 gm/30ml</i>	2	
LINZESS ORAL CAPSULE 145 MCG, 290 MCG, 72 MCG	3	QL (30 EA per 30 days)
<i>lubiprostone oral capsule 24 mcg, 8 mcg</i>	2	QL (60 EA per 30 days)
MOVANTIK ORAL TABLET 12.5 MG, 25 MG	3	QL (30 EA per 30 days)
<i>peg 3350-kcl-na bicarb-nacl oral solution reconstituted 420 gm</i>	2	
<i>peg-3350/electrolytes oral solution reconstituted 236 gm</i>	2	
RELISTOR ORAL TABLET 150 MG	4	PA
RELISTOR SUBCUTANEOUS SOLUTION 12 MG/0.6ML, 12 MG/0.6ML (0.6ML SYRINGE)	5	PA
RELISTOR SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 12 MG/0.6ML, 8 MG/0.4ML	5	PA
TRULANCE ORAL TABLET 3 MG	3	QL (30 EA per 30 days)
Anti-Diarrheal Agents		
<i>alosetron hcl oral tablet 0.5 mg</i>	2	QL (60 EA per 30 days)
<i>alosetron hcl oral tablet 1 mg</i>	4	QL (60 EA per 30 days)
<i>diphenoxylate-atropine oral liquid 2.5-0.025 mg/5ml</i>	2	PA

Name of Drug	Drug Tier	Requirements/Limits
<i>diphenoxylate-atropine oral tablet 2.5-0.025 mg</i>	2	PA
<i>loperamide hcl oral capsule 2 mg</i>	2	
XERMELO ORAL TABLET 250 MG	5	PA
XIFAXAN ORAL TABLET 200 MG	4	PA
XIFAXAN ORAL TABLET 550 MG	5	PA
Antispasmodics, Gastrointestinal		
<i>dicyclomine hcl oral capsule 10 mg</i>	1	
<i>dicyclomine hcl oral solution 10 mg/5ml</i>	2	
<i>dicyclomine hcl oral tablet 20 mg</i>	1	
<i>glycopyrrolate oral solution 1 mg/5ml</i>	2	
<i>glycopyrrolate oral tablet 1 mg, 2 mg</i>	2	
Gastrointestinal Agents, Other		
GATTEX SUBCUTANEOUS KIT 5 MG	5	PA
LIVMARLI ORAL SOLUTION 19 MG/ML, 9.5 MG/ML	5	PA
LIVMARLI ORAL TABLET 10 MG, 15 MG, 20 MG, 30 MG	5	PA
<i>ursodiol oral capsule 300 mg</i>	2	
<i>ursodiol oral tablet 250 mg, 500 mg</i>	2	
VOQUEZNA DUAL PAK ORAL THERAPY PACK 500-20 MG	4	QL (112 EA per 14 days)
VOQUEZNA ORAL TABLET 10 MG, 20 MG	4	QL (30 EA per 30 days)
VOQUEZNA TRIPLE PAK ORAL THERAPY PACK 500-500-20 MG	4	QL (112 EA per 14 days)
VOWST ORAL CAPSULE	5	PA
Histamine2 (H2) Receptor Antagonists		
<i>cimetidine oral tablet 200 mg, 300 mg, 400 mg, 800 mg</i>	2	
<i>famotidine oral tablet 20 mg, 40 mg</i>	1	
Protectants		
<i>misoprostol oral tablet 100 mcg, 200 mcg</i>	2	
<i>sucralfate oral tablet 1 gm</i>	1	
Proton Pump Inhibitors		
<i>esomeprazole magnesium oral capsule delayed release 20 mg</i>	1	QL (30 EA per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>esomeprazole magnesium oral capsule delayed release 40 mg</i>	1	QL (60 EA per 30 days)
<i>lansoprazole oral capsule delayed release 15 mg, 30 mg</i>	1	
<i>omeprazole oral capsule delayed release 10 mg, 20 mg</i>	1	QL (30 EA per 30 days)
<i>omeprazole oral capsule delayed release 40 mg</i>	1	QL (60 EA per 30 days)
<i>pantoprazole sodium oral tablet delayed release 20 mg</i>	1	QL (30 EA per 30 days)
<i>pantoprazole sodium oral tablet delayed release 40 mg</i>	1	QL (60 EA per 30 days)
Genetic Or Enzyme Or Protein Disorder: Replacement, Modifiers, Treatment - Products That Replace, Modify, Or Treat Genetic Or Enzyme Disorders		
Genetic Or Enzyme Or Protein Disorder: Replacement, Modifiers, Treatment		
ARALAST NP INTRAVENOUS SOLUTION RECONSTITUTED 1000 MG, 500 MG	5	PA
<i>betaine oral powder</i>	5	
CERDELGA ORAL CAPSULE 84 MG	5	PA
CHOLBAM ORAL CAPSULE 250 MG, 50 MG	5	PA
CREON ORAL CAPSULE DELAYED RELEASE PARTICLES 12000-38000 UNIT, 24000-76000 UNIT, 3000-9500 UNIT, 36000-114000 UNIT, 6000-19000 UNIT	3	
CYSTAGON ORAL CAPSULE 150 MG, 50 MG	4	PA
GALAFOLD ORAL CAPSULE 123 MG	5	PA
GLASSIA INTRAVENOUS SOLUTION 1000 MG/50ML	5	PA
GLASSIA INTRAVENOUS SOLUTION 4 GM/200ML, 5 GM/250ML	4	PA
<i>glycerol phenylbutyrate oral liquid 1.1 gm/ml</i>	5	PA
<i>l-glutamine oral packet 5 gm</i>	5	PA
<i>miglustat oral capsule 100 mg</i>	5	PA
<i>nitisinone oral capsule 10 mg, 2 mg, 20 mg, 5 mg</i>	5	PA

Name of Drug	Drug Tier	Requirements/Limits
ORFADIN ORAL SUSPENSION 4 MG/ML	5	PA
PROLASTIN-C INTRAVENOUS SOLUTION 1000 MG/20ML	5	PA
REVCovi INTRAMUSCULAR SOLUTION 2.4 MG/1.5ML	5	PA
<i>sapropterin dihydrochloride oral packet 100 mg, 500 mg</i>	5	PA
<i>sapropterin dihydrochloride oral tablet 100 mg</i>	5	PA
<i>sodium phenylbutyrate oral powder 3 gm/tsp</i>	5	PA
<i>sodium phenylbutyrate oral tablet 500 mg</i>	5	PA
SUCRAID ORAL SOLUTION 8500 UNIT/ML	5	PA
ZEMAIRA INTRAVENOUS SOLUTION RECONSTITUTED 1000 MG, 4000 MG, 5000 MG	5	PA
ZENPEP ORAL CAPSULE DELAYED RELEASE PARTICLES 10000-32000 UNIT, 15000-47000 UNIT, 20000-63000 UNIT, 25000-79000 UNIT, 3000-10000 UNIT, 40000-126000 UNIT, 5000-24000 UNIT, 60000-189600 UNIT	3	

Genitourinary Agents - Treatment Of Urinary Tract And Prostate Conditions

Antispasmodics, Urinary

<i>darifenacin hydrobromide er oral tablet extended release 24 hour 15 mg, 7.5 mg</i>	2	ST
<i>fesoterodine fumarate er oral tablet extended release 24 hour 4 mg, 8 mg</i>	2	QL (30 EA per 30 days)
<i>flavoxate hcl oral tablet 100 mg</i>	2	
GEMTESA ORAL TABLET 75 MG	3	QL (30 EA per 30 days)
MYRBETRIQ ORAL SUSPENSION RECONSTITUTED ER 8 MG/ML	3	QL (300 ML per 30 days)
MYRBETRIQ ORAL TABLET EXTENDED RELEASE 24 HOUR 25 MG, 50 MG	3	QL (30 EA per 30 days)
<i>oxybutynin chloride er oral tablet extended release 24 hour 10 mg, 15 mg</i>	1	QL (60 EA per 30 days)
<i>oxybutynin chloride er oral tablet extended release 24 hour 5 mg</i>	1	QL (30 EA per 30 days)
<i>oxybutynin chloride oral solution 5 mg/5ml</i>	1	
<i>oxybutynin chloride oral tablet 5 mg</i>	1	

Name of Drug	Drug Tier	Requirements/Limits
<i>solifenacin succinate oral tablet 10 mg, 5 mg</i>	2	QL (30 EA per 30 days)
<i>tolterodine tartrate er oral capsule extended release 24 hour 2 mg, 4 mg</i>	2	
<i>tolterodine tartrate oral tablet 1 mg, 2 mg</i>	2	
<i>tropium chloride er oral capsule extended release 24 hour 60 mg</i>	2	ST
<i>tropium chloride oral tablet 20 mg</i>	2	
Benign Prostatic Hypertrophy Agents		
<i>alfuzosin hcl er oral tablet extended release 24 hour 10 mg</i>	1	
<i>dutasteride oral capsule 0.5 mg</i>	2	
<i>finasteride oral tablet 5 mg</i>	1	
<i>tadalafil oral tablet 5 mg</i>	2	PA
<i>tamsulosin hcl oral capsule 0.4 mg</i>	1	
Genitourinary Agents, Other		
<i>bethanechol chloride oral tablet 10 mg, 25 mg, 5 mg, 50 mg</i>	2	
ELMIRON ORAL CAPSULE 100 MG	4	
FILSPARI ORAL TABLET 200 MG, 400 MG	5	PA
<i>tiopronin oral tablet 100 mg</i>	5	PA
<i>tiopronin oral tablet delayed release 100 mg, 300 mg</i>	5	PA
Hormonal Agents, Stimulant/ Replacement/ Modifying (Adrenal) - Treatment Of Conditions Requiring Steroids		
Hormonal Agents, Stimulant/ Replacement/ Modifying (Adrenal)		
CORTROPHIN GEL SUBCUTANEOUS PREFILLED SYRINGE 40 UNIT/0.5ML, 80 UNIT/ML	5	PA
CORTROPHIN INJECTION GEL 80 UNIT/ML	5	PA
<i>dexamethasone oral solution 0.5 mg/5ml</i>	2	
<i>dexamethasone oral tablet 0.5 mg, 0.75 mg, 1 mg, 1.5 mg, 2 mg, 4 mg, 6 mg</i>	1	
<i>fludrocortisone acetate oral tablet 0.1 mg</i>	2	
<i>hydrocortisone oral tablet 10 mg, 20 mg, 5 mg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
KYMBEE ORAL TABLET 18 MG, 30 MG, 36 MG, 6 MG	5	PA
<i>methylprednisolone oral tablet 16 mg, 32 mg, 4 mg, 8 mg</i>	2	
<i>methylprednisolone oral tablet therapy pack 4 mg</i>	2	
<i>prednisolone oral solution 15 mg/5ml</i>	2	
<i>prednisolone sodium phosphate oral solution 25 mg/5ml, 5 mg/5ml</i>	2	
Hormonal Agents, Stimulant/ Replacement/ Modifying (Pituitary) - Treatment Of Pituitary Gland Conditions		
Hormonal Agents, Stimulant/ Replacement/ Modifying (Pituitary)		
<i>desmopressin ace spray refrig nasal solution 0.01 %</i>	2	
<i>desmopressin acetate oral tablet 0.1 mg, 0.2 mg</i>	2	
<i>desmopressin acetate spray nasal solution 0.01 %</i>	2	
EGRIFTA SV SUBCUTANEOUS SOLUTION RECONSTITUTED 2 MG	5	PA
EGRIFTA WR SUBCUTANEOUS KIT 11.6 MG	5	PA
GENOTROPIN MINIQUICK SUBCUTANEOUS PREFILLED SYRINGE 0.2 MG	4	PA
GENOTROPIN MINIQUICK SUBCUTANEOUS PREFILLED SYRINGE 0.4 MG, 0.6 MG, 0.8 MG, 1 MG, 1.2 MG, 1.4 MG, 1.6 MG, 1.8 MG, 2 MG	5	PA
GENOTROPIN SUBCUTANEOUS CARTRIDGE 12 MG	5	PA
GENOTROPIN SUBCUTANEOUS CARTRIDGE 5 MG	4	PA
INCRELEX SUBCUTANEOUS SOLUTION 40 MG/4ML	5	PA
NGENLA SUBCUTANEOUS SOLUTION PEN-INJECTOR 24 MG/1.2ML, 60 MG/1.2ML	5	PA
OMNITROPE SUBCUTANEOUS SOLUTION CARTRIDGE 10 MG/1.5ML, 5 MG/1.5ML	5	PA

Name of Drug	Drug Tier	Requirements/Limits
OMNITROPE SUBCUTANEOUS SOLUTION RECONSTITUTED 5.8 MG	5	PA
SEROSTIM SUBCUTANEOUS SOLUTION RECONSTITUTED 4 MG, 5 MG, 6 MG	5	PA
SKYTROFA SUBCUTANEOUS CARTRIDGE 0.7 MG, 1.4 MG, 1.8 MG, 11 MG, 13.3 MG, 2.1 MG, 2.5 MG, 3 MG, 3.6 MG, 4.3 MG, 5.2 MG, 6.3 MG, 7.6 MG, 9.1 MG	5	PA
Hormonal Agents, Stimulant/ Replacement/ Modifying (Sex Hormones/ Modifiers) - For The Replacement Or Modification Of Sex Hormones		
Androgens		
<i>danazol oral capsule 100 mg, 200 mg, 50 mg</i>	2	
<i>methyltestosterone oral capsule 10 mg</i>	5	PA
<i>testosterone cypionate intramuscular solution 100 mg/ml, 200 mg/ml, 200 mg/ml (1 ml)</i>	2	
<i>testosterone enanthate intramuscular solution 200 mg/ml</i>	2	
<i>testosterone transdermal gel 1.62 %, 12.5 mg/act (1%), 20.25 mg/1.25gm (1.62%), 20.25 mg/act (1.62%), 25 mg/2.5gm (1%), 40.5 mg/2.5gm (1.62%), 50 mg/5gm (1%)</i>	2	PA
<i>testosterone transdermal solution 30 mg/act</i>	2	PA
Estrogens		
<i>estradiol oral tablet 0.5 mg, 1 mg, 2 mg</i>	1	
<i>estradiol transdermal patch twice weekly 0.025 mg/24hr, 0.0375 mg/24hr, 0.05 mg/24hr, 0.075 mg/24hr, 0.1 mg/24hr</i>	2	QL (8 EA per 28 days)
<i>estradiol transdermal patch weekly 0.025 mg/24hr, 0.0375 mg/24hr, 0.05 mg/24hr, 0.06 mg/24hr, 0.075 mg/24hr, 0.1 mg/24hr</i>	2	QL (4 EA per 28 days)
<i>estradiol vaginal cream 0.01 %</i>	2	
<i>estradiol vaginal tablet 10 mcg</i>	2	
<i>estradiol valerate intramuscular oil 10 mg/ml, 20 mg/ml, 40 mg/ml</i>	2	
PREMARIN ORAL TABLET 0.3 MG, 0.45 MG, 0.625 MG, 0.9 MG, 1.25 MG	3	

Name of Drug	Drug Tier	Requirements/Limits
PREMARIN VAGINAL CREAM 0.625 MG/GM	3	
Hormonal Agents, Stimulant/ Replacement/ Modifying (Sex Hormones/ Modifiers)		
ABIGALE LO ORAL TABLET 0.5-0.1 MG	2	
ABIGALE ORAL TABLET 1-0.5 MG	2	
ALTAVERA ORAL TABLET 0.15-30 MG-MCG	2	
<i>alyacen 1/35 oral tablet 1-35 mg-mcg</i>	2	
APRI ORAL TABLET 0.15-30 MG-MCG	2	
ARANELLE ORAL TABLET 0.5/1/0.5-35 MG-MCG	2	
ASHLYNA ORAL TABLET 0.15-0.03 & 0.01 MG	2	
AUBRA EQ ORAL TABLET 0.1-20 MG-MCG	2	
AUROVELA 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
AUROVELA 1/20 ORAL TABLET 1-20 MG-MCG	2	
AUROVELA 24 FE ORAL TABLET 1-20 MG-MCG(24)	2	
AUROVELA FE 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
AUROVELA FE 1/20 ORAL TABLET 1-20 MG-MCG	2	
AVIANE ORAL TABLET 0.1-20 MG-MCG	2	
AYUNA ORAL TABLET 0.15-30 MG-MCG	2	
AZURETTE ORAL TABLET 0.15-0.02/0.01 MG (21/5)	2	
BALZIVA ORAL TABLET 0.4-35 MG-MCG	2	
BLISOVI 24 FE ORAL TABLET 1-20 MG-MCG(24)	2	
BLISOVI FE 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
BLISOVI FE 1/20 ORAL TABLET 1-20 MG-MCG	2	
<i>briellyn oral tablet 0.4-35 mg-mcg</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
CHARLOTTE 24 FE ORAL TABLET CHEWABLE 1-20 MG-MCG(24)	2	
CHATEAL EQ ORAL TABLET 0.15-30 MG-MCG	2	
COMBIPATCH TRANSDERMAL PATCH TWICE WEEKLY 0.05-0.14 MG/DAY, 0.05-0.25 MG/DAY	4	
CRYSELLE ORAL TABLET 0.3-30 MG-MCG	2	
CRYSELLE-28 ORAL TABLET 0.3-30 MG-MCG	2	
CYRED EQ ORAL TABLET 0.15-30 MG-MCG	2	
DASETTA 1/35 (28) ORAL TABLET 1-35 MG-MCG	2	
DASETTA 7/7/7 ORAL TABLET 0.5/0.75/1-35 MG-MCG	2	
DAYSEE ORAL TABLET 0.15-0.03 & 0.01 MG	2	
DEPO-PROVERA INTRAMUSCULAR SUSPENSION 150 MG/ML	3	
DEPO-PROVERA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 150 MG/ML	3	
<i>drospirenone-ethinyl estradiol oral tablet 3-0.02 mg, 3-0.03 mg</i>	2	
ELINEST ORAL TABLET 0.3-30 MG-MCG	2	
ELURYNG VAGINAL RING 0.12-0.015 MG/24HR	2	
EMZAHH ORAL TABLET 0.35 MG	2	
ENILLORING VAGINAL RING 0.12-0.015 MG/24HR	2	
ENSKYCE ORAL TABLET 0.15-30 MG-MCG	2	
ESTARYLLA ORAL TABLET 0.25-35 MG-MCG	2	
<i>estradiol-norethindrone acet oral tablet 0.5-0.1 mg, 1-0.5 mg</i>	2	
<i>etonogestrel-ethinyl estradiol vaginal ring 0.12-0.015 mg/24hr</i>	2	
FALMINA ORAL TABLET 0.1-20 MG-MCG	2	
FINZALA ORAL TABLET CHEWABLE 1-20 MG-MCG(24)	2	

Name of Drug	Drug Tier	Requirements/Limits
FYAVOLV ORAL TABLET 0.5-2.5 MG-MCG, 1-5 MG-MCG	2	
HAILEY 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
HAILEY 24 FE ORAL TABLET 1-20 MG-MCG(24)	2	
HAILEY FE 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
HAILEY FE 1/20 ORAL TABLET 1-20 MG-MCG	2	
HALOETTE VAGINAL RING 0.12-0.015 MG/24HR	2	
HEATHER ORAL TABLET 0.35 MG	2	
ICLEVIA ORAL TABLET 0.15-0.03 MG	2	
INTROVALE ORAL TABLET 0.15-0.03 MG	2	
ISIBLOOM ORAL TABLET 0.15-30 MG-MCG	2	
JAIMIESS ORAL TABLET 0.15-0.03 & 0.01 MG	2	
JASMIEL ORAL TABLET 3-0.02 MG	2	
JENCYCLA ORAL TABLET 0.35 MG	2	
JINTELI ORAL TABLET 1-5 MG-MCG	2	
JOLESSA ORAL TABLET 0.15-0.03 MG	2	
JULEBER ORAL TABLET 0.15-30 MG-MCG	2	
JUNEL 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
JUNEL 1/20 ORAL TABLET 1-20 MG-MCG	2	
JUNEL FE 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
JUNEL FE 1/20 ORAL TABLET 1-20 MG-MCG	2	
JUNEL FE 24 ORAL TABLET 1-20 MG-MCG(24)	2	
KALLIGA ORAL TABLET 0.15-30 MG-MCG	2	
KARIVA ORAL TABLET 0.15-0.02/0.01 MG (21/5)	2	
KELNOR 1/35 ORAL TABLET 1-35 MG-MCG	2	
KURVELO ORAL TABLET 0.15-30 MG-MCG	2	
KYLEENA INTRAUTERINE INTRAUTERINE DEVICE 19.5 MG	4	

Name of Drug	Drug Tier	Requirements/Limits
LARIN 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
LARIN 1/20 ORAL TABLET 1-20 MG-MCG	2	
LARIN 24 FE ORAL TABLET 1-20 MG-MCG(24)	2	
LARIN FE 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
LARIN FE 1/20 ORAL TABLET 1-20 MG-MCG	2	
LESSINA ORAL TABLET 0.1-20 MG-MCG	2	
LEVONEST ORAL TABLET 50-30/75-40/ 125-30 MCG	2	
<i>levonorgest-eth estrad 91-day oral tablet 0.1-0.02 & 0.01 mg, 0.15-0.03 & 0.01 mg, 0.15-0.03 mg</i>	2	
<i>levonorgestrel-ethinyl estrad oral tablet 0.1-20 mg-mcg</i>	2	
<i>levonorg-eth estrad triphasic oral tablet 50-30/75-40/ 125-30 mcg</i>	2	
LILETTA (52 MG) INTRAUTERINE INTRAUTERINE DEVICE 20.1 MCG/DAY	4	
LOJAIMIESS ORAL TABLET 0.1-0.02 & 0.01 MG	2	
LORYNA ORAL TABLET 3-0.02 MG	2	
LOW-OGESTREL ORAL TABLET 0.3-30 MG-MCG	2	
LO-ZUMANDIMINE ORAL TABLET 3-0.02 MG	2	
LUIZZA 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
LUIZZA 1/20 ORAL TABLET 1-20 MG-MCG	2	
LUTERA ORAL TABLET 0.1-20 MG-MCG	2	
LYLEQ ORAL TABLET 0.35 MG	2	
<i>marlissa oral tablet 0.15-30 mg-mcg</i>	2	
MIBELAS 24 FE ORAL TABLET CHEWABLE 1-20 MG-MCG(24)	2	
MICROGESTIN 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
MICROGESTIN 1/20 ORAL TABLET 1-20 MG-MCG	2	

Name of Drug	Drug Tier	Requirements/Limits
MICROGESTIN FE 1.5/30 ORAL TABLET 1.5-30 MG-MCG	2	
MICROGESTIN FE 1/20 ORAL TABLET 1-20 MG-MCG	2	
MILI ORAL TABLET 0.25-35 MG-MCG	2	
MIMVEY ORAL TABLET 1-0.5 MG	2	
MIRENA (52 MG) INTRAUTERINE INTRAUTERINE DEVICE 21 MCG/DAY	4	
MONO-LINYAH ORAL TABLET 0.25-35 MG-MCG	2	
NECON 0.5/35 (28) ORAL TABLET 0.5-35 MG-MCG	2	
NEXPLANON SUBCUTANEOUS IMPLANT 68 MG	3	
NIKKI ORAL TABLET 3-0.02 MG	2	
<i>norelgestromin-eth estradiol transdermal patch weekly 150-35 mcg/24hr</i>	2	
<i>norethin ace-eth estrad-fe oral tablet chewable 1-20 mg-mcg(24)</i>	2	
<i>norethindrone acet-ethinyl est oral tablet 1-20 mg-mcg</i>	2	
<i>norethindrone-eth estradiol oral tablet 0.5-2.5 mg-mcg, 1-5 mg-mcg</i>	2	
<i>norgestimate-eth estradiol oral tablet 0.25-35 mg-mcg</i>	2	
<i>norgestim-eth estrad triphasic oral tablet 0.18/0.215/0.25 mg-25 mcg, 0.18/0.215/0.25 mg-35 mcg</i>	2	
NORTREL 0.5/35 (28) ORAL TABLET 0.5-35 MG-MCG	2	
NORTREL 1/35 (21) ORAL TABLET 1-35 MG-MCG	2	
NORTREL 1/35 (28) ORAL TABLET 1-35 MG-MCG	2	
NORTREL 7/7/7 ORAL TABLET 0.5/0.75/1-35 MG-MCG	2	
NYLIA 1/35 ORAL TABLET 1-35 MG-MCG	2	
NYLIA 7/7/7 ORAL TABLET 0.5/0.75/1-35 MG-MCG	2	

Name of Drug	Drug Tier	Requirements/Limits
PHILITH ORAL TABLET 0.4-35 MG-MCG	2	
PIMTREA ORAL TABLET 0.15-0.02/0.01 MG (21/5)	2	
PORTIA-28 ORAL TABLET 0.15-30 MG-MCG	2	
PREMPHASE ORAL TABLET 0.625-5 MG	3	
PREMPRO ORAL TABLET 0.3-1.5 MG, 0.45-1.5 MG, 0.625-2.5 MG, 0.625-5 MG	3	
RECLIPSEN ORAL TABLET 0.15-30 MG-MCG	2	
SETLAKIN ORAL TABLET 0.15-0.03 MG	2	
SIMLIYA ORAL TABLET 0.15-0.02/0.01 MG (21/5)	2	
SIMPESSE ORAL TABLET 0.15-0.03 & 0.01 MG	2	
SKYLA INTRAUTERINE INTRAUTERINE DEVICE 13.5 MG	3	
SPRINTEC 28 ORAL TABLET 0.25-35 MG-MCG	2	
SYEDA ORAL TABLET 3-0.03 MG	2	
TARINA 24 FE ORAL TABLET 1-20 MG-MCG(24)	2	
TARINA FE 1/20 EQ ORAL TABLET 1-20 MG-MCG	2	
TILIA FE ORAL TABLET 1-20/1-30/1-35 MG-MCG	2	
TRI-ESTARYLLA ORAL TABLET 0.18/0.215/0.25 MG-35 MCG	2	
TRI-LEGEST FE ORAL TABLET 1-20/1-30/1-35 MG-MCG	2	
TRI-LINYAH ORAL TABLET 0.18/0.215/0.25 MG-35 MCG	2	
TRI-LO-ESTARYLLA ORAL TABLET 0.18/0.215/0.25 MG-25 MCG	2	
TRI-LO-MARZIA ORAL TABLET 0.18/0.215/0.25 MG-25 MCG	2	
TRI-LO-MILI ORAL TABLET 0.18/0.215/0.25 MG-25 MCG	2	
TRI-LO-SPRINTEC ORAL TABLET 0.18/0.215/0.25 MG-25 MCG	2	

Name of Drug	Drug Tier	Requirements/Limits
TRI-MILI ORAL TABLET 0.18/0.215/0.25 MG-35 MCG	2	
TRI-SPRINTEC ORAL TABLET 0.18/0.215/0.25 MG-35 MCG	2	
TRI-VYLIBRA LO ORAL TABLET 0.18/0.215/0.25 MG-25 MCG	2	
TRI-VYLIBRA ORAL TABLET 0.18/0.215/0.25 MG-35 MCG	2	
TURQOZ ORAL TABLET 0.3-30 MG-MCG	2	
VELIVET ORAL TABLET 0.1/0.125/0.15 -0.025 MG	2	
VESTURA ORAL TABLET 3-0.02 MG	2	
VIENVA ORAL TABLET 0.1-20 MG-MCG	2	
<i>viorele oral tablet 0.15-0.02/0.01 mg (21/5)</i>	2	
VOLNEA ORAL TABLET 0.15-0.02/0.01 MG (21/5)	2	
VYFEMLA ORAL TABLET 0.4-35 MG-MCG	2	
VYLIBRA ORAL TABLET 0.25-35 MG-MCG	2	
WERA ORAL TABLET 0.5-35 MG-MCG	2	
WYMZYA FE ORAL TABLET CHEWABLE 0.4-35 MG-MCG	2	
XULANE TRANSDERMAL PATCH WEEKLY 150-35 MCG/24HR	2	
ZAFEMY TRANSDERMAL PATCH WEEKLY 150-35 MCG/24HR	2	
ZOVIA 1/35 (28) ORAL TABLET 1-35 MG-MCG	2	
ZUMANDIMINE ORAL TABLET 3-0.03 MG	2	
Progestins		
CAMILA ORAL TABLET 0.35 MG	2	
DEBLITANE ORAL TABLET 0.35 MG	2	
DEPO-SUBQ PROVERA 104 SUBCUTANEOUS SUSPENSION PREFILLED SYRINGE 104 MG/0.65ML	3	
ERRIN ORAL TABLET 0.35 MG	2	
INCASSIA ORAL TABLET 0.35 MG	2	
LYZA ORAL TABLET 0.35 MG	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>medroxyprogesterone acetate intramuscular suspension 150 mg/ml</i>	2	
<i>medroxyprogesterone acetate intramuscular suspension prefilled syringe 150 mg/ml</i>	2	
<i>medroxyprogesterone acetate oral tablet 10 mg, 2.5 mg, 5 mg</i>	1	
<i>megestrol acetate oral suspension 40 mg/ml, 400 mg/10ml, 625 mg/5ml</i>	2	PA
<i>megestrol acetate oral tablet 20 mg, 40 mg</i>	2	PA
MELEYA ORAL TABLET 0.35 MG	2	
NORA-BE ORAL TABLET 0.35 MG	2	
<i>norethindrone acetate oral tablet 5 mg</i>	2	
<i>norethindrone oral tablet 0.35 mg</i>	2	
NORLYROC ORAL TABLET 0.35 MG	2	
ORQUIDEA ORAL TABLET 0.35 MG	2	
<i>progesterone oral capsule 100 mg, 200 mg</i>	2	
SHAROBEL ORAL TABLET 0.35 MG	2	
Selective Estrogen Receptor Modifying Agents		
DUAVEE ORAL TABLET 0.45-20 MG	3	
<i>raloxifene hcl oral tablet 60 mg</i>	2	
Hormonal Agents, Stimulant/ Replacement/ Modifying (Thyroid) - Treatment Of Thyroid Conditions		
Hormonal Agents, Stimulant/ Replacement/ Modifying (Thyroid)		
LEVO-T ORAL TABLET 100 MCG, 112 MCG, 125 MCG, 137 MCG, 150 MCG, 175 MCG, 200 MCG, 25 MCG, 300 MCG, 50 MCG, 75 MCG, 88 MCG	4	
<i>levothyroxine sodium oral tablet 100 mcg, 112 mcg, 125 mcg, 137 mcg, 150 mcg, 175 mcg, 200 mcg, 25 mcg, 300 mcg, 50 mcg, 75 mcg, 88 mcg</i>	1	
LEVOXYL ORAL TABLET 100 MCG, 112 MCG, 125 MCG, 137 MCG, 150 MCG, 175 MCG, 200 MCG, 25 MCG, 50 MCG, 75 MCG, 88 MCG	4	

Name of Drug	Drug Tier	Requirements/Limits
<i>lithyronine sodium oral tablet 25 mcg, 5 mcg, 50 mcg</i>	2	
REZDIFFRA ORAL TABLET 100 MG, 60 MG, 80 MG	5	PA; QL (30 EA per 30 days)
SYNTHROID ORAL TABLET 100 MCG, 112 MCG, 125 MCG, 137 MCG, 150 MCG, 175 MCG, 200 MCG, 25 MCG, 300 MCG, 50 MCG, 75 MCG, 88 MCG	4	
UNITHROID ORAL TABLET 100 MCG, 112 MCG, 125 MCG, 137 MCG, 150 MCG, 175 MCG, 200 MCG, 25 MCG, 300 MCG, 50 MCG, 75 MCG, 88 MCG	4	
Hormonal Agents, Suppressant (Pituitary) - Treatment Of Or Modification Of Pituitary Hormone Secretion		
Hormonal Agents, Suppressant (Pituitary)		
<i>cabergoline oral tablet 0.5 mg</i>	2	
ELIGARD SUBCUTANEOUS KIT 22.5 MG, 30 MG, 45 MG, 7.5 MG	4	PA
FIRMAGON (240 MG DOSE) SUBCUTANEOUS SOLUTION RECONSTITUTED 120 MG/VIAL	5	PA
FIRMAGON SUBCUTANEOUS SOLUTION RECONSTITUTED 80 MG	4	PA
<i>leuprolide acetate (3 month) intramuscular injectable 22.5 mg</i>	2	PA
<i>leuprolide acetate injection kit 1 mg/0.2ml</i>	4	PA
LUPRON DEPOT (1-MONTH) INTRAMUSCULAR KIT 3.75 MG, 7.5 MG	5	PA
LUPRON DEPOT (3-MONTH) INTRAMUSCULAR KIT 11.25 MG, 22.5 MG	5	PA
LUPRON DEPOT (4-MONTH) INTRAMUSCULAR KIT 30 MG	5	PA
LUPRON DEPOT (6-MONTH) INTRAMUSCULAR KIT 45 MG	5	PA
LUTRATE DEPOT INTRAMUSCULAR INJECTABLE 22.5 MG	4	PA

Name of Drug	Drug Tier	Requirements/Limits
MYFEMBREE ORAL TABLET 40-1-0.5 MG	5	PA
<i>octreotide acetate injection solution 100 mcg/ml, 200 mcg/ml, 50 mcg/ml</i>	2	PA
<i>octreotide acetate injection solution 1000 mcg/ml, 500 mcg/ml</i>	5	PA
<i>octreotide acetate intramuscular kit 10 mg, 20 mg, 30 mg</i>	5	PA
<i>octreotide acetate subcutaneous solution prefilled syringe 100 mcg/ml, 50 mcg/ml</i>	2	
<i>octreotide acetate subcutaneous solution prefilled syringe 500 mcg/ml</i>	5	
ORGOVYX ORAL TABLET 120 MG	5	PA
ORIAHNN ORAL CAPSULE THERAPY PACK 300-1-0.5 & 300 MG	5	PA
ORILISSA ORAL TABLET 150 MG, 200 MG	5	PA
RECORLEV ORAL TABLET 150 MG	5	PA
SIGNIFOR SUBCUTANEOUS SOLUTION 0.3 MG/ML, 0.6 MG/ML, 0.9 MG/ML	5	PA
SOMAVERT SUBCUTANEOUS SOLUTION RECONSTITUTED 10 MG, 15 MG, 20 MG, 25 MG, 30 MG	5	PA
SYNAREL NASAL SOLUTION 2 MG/ML	5	PA
TARPEYO ORAL CAPSULE DELAYED RELEASE 4 MG	5	PA
TRELSTAR MIXJECT INTRAMUSCULAR SUSPENSION RECONSTITUTED 11.25 MG, 22.5 MG, 3.75 MG	4	PA
Hormonal Agents, Suppressant (Thyroid) - Treatment For Overactive Thyroid		
Antithyroid Agents		
<i>methimazole oral tablet 10 mg, 5 mg</i>	1	
<i>propylthiouracil oral tablet 50 mg</i>	2	
Immunological Agents - Medications That Alter The Immune System Including Vaccinations		
Angioedema Agents		

Name of Drug	Drug Tier	Requirements/Limits
CINRYZE INTRAVENOUS SOLUTION RECONSTITUTED 500 UNIT	5	PA
HAEGARDA SUBCUTANEOUS SOLUTION RECONSTITUTED 2000 UNIT, 3000 UNIT	5	PA
<i>icatibant acetate subcutaneous solution prefilled syringe 30 mg/3ml</i>	5	PA
ORLADEYO ORAL CAPSULE 110 MG, 150 MG	5	PA
ORLADEYO ORAL PACKET 108 MG, 132 MG, 72 MG, 96 MG	5	PA
Immunoglobulins		
GAMMAGARD ERC INJECTION SOLUTION 10 GM/100ML, 5 GM/50ML	5	B/D
GAMMAGARD INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 2.5 GM/25ML, 20 GM/200ML, 30 GM/300ML, 5 GM/50ML	5	B/D
GAMMAGARD S/D LESS IGA INTRAVENOUS SOLUTION RECONSTITUTED 10 GM, 5 GM	5	B/D
GAMMAKED INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 20 GM/200ML, 5 GM/50ML	5	B/D
GAMMAPLEX INTRAVENOUS SOLUTION 10 GM/100ML, 10 GM/200ML, 20 GM/200ML, 20 GM/400ML, 5 GM/100ML, 5 GM/50ML	5	B/D
GAMUNEX-C INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 2.5 GM/25ML, 20 GM/200ML, 40 GM/400ML, 5 GM/50ML	5	B/D
OCTAGAM INTRAVENOUS SOLUTION 1 GM/20ML, 10 GM/100ML, 10 GM/200ML, 2 GM/20ML, 2.5 GM/50ML, 20 GM/200ML, 30 GM/300ML, 5 GM/100ML, 5 GM/50ML	5	B/D
PRIVIGEN INTRAVENOUS SOLUTION 10 GM/100ML, 20 GM/200ML, 40 GM/400ML, 5 GM/50ML	5	B/D
Immunological Agents, Other		
ACTEMRA ACTPEN SUBCUTANEOUS SOLUTION AUTO-INJECTOR 162 MG/0.9ML	5	PA; QL (3.6 ML per 28 days)
ACTEMRA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 162 MG/0.9ML	5	PA; QL (3.6 ML per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
ARCALYST SUBCUTANEOUS SOLUTION RECONSTITUTED 220 MG	5	PA
AVTOZMA SUBCUTANEOUS SOLUTION AUTO-INJECTOR 162 MG/0.9ML	5	PA; QL (3.6 ML per 30 days)
AVTOZMA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 162 MG/0.9ML	5	PA; QL (3.6 ML per 30 days)
BENLYSTA SUBCUTANEOUS SOLUTION AUTO-INJECTOR 200 MG/ML	5	PA
BENLYSTA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 200 MG/ML	5	PA
CIBINQO ORAL TABLET 100 MG, 200 MG, 50 MG	5	PA; QL (30 EA per 30 days)
COSENTYX (300 MG DOSE) SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 150 MG/ML	5	PA; QL (10 ML per 28 days)
COSENTYX INTRAVENOUS SOLUTION 125 MG/5ML	5	PA
COSENTYX SENSOREADY (300 MG) SUBCUTANEOUS SOLUTION AUTO-INJECTOR 150 MG/ML	5	PA; QL (10 ML per 28 days)
COSENTYX SENSOREADY PEN SUBCUTANEOUS SOLUTION AUTO-INJECTOR 150 MG/ML	5	PA; QL (10 ML per 28 days)
COSENTYX SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 150 MG/ML	5	PA; QL (10 ML per 28 days)
COSENTYX SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 75 MG/0.5ML	5	PA; QL (2.5 ML per 28 days)
COSENTYX UNOREADY SUBCUTANEOUS SOLUTION AUTO-INJECTOR 300 MG/2ML	5	PA; QL (10 ML per 28 days)
ENTYVIO PEN SUBCUTANEOUS SOLUTION AUTO-INJECTOR 108 MG/0.68ML	5	PA
FABHALTA ORAL CAPSULE 200 MG	5	PA
ILARIS SUBCUTANEOUS SOLUTION 150 MG/ML	5	PA
ILUMYA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 100 MG/ML	5	PA
IMULDOSA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
IMULDOSA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 90 MG/ML	3	PA; QL (1 ML per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
KEVZARA SUBCUTANEOUS SOLUTION AUTO-INJECTOR 150 MG/1.14ML, 200 MG/1.14ML	5	PA; QL (2.28 ML per 28 days)
KEVZARA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 150 MG/1.14ML, 200 MG/1.14ML	5	PA; QL (2.28 ML per 28 days)
KINERET SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 100 MG/0.67ML	5	PA
LEQSELVI ORAL TABLET 8 MG	5	PA; QL (60 EA per 30 days)
LITFULO ORAL CAPSULE 50 MG	5	PA
ORENCIA CLICKJECT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 125 MG/ML	5	PA; QL (4 ML per 28 days)
ORENCIA INTRAVENOUS SOLUTION RECONSTITUTED 250 MG	5	PA; QL (4 EA per 28 days)
ORENCIA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 125 MG/ML	5	PA; QL (4 ML per 28 days)
ORENCIA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 50 MG/0.4ML	5	PA; QL (1.6 ML per 28 days)
ORENCIA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 87.5 MG/0.7ML	5	PA; QL (2.8 ML per 28 days)
SELARSDI INTRAVENOUS SOLUTION 130 MG/26ML	3	PA; QL (104 ML per 180 days)
SELARSDI SUBCUTANEOUS SOLUTION 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
SELARSDI SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
SELARSDI SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 90 MG/ML	3	PA; QL (1 ML per 28 days)
SILIQ SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 210 MG/1.5ML	5	PA; QL (4.5 ML per 28 days)
SOTYKTU ORAL TABLET 6 MG	5	PA
STARJEMZA INTRAVENOUS SOLUTION 130 MG/26ML	3	PA; QL (104 ML per 180 days)
STARJEMZA SUBCUTANEOUS SOLUTION 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
STARJEMZA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
STARJEMZA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 90 MG/ML	3	PA; QL (1 ML per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
STELARA INTRAVENOUS SOLUTION 130 MG/26ML	3	PA; QL (104 ML per 180 days)
STELARA SUBCUTANEOUS SOLUTION 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
STELARA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
STELARA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 90 MG/ML	3	PA; QL (1 ML per 28 days)
STEQEYMA INTRAVENOUS SOLUTION 130 MG/26ML	3	PA; QL (104 ML per 180 days)
STEQEYMA SUBCUTANEOUS SOLUTION 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
STEQEYMA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
STEQEYMA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 90 MG/ML	3	PA; QL (1 ML per 28 days)
TALTZ SUBCUTANEOUS SOLUTION AUTO-INJECTOR 80 MG/ML	5	PA; QL (3 ML per 28 days)
TALTZ SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 20 MG/0.25ML	5	PA; QL (0.75 ML per 28 days)
TALTZ SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 40 MG/0.5ML	5	PA; QL (1.5 ML per 28 days)
TALTZ SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 80 MG/ML	5	PA; QL (3 ML per 28 days)
<i>tofacitinib citrate er oral tablet extended release 24 hour 11 mg</i>	2	PA; QL (30 EA per 30 days)
<i>tofacitinib citrate er oral tablet extended release 24 hour 22 mg</i>	5	PA; QL (30 EA per 30 days)
<i>tofacitinib citrate oral solution 1 mg/ml</i>	5	PA; QL (480 ML per 24 days)
<i>tofacitinib citrate oral tablet 10 mg, 5 mg</i>	2	PA; QL (60 EA per 30 days)
TREMFYA ONE-PRESS SUBCUTANEOUS SOLUTION PEN-INJECTOR 100 MG/ML	5	PA; QL (1 ML per 28 days)
TREMFYA PEN SUBCUTANEOUS SOLUTION AUTO-INJECTOR 100 MG/ML	5	PA; QL (1 ML per 28 days)
TREMFYA PEN SUBCUTANEOUS SOLUTION AUTO-INJECTOR 200 MG/2ML	5	PA; QL (4 ML per 28 days)
TREMFYA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 100 MG/ML	5	PA; QL (1 ML per 28 days)

Name of Drug	Drug Tier	Requirements/Limits
TREMFYA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 200 MG/2ML	5	PA; QL (4 ML per 28 days)
TREMFYA-CD/UC INDUCTION SUBCUTANEOUS SOLUTION AUTO- INJECTOR 200 MG/2ML	5	PA
<i>ustekinumab subcutaneous solution 45 mg/0.5ml</i>	3	PA; QL (0.5 ML per 28 days)
<i>ustekinumab subcutaneous solution prefilled syringe 45 mg/0.5ml</i>	3	PA; QL (0.5 ML per 28 days)
<i>ustekinumab subcutaneous solution prefilled syringe 90 mg/ml</i>	3	PA; QL (1 ML per 28 days)
<i>ustekinumab-aauz subcutaneous solution prefilled syringe 45 mg/0.5ml</i>	3	PA; QL (0.5 ML per 28 days)
<i>ustekinumab-aauz subcutaneous solution prefilled syringe 90 mg/ml</i>	3	PA; QL (1 ML per 28 days)
<i>ustekinumab-aekn subcutaneous solution prefilled syringe 45 mg/0.5ml</i>	3	PA; QL (0.5 ML per 28 days)
<i>ustekinumab-aekn subcutaneous solution prefilled syringe 90 mg/ml</i>	3	PA; QL (1 ML per 28 days)
XELJANZ ORAL SOLUTION 1 MG/ML	5	PA; QL (480 ML per 24 days)
XELJANZ ORAL TABLET 10 MG, 5 MG	5	PA; QL (60 EA per 30 days)
XELJANZ XR ORAL TABLET EXTENDED RELEASE 24 HOUR 11 MG, 22 MG	5	PA; QL (30 EA per 30 days)
YESINTEK INTRAVENOUS SOLUTION 130 MG/26ML	3	PA; QL (104 ML per 180 days)
YESINTEK SUBCUTANEOUS SOLUTION 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
YESINTEK SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML	3	PA; QL (0.5 ML per 28 days)
YESINTEK SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 90 MG/ML	3	PA; QL (1 ML per 28 days)
ZILBRYSQ SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 16.6 MG/0.416ML, 23 MG/0.574ML, 32.4 MG/0.81ML	5	PA
Immunostimulants		
ACTIMMUNE SUBCUTANEOUS SOLUTION 100 MCG/0.5ML	5	
PEGASYS SUBCUTANEOUS SOLUTION 180 MCG/ML	5	PA

Name of Drug	Drug Tier	Requirements/Limits
PEGASYS SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 180 MCG/0.5ML	5	PA
Immunosuppressants		
<i>adalimumab-fkjp (2 pen) subcutaneous auto-injector kit 40 mg/0.8ml</i>	3	PA; QL (6 EA per 28 days)
<i>adalimumab-fkjp (2 syringe) subcutaneous prefilled syringe kit 20 mg/0.4ml</i>	3	PA; QL (4 EA per 28 days)
<i>adalimumab-fkjp (2 syringe) subcutaneous prefilled syringe kit 40 mg/0.8ml</i>	3	PA; QL (6 EA per 28 days)
ASTAGRAF XL ORAL CAPSULE EXTENDED RELEASE 24 HOUR 0.5 MG, 1 MG	4	B/D
ASTAGRAF XL ORAL CAPSULE EXTENDED RELEASE 24 HOUR 5 MG	5	B/D
<i>azathioprine oral tablet 50 mg</i>	2	B/D
CIMZIA (1 SYRINGE) SUBCUTANEOUS PREFILLED SYRINGE KIT 200 MG/ML	5	PA; QL (2 EA per 28 days)
CIMZIA (2 SYRINGE) SUBCUTANEOUS PREFILLED SYRINGE KIT 200 MG/ML	5	PA; QL (2 EA per 28 days)
CIMZIA SUBCUTANEOUS KIT 2 X 200 MG	5	PA; QL (2 EA per 28 days)
CIMZIA-STARTER SUBCUTANEOUS PREFILLED SYRINGE KIT 200 MG/ML	5	PA; QL (3 EA per 28 days)
<i>cyclosporine modified oral capsule 100 mg, 25 mg, 50 mg</i>	2	B/D
<i>cyclosporine modified oral solution 100 mg/ml</i>	2	B/D
<i>cyclosporine oral capsule 100 mg, 25 mg</i>	2	B/D
ENBREL MINI SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	5	PA; QL (8 ML per 28 days)
ENBREL SUBCUTANEOUS SOLUTION 25 MG/0.5ML	5	PA; QL (8 ML per 28 days)
ENBREL SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 25 MG/0.5ML, 50 MG/ML	5	PA; QL (8 ML per 28 days)
ENBREL SURECLICK SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML	5	PA; QL (8 ML per 28 days)
ENVARUS XR ORAL TABLET EXTENDED RELEASE 24 HOUR 0.75 MG, 1 MG	4	B/D
ENVARUS XR ORAL TABLET EXTENDED RELEASE 24 HOUR 4 MG	5	B/D
<i>everolimus oral tablet 0.25 mg, 0.5 mg</i>	2	B/D

Name of Drug	Drug Tier	Requirements/Limits
<i>everolimus oral tablet 0.75 mg, 1 mg</i>	5	B/D
GENGRAF ORAL CAPSULE 100 MG, 25 MG	2	B/D
<i>leflunomide oral tablet 10 mg, 20 mg</i>	2	
LUPKYNIS ORAL CAPSULE 7.9 MG	5	PA
<i>methotrexate sodium (pf) injection solution 50 mg/2ml</i>	2	
<i>methotrexate sodium injection solution 250 mg/10ml, 50 mg/2ml</i>	2	
<i>methotrexate sodium oral tablet 2.5 mg</i>	1	
<i>mycophenolate mofetil oral capsule 250 mg</i>	2	B/D
<i>mycophenolate mofetil oral suspension reconstituted 200 mg/ml</i>	2	B/D
<i>mycophenolate mofetil oral tablet 500 mg</i>	2	B/D
<i>mycophenolate sodium oral tablet delayed release 180 mg, 360 mg</i>	2	B/D
PROGRAF INTRAVENOUS SOLUTION 5 MG/ML	5	B/D
PROGRAF ORAL PACKET 0.2 MG, 1 MG	4	B/D
REZUROCK ORAL TABLET 200 MG	5	PA
SIMLANDI (1 PEN) SUBCUTANEOUS AUTO-INJECTOR KIT 40 MG/0.4ML	3	PA; QL (6 EA per 28 days)
SIMLANDI (1 PEN) SUBCUTANEOUS AUTO-INJECTOR KIT 80 MG/0.8ML	3	PA; QL (3 EA per 28 days)
SIMLANDI (1 SYRINGE) SUBCUTANEOUS PREFILLED SYRINGE KIT 80 MG/0.8ML	3	PA; QL (3 EA per 28 days)
SIMLANDI (2 PEN) SUBCUTANEOUS AUTO-INJECTOR KIT 40 MG/0.4ML	3	PA; QL (6 EA per 28 days)
SIMLANDI (2 SYRINGE) SUBCUTANEOUS PREFILLED SYRINGE KIT 20 MG/0.2ML	3	PA; QL (4 EA per 28 days)
SIMLANDI (2 SYRINGE) SUBCUTANEOUS PREFILLED SYRINGE KIT 40 MG/0.4ML	3	PA; QL (6 EA per 28 days)
SIMPONI SUBCUTANEOUS SOLUTION AUTO-INJECTOR 100 MG/ML, 50 MG/0.5ML	5	PA
SIMPONI SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 100 MG/ML, 50 MG/0.5ML	5	PA
<i>sirolimus oral solution 1 mg/ml</i>	4	B/D
<i>sirolimus oral tablet 0.5 mg, 1 mg, 2 mg</i>	2	B/D

Name of Drug	Drug Tier	Requirements/Limits
<i>tacrolimus er oral capsule extended release 24 hour 0.5 mg, 1 mg</i>	2	B/D
<i>tacrolimus er oral capsule extended release 24 hour 5 mg</i>	4	B/D
<i>tacrolimus intravenous solution 5 mg/ml</i>	5	B/D
<i>tacrolimus oral capsule 0.5 mg, 1 mg, 5 mg</i>	2	B/D
Vaccines		
ABRYSVO INTRAMUSCULAR SOLUTION RECONSTITUTED 120 MCG/0.5ML	6	
ACTHIB INTRAMUSCULAR SOLUTION RECONSTITUTED	6	
ADACEL INTRAMUSCULAR SUSPENSION 5-2-15.5 (PREFILLED SYRINGE), 5-2-15.5 LF-MCG/0.5	6	
ADACEL INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 5-2-15.5 LF-MCG/0.5	6	
AREXVY INTRAMUSCULAR SUSPENSION RECONSTITUTED 120 MCG/0.5ML	6	
<i>bcg vaccine injection solution reconstituted 50 mg</i>	6	
BEXSERO INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 0.5 ML	6	
BEYFORTUS INTRAMUSCULAR SOLUTION PREFILLED SYRINGE 100 MG/ML, 50 MG/0.5ML	6	
BOOSTRIX INTRAMUSCULAR SUSPENSION 5-2.5-18.5 LF-MCG/0.5	6	
BOOSTRIX INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 5-2.5-18.5 LF-MCG/0.5	6	
DAPTACEL INTRAMUSCULAR SUSPENSION 23-15-5	6	
ENFLONSIA INTRAMUSCULAR SOLUTION PREFILLED SYRINGE 105 MG/0.7ML	6	
ENGERIX-B INJECTION SUSPENSION 20 MCG/ML	6	B/D
ENGERIX-B INJECTION SUSPENSION PREFILLED SYRINGE 10 MCG/0.5ML, 20 MCG/ML	6	B/D
ERVEBO INTRAMUSCULAR SUSPENSION	6	

Name of Drug	Drug Tier	Requirements/Limits
GARDASIL 9 INTRAMUSCULAR SUSPENSION 0.5 ML	6	
GARDASIL 9 INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 0.5 ML	6	
HAVRIX INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 1440 EL U/ML, 720 EL U/0.5ML	6	
HEPLISAV-B INTRAMUSCULAR SOLUTION PREFILLED SYRINGE 20 MCG/0.5ML	6	B/D
HIBERIX INJECTION SOLUTION RECONSTITUTED 10 MCG	6	
IMOVAX RABIES INTRAMUSCULAR SUSPENSION RECONSTITUTED 2.5 UNIT/ML	6	B/D
INFANRIX INTRAMUSCULAR SUSPENSION 25-58-10	6	
IPOL INJECTION SUSPENSION	6	
IXIARO INTRAMUSCULAR SUSPENSION	6	
JYNNEOS SUBCUTANEOUS SUSPENSION 0.5 ML	6	
KINRIX INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 0.5 ML	6	
MENQUADFI INTRAMUSCULAR SOLUTION 0.5 ML	6	
MENVEO INTRAMUSCULAR SOLUTION	6	
MENVEO INTRAMUSCULAR SOLUTION RECONSTITUTED	6	
M-M-R II INJECTION SOLUTION RECONSTITUTED	6	
MRESVIA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 50 MCG/0.5ML	6	
PEDIARIX INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE	6	
PEDVAX HIB INTRAMUSCULAR SUSPENSION 7.5 MCG/0.5ML	6	
PENBRAYA INTRAMUSCULAR SUSPENSION RECONSTITUTED	6	
<i>penmenvay intramuscular suspension reconstituted</i>	6	

Name of Drug	Drug Tier	Requirements/Limits
PENTACEL INTRAMUSCULAR SUSPENSION RECONSTITUTED	6	
PRIORIX SUBCUTANEOUS SUSPENSION RECONSTITUTED	6	
PROQUAD SUBCUTANEOUS SUSPENSION RECONSTITUTED	6	
QUADRACEL INTRAMUSCULAR SUSPENSION	6	
QUADRACEL INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 0.5 ML	6	
RABAERT INTRAMUSCULAR SUSPENSION RECONSTITUTED	6	B/D
RECOMBIVAX HB INJECTION SUSPENSION 10 MCG/ML, 40 MCG/ML, 5 MCG/0.5ML	6	B/D
RECOMBIVAX HB INJECTION SUSPENSION PREFILLED SYRINGE 10 MCG/ML, 5 MCG/0.5ML	6	B/D
ROTARIX ORAL SUSPENSION	6	
ROTATEQ ORAL SOLUTION	6	
SHINGRIX INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 50 MCG/0.5ML	6	QL (2 ML per 999 days)
SHINGRIX INTRAMUSCULAR SUSPENSION RECONSTITUTED 50 MCG/0.5ML	6	QL (2 EA per 999 days)
TENIVAC INTRAMUSCULAR INJECTABLE 5-2 LFU (INJECTION)	6	B/D
TENIVAC INTRAMUSCULAR SUSPENSION 5-2 LF/0.5ML	6	B/D
TICOVAC INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 1.2 MCG/0.25ML, 2.4 MCG/0.5ML	6	
TRUMENBA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 0.5 ML	6	
TWINRIX INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 720-20 ELU-MCG/ML	6	
TYPHIM VI INTRAMUSCULAR SOLUTION 25 MCG/0.5ML	6	
TYPHIM VI INTRAMUSCULAR SOLUTION PREFILLED SYRINGE 25 MCG/0.5ML	6	

Name of Drug	Drug Tier	Requirements/Limits
VAQTA INTRAMUSCULAR SUSPENSION 25 UNIT/0.5ML, 25 UNIT/0.5ML 0.5 ML, 50 UNIT/ML, 50 UNIT/ML 1 ML	6	
VAQTA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 25 UNIT/0.5ML, 50 UNIT/ML	6	
VARIVAX INJECTION SUSPENSION RECONSTITUTED 1350 PFU/0.5ML	6	
VAXCHORA ORAL SUSPENSION RECONSTITUTED	6	
VIMKUNYA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 40 MCG/0.8ML	6	
VIVOTIF ORAL CAPSULE DELAYED RELEASE	6	
YF-VAX SUBCUTANEOUS SUSPENSION RECONSTITUTED	6	
Inflammatory Bowel Disease Agents - Treatment Of Ulcerative Colitis Or Crohn's Disease		
Aminosalicylates		
<i>balsalazide disodium oral capsule 750 mg</i>	2	
<i>mesalamine er oral capsule extended release 24 hour 0.375 gm</i>	2	
<i>mesalamine oral capsule delayed release 400 mg</i>	2	
<i>mesalamine oral tablet delayed release 1.2 gm</i>	2	
<i>mesalamine rectal enema 4 gm</i>	2	
<i>mesalamine rectal suppository 1000 mg</i>	2	
<i>sulfasalazine oral tablet 500 mg</i>	2	
<i>sulfasalazine oral tablet delayed release 500 mg</i>	2	
Glucocorticoids		
<i>budesonide er oral tablet extended release 24 hour 9 mg</i>	4	
<i>budesonide oral capsule delayed release particles 3 mg</i>	2	
DEXAMETHASONE INTENSOL ORAL CONCENTRATE 1 MG/ML	2	
<i>dexamethasone oral elixir 0.5 mg/5ml</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>dexamethasone sodium phosphate injection solution 120 mg/30ml, 20 mg/5ml, 4 mg/ml</i>	2	
<i>hydrocortisone rectal enema 100 mg/60ml</i>	2	
<i>methylprednisolone acetate injection suspension 40 mg/ml, 80 mg/ml</i>	2	
<i>prednisolone sodium phosphate oral solution 15 mg/5ml</i>	2	
PREDNISONE INTENSOL ORAL CONCENTRATE 5 MG/ML	2	
<i>prednisone oral solution 5 mg/5ml</i>	2	
<i>prednisone oral tablet 1 mg, 10 mg, 2.5 mg, 20 mg, 5 mg, 50 mg</i>	1	
<i>prednisone oral tablet therapy pack 10 mg (21)</i>	1	
<i>prednisone oral tablet therapy pack 10 mg (48), 5 mg (21), 5 mg (48)</i>	2	

Metabolic Bone Disease Agents - Treatment Of Bone Diseases Including Osteoporosis

Metabolic Bone Disease Agents

<i>alendronate sodium oral tablet 10 mg</i>	1	QL (30 EA per 30 days)
<i>alendronate sodium oral tablet 35 mg, 70 mg</i>	1	QL (4 EA per 28 days)
BONSITY SUBCUTANEOUS SOLUTION PEN-INJECTOR 560 MCG/2.24ML	5	PA
<i>calcitonin (salmon) nasal solution 200 unit/act</i>	2	
<i>calcitriol oral capsule 0.25 mcg, 0.5 mcg</i>	2	
<i>calcitriol oral solution 1 mcg/ml</i>	2	
<i>cinacalcet hcl oral tablet 30 mg, 60 mg</i>	2	QL (60 EA per 30 days)
<i>cinacalcet hcl oral tablet 90 mg</i>	2	QL (120 EA per 30 days)
<i>doxercalciferol oral capsule 0.5 mcg, 1 mcg, 2.5 mcg</i>	2	
ENOBY SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 60 MG/ML	3	QL (1 ML per 180 days)
<i>ibandronate sodium oral tablet 150 mg</i>	1	
JUBBONTI SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 60 MG/ML	3	
OSENVELT SUBCUTANEOUS SOLUTION 120 MG/1.7ML	5	

Name of Drug	Drug Tier	Requirements/Limits
<i>paricalcitol oral capsule 1 mcg, 2 mcg, 4 mcg</i>	2	
<i>risedronate sodium oral tablet 150 mg, 35 mg, 35 mg (12 pack), 35 mg (4 pack), 5 mg</i>	2	
<i>risedronate sodium oral tablet 30 mg</i>	4	
STOBOCLO SUBCUTANEOUS SOLUTION REFILLED SYRINGE 60 MG/ML	3	
<i>teriparatide subcutaneous solution pen-injector 560 mcg/2.24ml</i>	5	PA
TYMLOS SUBCUTANEOUS SOLUTION PEN- INJECTOR 3120 MCG/1.56ML	5	PA
WYOST SUBCUTANEOUS SOLUTION 120 MG/1.7ML	5	
XTRENBO SUBCUTANEOUS SOLUTION 120 MG/1.7ML	4	
YORVIPATH SUBCUTANEOUS SOLUTION PEN-INJECTOR 168 MCG/0.56ML, 294 MCG/0.98ML, 420 MCG/1.4ML	5	PA
Ophthalmic Agents - Treatment Of Eye Conditions		
Ophthalmic Agents, Other		
<i>atropine sulfate ophthalmic solution 1 %</i>	2	
<i>brimonidine tartrate-timolol ophthalmic solution 0.2-0.5 %</i>	2	
<i>cyclosporine (pf) ophthalmic emulsion 0.05 %</i>	2	QL (60 EA per 30 days)
CYSTARAN OPHTHALMIC SOLUTION 0.44 %	5	PA
<i>dorzolamide hcl-timolol mal ophthalmic solution 2-0.5 %</i>	2	
<i>dorzolamide hcl-timolol mal pf ophthalmic solution 2-0.5 %</i>	2	ST
<i>neomycin-polymyxin-dexameth ophthalmic ointment 3.5-10000-0.1</i>	1	
<i>neomycin-polymyxin-dexameth ophthalmic suspension 0.1 %, 3.5-10000-0.1</i>	1	
<i>neomycin-polymyxin-gramicidin ophthalmic solution 1.75-10000-.025</i>	2	
OXERVATE OPHTHALMIC SOLUTION 0.002 %	5	PA

Name of Drug	Drug Tier	Requirements/Limits
<i>sulfacetamide-prednisolone ophthalmic solution 10-0.23 %</i>	2	
TEPEZZA INTRAVENOUS SOLUTION RECONSTITUTED 500 MG	5	PA
<i>tobramycin-dexamethasone ophthalmic suspension 0.3-0.1 %</i>	2	
XDEMVY OPHTHALMIC SOLUTION 0.25 %	5	PA
Ophthalmic Anti-Allergy Agents		
<i>azelastine hcl ophthalmic solution 0.05 %</i>	1	
<i>cromolyn sodium ophthalmic solution 4 %</i>	1	
Ophthalmic Anti-Infectives		
<i>bacitracin-polymyxin b ophthalmic ointment 500-10000 unit/gm</i>	1	
<i>ciprofloxacin hcl ophthalmic solution 0.3 %</i>	2	
<i>erythromycin ophthalmic ointment 5 mg/gm</i>	1	
<i>gentamicin sulfate ophthalmic solution 0.3 %</i>	1	
<i>moxifloxacin hcl ophthalmic solution 0.5 %</i>	2	
NATACYN OPHTHALMIC SUSPENSION 5 %	4	
<i>ofloxacin ophthalmic solution 0.3 %</i>	2	
<i>polymyxin b-trimethoprim ophthalmic solution 10000-0.1 unit/ml-%</i>	1	
<i>sulfacetamide sodium ophthalmic solution 10 %</i>	2	
<i>tobramycin ophthalmic solution 0.3 %</i>	1	
Ophthalmic Anti-Inflammatories		
<i>dexamethasone sodium phosphate ophthalmic solution 0.1 %</i>	2	
<i>diclofenac sodium ophthalmic solution 0.1 %</i>	2	
<i>difluprednate ophthalmic emulsion 0.05 %</i>	2	
<i>fluorometholone ophthalmic suspension 0.1 %</i>	2	
<i>flurbiprofen sodium ophthalmic solution 0.03 %</i>	2	
<i>ketorolac tromethamine ophthalmic solution 0.4 %, 0.5 %</i>	2	
<i>prednisolone acetate ophthalmic suspension 1 %</i>	2	
<i>prednisolone sodium phosphate ophthalmic solution 1 %</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
Ophthalmic Beta-Adrenergic Blocking Agents		
<i>carteolol hcl ophthalmic solution 1 %</i>	2	
<i>levobunolol hcl ophthalmic solution 0.5 %</i>	1	
<i>timolol maleate ophthalmic solution 0.25 %, 0.5 %</i>	1	
Ophthalmic Intraocular Pressure Lowering Agents, Other		
<i>acetazolamide er oral capsule extended release 12 hour 500 mg</i>	2	
<i>brimonidine tartrate ophthalmic solution 0.1 %</i>	2	
<i>brimonidine tartrate ophthalmic solution 0.2 %</i>	1	
<i>brinzolamide ophthalmic suspension 1 %</i>	2	ST
<i>dorzolamide hcl ophthalmic solution 2 %</i>	2	
<i>methazolamide oral tablet 25 mg, 50 mg</i>	2	
<i>pilocarpine hcl ophthalmic solution 1 %, 1.25 %, 2 %, 4 %</i>	2	
RHOPRESSA OPHTHALMIC SOLUTION 0.02 %	3	ST
ROCKLATAN OPHTHALMIC SOLUTION 0.02-0.005 %	3	ST
SIMBRINZA OPHTHALMIC SUSPENSION 1-0.2 %	3	
Ophthalmic Prostaglandin And Prostamide Analogs		
<i>latanoprost ophthalmic solution 0.005 %</i>	1	
LUMIGAN OPHTHALMIC SOLUTION 0.01 %	3	
<i>travoprost (bak free) ophthalmic solution 0.004 %</i>	2	
Otic Agents - Treatment Of Ear Conditions		
Otic Agents		
<i>acetic acid otic solution 2 %</i>	2	
<i>hydrocortisone-acetic acid otic solution 1-2 %</i>	2	
<i>neomycin-polymyxin-hc otic solution 1 %, 3.5-10000-1</i>	2	

Name of Drug	Drug Tier	Requirements/Limits
<i>neomycin-polymyxin-hc otic suspension 3.5-10000-1</i>	2	
<i>ofloxacin otic solution 0.3 %</i>	2	
Respiratory Tract/ Pulmonary Agents - Treatment Of Breathing Conditions		
Antihistamines		
<i>azelastine hcl nasal solution 0.1 %, 137 mcg/spray</i>	1	
<i>cetirizine hcl oral solution 1 mg/ml, 5 mg/5ml</i>	1	
<i>clemastine fumarate oral tablet 2.68 mg</i>	2	
<i>cyproheptadine hcl oral syrup 2 mg/5ml</i>	2	PA
<i>cyproheptadine hcl oral tablet 4 mg</i>	2	PA
<i>hydroxyzine hcl oral syrup 10 mg/5ml</i>	2	PA
<i>hydroxyzine hcl oral tablet 10 mg, 25 mg, 50 mg</i>	1	PA
<i>levocetirizine dihydrochloride oral solution 2.5 mg/5ml</i>	2	
<i>levocetirizine dihydrochloride oral tablet 5 mg</i>	1	
<i>promethazine hcl oral solution 6.25 mg/5ml</i>	2	PA
Anti-Inflammatories, Inhaled Corticosteroids		
ARNUITY ELLIPTA INHALATION AEROSOL POWDER BREATH ACTIVATED 100 MCG/ACT, 200 MCG/ACT, 50 MCG/ACT	3	
<i>budesonide inhalation suspension 0.25 mg/2ml, 0.5 mg/2ml</i>	2	B/D; QL (120 ML per 30 days)
<i>budesonide inhalation suspension 1 mg/2ml</i>	2	B/D; QL (60 ML per 30 days)
<i>flunisolide nasal solution 25 mcg/act (0.025%)</i>	2	
<i>fluticasone propionate diskus inhalation aerosol powder breath activated 100 mcg/act, 250 mcg/act, 50 mcg/act</i>	2	
<i>fluticasone propionate hfa inhalation aerosol 110 mcg/act</i>	2	QL (12 GM per 30 days)
<i>fluticasone propionate hfa inhalation aerosol 220 mcg/act</i>	2	QL (24 GM per 30 days)
<i>fluticasone propionate hfa inhalation aerosol 44 mcg/act</i>	2	QL (10.6 GM per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
<i>fluticasone propionate nasal suspension 50 mcg/act</i>	2	QL (16 GM per 30 days)
<i>mometasone furoate nasal suspension 50 mcg/act</i>	2	
Bronchodilators, Anticholinergic		
INCRUSE ELLIPTA INHALATION AEROSOL POWDER BREATH ACTIVATED 62.5 MCG/ACT	3	QL (30 EA per 30 days)
<i>ipratropium bromide hfa inhalation aerosol solution 17 mcg/act</i>	2	QL (25.8 GM per 30 days)
<i>ipratropium bromide inhalation solution 0.02 %</i>	2	B/D
<i>ipratropium bromide nasal solution 0.03 %, 0.06 %</i>	2	
SPIRIVA RESPIMAT INHALATION AEROSOL SOLUTION 1.25 MCG/ACT, 2.5 MCG/ACT	3	QL (4 GM per 30 days)
<i>tiotropium bromide inhalation capsule 18 mcg</i>	2	QL (90 EA per 90 days)
Bronchodilators, Sympathomimetic		
<i>albuterol sulfate hfa inhalation aerosol solution 108 (90 base) mcg/act, 108 (90 base) mcg/act (nda020503), 108 (90 base) mcg/act (nda020983)</i>	1	QL (36 GM per 30 days)
<i>albuterol sulfate inhalation nebulization solution (2.5 mg/3ml) 0.083%, (5 mg/ml) 0.5%, 0.63 mg/3ml, 1.25 mg/3ml, 2.5 mg/0.5ml</i>	1	B/D
<i>albuterol sulfate oral syrup 2 mg/5ml</i>	1	
<i>albuterol sulfate oral tablet 2 mg, 4 mg</i>	2	
<i>epinephrine injection solution auto-injector 0.15 mg/0.15ml, 0.15 mg/0.3ml, 0.3 mg/0.3ml</i>	2	QL (2 EA per 30 days)
<i>formoterol fumarate inhalation nebulization solution 20 mcg/2ml</i>	2	B/D
<i>levalbuterol hcl inhalation nebulization solution 0.31 mg/3ml, 0.63 mg/3ml, 1.25 mg/3ml</i>	2	B/D
SEREVENT DISKUS INHALATION AEROSOL POWDER BREATH ACTIVATED 50 MCG/ACT	3	QL (60 EA per 30 days)
<i>terbutaline sulfate oral tablet 2.5 mg, 5 mg</i>	2	
VENTOLIN HFA INHALATION AEROSOL SOLUTION 108 (90 BASE) MCG/ACT	3	QL (36 GM per 30 days)
Cystic Fibrosis Agents		

Name of Drug	Drug Tier	Requirements/Limits
ALYFTREK ORAL TABLET 10-50-125 MG	5	PA; QL (56 EA per 28 days)
ALYFTREK ORAL TABLET 4-20-50 MG	5	PA; QL (84 EA per 28 days)
CAYSTON INHALATION SOLUTION RECONSTITUTED 75 MG	5	PA
KALYDECO ORAL PACKET 13.4 MG, 25 MG, 5.8 MG, 50 MG, 75 MG	5	PA
KALYDECO ORAL TABLET 150 MG	5	PA
ORKAMBI ORAL PACKET 100-125 MG, 150- 188 MG, 75-94 MG	5	PA
ORKAMBI ORAL TABLET 100-125 MG, 200- 125 MG	5	PA
PULMOZYME INHALATION SOLUTION 2.5 MG/2.5ML	5	B/D
SYMDEKO ORAL TABLET THERAPY PACK 100-150 & 150 MG, 50-75 & 75 MG	5	PA
TOBI PODHALER INHALATION CAPSULE 28 MG	5	PA; QL (224 EA per 56 days)
<i>tobramycin inhalation nebulization solution 300 mg/4ml</i>	3	B/D
<i>tobramycin inhalation nebulization solution 300 mg/5ml</i>	3	B/D; QL (280 ML per 56 days)
TRIKAFTA ORAL TABLET THERAPY PACK 100-50-75 & 150 MG, 50-25-37.5 & 75 MG	5	PA
TRIKAFTA ORAL THERAPY PACK 100-50-75 & 75 MG, 80-40-60 & 59.5 MG	5	PA
Mast Cell Stabilizers		
<i>cromolyn sodium inhalation nebulization solution 20 mg/2ml</i>	3	B/D
<i>cromolyn sodium oral concentrate 100 mg/5ml</i>	2	
Phosphodiesterase Inhibitors, Airways Disease		
<i>roflumilast oral tablet 250 mcg, 500 mcg</i>	2	
<i>theophylline er oral tablet extended release 12 hour 100 mg, 200 mg, 300 mg, 450 mg</i>	2	
<i>theophylline er oral tablet extended release 24 hour 400 mg, 600 mg</i>	2	
<i>theophylline oral solution 80 mg/15ml</i>	2	
Pulmonary Antihypertensives		

Name of Drug	Drug Tier	Requirements/Limits
ADEMPAS ORAL TABLET 0.5 MG, 1 MG, 1.5 MG, 2 MG, 2.5 MG	5	PA
<i>ambrisentan oral tablet 10 mg, 5 mg</i>	5	PA
<i>bosentan oral tablet 125 mg, 62.5 mg</i>	5	PA
OPSUMIT ORAL TABLET 10 MG	5	PA; QL (30 EA per 30 days)
<i>sildenafil citrate oral suspension reconstituted 10 mg/ml</i>	4	PA; QL (720 ML per 30 days)
<i>sildenafil citrate oral tablet 20 mg</i>	2	PA
<i>tadalafil (pah) oral tablet 20 mg</i>	2	PA
TADLIQ ORAL SUSPENSION 20 MG/5ML	5	PA
TYVASO DPI MAINTENANCE KIT INHALATION POWDER 112 X 32MCG & 112 X64MCG, 112 X 48MCG & 112 X64MCG, 112 X 48MCG & 112 X80MCG, 16 MCG, 32 MCG, 48 MCG, 64 MCG, 80 MCG	5	PA
TYVASO DPI TITRATION KIT INHALATION POWDER 16 & 32 & 48 MCG	5	PA
UPTRAVI ORAL TABLET 1000 MCG, 1200 MCG, 1400 MCG, 1600 MCG, 200 MCG, 400 MCG, 600 MCG, 800 MCG	5	PA
UPTRAVI TITRATION ORAL TABLET THERAPY PACK 200 & 800 MCG	5	PA
WINREVAIR SUBCUTANEOUS KIT 2 X 45 MG, 2 X 60 MG, 45 MG, 60 MG	5	PA
YUTREPIA INHALATION CAPSULE 106 MCG, 26.5 MCG, 53 MCG, 79.5 MCG	5	PA
Pulmonary Fibrosis Agents		
<i>nintedanib esylate oral capsule 100 mg, 150 mg</i>	5	PA
OFEV ORAL CAPSULE 100 MG, 150 MG	5	PA
<i>pirfenidone oral capsule 267 mg</i>	5	PA
<i>pirfenidone oral tablet 267 mg</i>	2	PA
<i>pirfenidone oral tablet 534 mg, 801 mg</i>	5	PA
Respiratory Tract Agents, Other		
<i>acetylcysteine inhalation solution 10 %, 20 %</i>	2	B/D
ADVAIR HFA INHALATION AEROSOL 115-21 MCG/ACT, 230-21 MCG/ACT, 45-21 MCG/ACT	3	QL (12 GM per 30 days)

Name of Drug	Drug Tier	Requirements/Limits
ANORO ELLIPTA INHALATION AEROSOL POWDER BREATH ACTIVATED 62.5-25 MCG/ACT	3	QL (60 EA per 30 days)
BEVESPI AEROSPHERE INHALATION AEROSOL 9-4.8 MCG/ACT	3	
BREO ELLIPTA INHALATION AEROSOL POWDER BREATH ACTIVATED 100-25 MCG/ACT, 200-25 MCG/ACT, 50-25 MCG/INH	3	QL (60 EA per 30 days)
BREZTRI AEROSPHERE INHALATION AEROSOL 160-18-4.8 MCG/ACT, 160-9-4.8 MCG/ACT	3	QL (10.7 GM per 30 days)
BRINSUPRI ORAL TABLET 10 MG, 25 MG	5	PA; QL (30 EA per 30 days)
<i>budesonide-formoterol fumarate inhalation aerosol 160-4.5 mcg/act, 80-4.5 mcg/act</i>	2	QL (10.2 GM per 30 days)
COMBIVENT RESPIMAT INHALATION AEROSOL SOLUTION 20-100 MCG/ACT	4	QL (8 GM per 30 days)
DUPIXENT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 200 MG/1.14ML, 300 MG/2ML	5	PA
DUPIXENT SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 200 MG/1.14ML, 300 MG/2ML	5	PA
FASENRA PEN SUBCUTANEOUS SOLUTION AUTO-INJECTOR 30 MG/ML	5	PA
FASENRA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 10 MG/0.5ML, 30 MG/ML	5	PA
<i>fluticasone-salmeterol inhalation aerosol powder breath activated 100-50 mcg/act, 113-14 mcg/act, 232-14 mcg/act, 250-50 mcg/act, 500-50 mcg/act, 55-14 mcg/act</i>	2	QL (60 EA per 30 days)
<i>ipratropium-albuterol inhalation solution 0.5-2.5 (3) mg/3ml</i>	2	B/D
<i>montelukast sodium oral packet 4 mg</i>	2	
<i>montelukast sodium oral tablet 10 mg</i>	1	
<i>montelukast sodium oral tablet chewable 4 mg, 5 mg</i>	1	
NUCALA SUBCUTANEOUS SOLUTION AUTO-INJECTOR 100 MG/ML	5	PA

Name of Drug	Drug Tier	Requirements/Limits
NUCALA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 100 MG/ML, 40 MG/0.4ML	5	PA
NUCALA SUBCUTANEOUS SOLUTION RECONSTITUTED 100 MG	5	PA
<i>promethazine-phenylephrine oral syrup 6.25-5 mg/5ml</i>	2	PA
STIOLTO RESPIMAT INHALATION AEROSOL SOLUTION 2.5-2.5 MCG/ACT	3	QL (4 GM per 30 days)
TRELEGY ELLIPTA INHALATION AEROSOL POWDER BREATH ACTIVATED 100-62.5-25 MCG/ACT, 200-62.5-25 MCG/ACT	3	QL (60 EA per 30 days)
WIXELA INHUB INHALATION AEROSOL POWDER BREATH ACTIVATED 100-50 MCG/ACT, 250-50 MCG/ACT, 500-50 MCG/ACT	2	QL (60 EA per 30 days)
XOLAIR SUBCUTANEOUS SOLUTION AUTO-INJECTOR 150 MG/ML, 300 MG/2ML, 75 MG/0.5ML	5	PA
XOLAIR SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 150 MG/ML, 300 MG/2ML, 75 MG/0.5ML	5	PA
XOLAIR SUBCUTANEOUS SOLUTION RECONSTITUTED 150 MG	5	PA

Skeletal Muscle Relaxants - Treatment Of Muscle Tightness

Skeletal Muscle Relaxants

<i>carisoprodol oral tablet 250 mg, 350 mg</i>	2	PA; QL (90 EA per 30 days)
<i>chlorzoxazone oral tablet 500 mg</i>	2	PA; QL (180 EA per 30 days)
<i>cyclobenzaprine hcl oral tablet 10 mg, 5 mg</i>	1	QL (90 EA per 30 days)
<i>metaxalone oral tablet 800 mg</i>	2	PA; QL (120 EA per 30 days)
<i>methocarbamol oral tablet 500 mg, 750 mg</i>	1	PA
<i>orphenadrine citrate er oral tablet extended release 12 hour 100 mg</i>	2	PA

Sleep Disorder Agents - Treatment Of Insomnia

Sleep Promoting Agents

<i>doxepin hcl oral tablet 3 mg, 6 mg</i>	2	QL (30 EA per 30 days)
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Name of Drug	Drug Tier	Requirements/Limits
<i>eszopiclone oral tablet 1 mg, 2 mg, 3 mg</i>	2	PA; QL (30 EA per 30 days)
HETLIOZ LQ ORAL SUSPENSION 4 MG/ML	5	PA
<i>ramelteon oral tablet 8 mg</i>	2	QL (30 EA per 30 days)
<i>tasimelteon oral capsule 20 mg</i>	5	PA
<i>temazepam oral capsule 15 mg, 22.5 mg, 30 mg, 7.5 mg</i>	2	PA; QL (30 EA per 30 days)
<i>zaleplon oral capsule 10 mg</i>	2	PA; QL (60 EA per 30 days)
<i>zaleplon oral capsule 5 mg</i>	2	PA; QL (30 EA per 30 days)
ZEPBOUND KWIKPEN SUBCUTANEOUS SOLUTION PEN-INJECTOR 10 MG/0.6ML, 12.5 MG/0.6ML, 15 MG/0.6ML, 2.5 MG/0.6ML, 5 MG/0.6ML, 7.5 MG/0.6ML	4	PA; QL (2.4 ML per 28 days)
ZEPBOUND SUBCUTANEOUS SOLUTION AUTO-INJECTOR 10 MG/0.5ML, 12.5 MG/0.5ML, 15 MG/0.5ML, 2.5 MG/0.5ML, 5 MG/0.5ML, 7.5 MG/0.5ML	5	PA; QL (2 ML per 28 days)
<i>zolpidem tartrate er oral tablet extended release 12.5 mg, 6.25 mg</i>	2	PA; QL (30 EA per 30 days)
<i>zolpidem tartrate oral tablet 10 mg</i>	2	PA; QL (30 EA per 30 days)
<i>zolpidem tartrate oral tablet 5 mg</i>	2	QL (30 EA per 30 days)
Wakefulness Promoting Agents		
<i>armodafinil oral tablet 150 mg, 200 mg, 250 mg, 50 mg</i>	2	PA
<i>modafinil oral tablet 100 mg, 200 mg</i>	2	PA
<i>sodium oxybate oral solution 500 mg/ml</i>	5	PA; QL (540 ML per 30 days)
XYWAV ORAL SOLUTION 500 MG/ML	5	PA

Index

A	<i>abacavir sulfate</i>	49	<i>allopurinol</i>	31	AQNEURSA	72
<i>abacavir sulfate-lamivudine</i> ...	49	<i>alosetron hcl</i>	83	ARALAST NP.....	85	
ABIGALE	90	<i>alprazolam</i>	52	ARANELLE	90	
ABIGALE LO	90	ALPRAZOLAM INTENSOL	52	ARANESP (ALBUMIN FREE)		
ABILIFY ASIMTUFII	43	ALTAVERA	90		61	
ABILIFY MAINTENA.....	43	ALUNBRIG	35	ARCALYST	101	
<i>abiraterone acetate</i>	33	<i>alyacen 1/35</i>	90	AREXVY	107	
ABIRTEGA.....	33	ALYFTREK	117	ARIKAYCE	14	
ABRYSVO.....	107	<i>amantadine hcl</i>	41, 42	<i>aripiprazole</i>	44	
<i>acamprosate calcium</i>	13	<i>ambrisentan</i>	118	ARISTADA.....	44	
<i>acarbose</i>	53	<i>amikacin sulfate</i>	14	ARISTADA INITIO.....	44	
<i>acebutolol hcl</i>	64	<i>amiloride hcl</i>	68	<i>armodafinil</i>	121	
<i>acetaminophen-codeine</i>	12	<i>amiloride-hydrochlorothiazide</i>		ARNUITY ELLIPTA.....	115	
<i>acetazolamide</i>	66		66	<i>asenapine maleate</i>	44	
<i>acetazolamide er</i>	114	<i>amiodarone hcl</i>	64	ASHLYNA.....	90	
<i>acetic acid</i>	114	<i>amitriptyline hcl</i>	27	<i>aspirin-dipyridamole er</i>	62	
<i>acetylcysteine</i>	118	<i>amlodipine besy-benazepril hcl</i>		ASTAGRAF XL.....	105	
<i>acitretin</i>	76		66	<i>atazanavir sulfate</i>	51	
ACTEMRA	100	<i>amlodipine besylate</i>	65	<i>atenolol</i>	64	
ACTEMRA ACTPEN.....	100	<i>amlodipine besylate-valsartan</i>	66	<i>atenolol-chlorthalidone</i>	66	
ACTHIB	107	<i>amlodipine-atorvastatin</i>	66	<i>atomoxetine hcl</i>	71	
ACTIMMUNE	104	<i>amlodipine-olmesartan</i>	66	<i>atorvastatin calcium</i>	69	
<i>acyclovir</i>	48, 79	<i>amlodipine-valsartan-hctz</i>	66	<i>atovaquone</i>	41	
<i>acyclovir sodium</i>	49	<i>ammonium lactate</i>	76	<i>atovaquone-proguanil hcl</i>	41	
ADACEL.....	107	AMNESTEEM	76	<i>atropine sulfate</i>	112	
<i>adalimumab-fkjp (2 pen)</i>	105	<i>amoxapine</i>	28	AUBRA EQ.....	90	
<i>adalimumab-fkjp (2 syringe)</i>	105	<i>amoxicillin</i>	18	AUGTYRO.....	35	
<i>adapalene</i>	76	<i>amoxicillin-pot clavulanate</i>	18	AUROVELA 1.5/30	90	
<i>adapalene-benzoyl peroxide</i> ...	76	<i>amoxicillin-pot clavulanate er</i>	18	AUROVELA 1/20	90	
<i>adefovir dipivoxil</i>	48	<i>amphetamine-dextroamphet er</i>		AUROVELA 24 FE	90	
ADEMPAS	118		71	AUROVELA FE 1.5/30	90	
ADVAIR HFA	118	<i>amphetamine-</i>		AUROVELA FE 1/20	90	
AIMOVIG.....	31	<i>dextroamphetamine</i>	71	AUSTEDO	72	
AKEEGA	34	<i>amphotericin b</i>	29	AUSTEDO XR.....	72	
<i>albendazole</i>	41	<i>amphotericin b liposome</i>	29	AUSTEDO XR PATIENT		
<i>albuterol sulfate</i>	116	<i>ampicillin</i>	18	TITRATION.....	73	
<i>albuterol sulfate hfa</i>	116	<i>ampicillin sodium</i>	18	AUVELITY	25	
<i>alclometasone dipropionate</i> ...	76	<i>ampicillin-sulbactam sodium</i> .	18	AUVELITY TITRATION		
<i>alcohol</i>	78, 79	<i>anagrelide hcl</i>	61	PACK	25	
ALCOHOL.....	78	<i>anastrozole</i>	35	AVIANE.....	90	
ALECENSA	35	ANORO ELLIPTA.....	119	AVMAPKI FAKZYNJA CO-		
<i>alendronate sodium</i>	111	<i>apomorphine hcl</i>	42	PACK	34	
<i>alfuzosin hcl er</i>	87	<i>aprepitant</i>	29	AVTOZMA	101	
<i>aliskiren fumarate</i>	66	APRI.....	90	AYUNA.....	90	
		APTIVUS	51	AYVAKIT	36	

<i>azathioprine</i>	105
<i>azelastine hcl</i>	113, 115
<i>azithromycin</i>	19
<i>aztreonam</i>	14
AZURETTE.....	90
B	
BAC (BUTALBITAL- ACETAMIN-CAFF).....	10
<i>bacitracin-polymyxin b</i>	113
<i>baclofen</i>	47
BAFIERTAM.....	74
<i>balsalazide disodium</i>	110
BALVERSA.....	36
BALZIVA.....	90
BAQSIMI ONE PACK.....	56
BAQSIMI TWO PACK.....	56
BARACLUDE.....	48
<i>bcg vaccine</i>	107
<i>benazepril hcl</i>	63
<i>benazepril-hydrochlorothiazide</i>	66
BENLYSTA.....	101
<i>benzoyl peroxide-erythromycin</i>	76
<i>benztropine mesylate</i>	41
BESREMI.....	34
<i>betaine</i>	85
<i>betamethasone dipropionate</i> ..	77
<i>betamethasone dipropionate aug</i>	76, 77
<i>betamethasone valerate</i>	77
BETASERON.....	74
<i>betaxolol hcl</i>	64
<i>bethanechol chloride</i>	87
BEVESPI AEROSPHERE... 119	
<i>bexarotene</i>	40
BEXSERO.....	107
BEYFORTUS.....	107
<i>bicalutamide</i>	33
BICILLIN L-A.....	18
BIKTARVY.....	50
<i>bisoprolol fumarate</i>	64
<i>bisoprolol-hydrochlorothiazide</i>	67
BLISOVI 24 FE.....	90
BLISOVI FE 1.5/30.....	90
BLISOVI FE 1/20.....	90
BONSITY.....	111

H4676_26DCF_C V14

BOOSTRIX.....	107
<i>bosentan</i>	118
BOSULIF.....	36
<i>bosutinib</i>	36
BRAFTOVI.....	36
BREO ELLIPTA.....	119
BREZTRI AEROSPHERE... 119	
<i>briellyn</i>	90
BRILINTA.....	62
<i>brimonidine tartrate</i>	114
<i>brimonidine tartrate-timolol</i> ..	112
BRINSUPRI.....	119
<i>brinzolamide</i>	114
<i>brivaracetam</i>	21
<i>bromocriptine mesylate</i>	42
BRUKINSA.....	36
<i>budesonide</i>	110, 115
<i>budesonide er</i>	110
<i>budesonide-formoterol fumarate</i>	119
<i>bumetanide</i>	68
<i>buprenorphine</i>	11
<i>buprenorphine hcl</i>	13
<i>buprenorphine hcl-naloxone hcl</i>	13
<i>bupropion hcl</i>	26
<i>bupropion hcl er (smoking det)</i>	14
<i>bupropion hcl er (sr)</i>	26
<i>bupropion hcl er (xl)</i>	26
<i>buspirone hcl</i>	52
<i>butalbital-acetaminophen</i>	10
<i>butalbital-apap-caff-cod</i>	10
<i>butalbital-apap-caffeine</i>	10
<i>butalbital-asa-caff-codeine</i> ...	10
<i>butalbital-aspirin-caffeine</i>	10
<i>butorphanol tartrate</i>	12
BYSANTI.....	73
BYSANTI TITRATION PACK A.....	73
BYSANTI TITRATION PACK B.....	73
BYSANTI TITRATION PACK C.....	73
C	
<i>cabergoline</i>	98
CABOMETYX.....	36
<i>calcipotriene</i>	79

Formulary ID 26314, Version 14

Page 123

<i>calcitonin (salmon)</i>	111
<i>calcitriol</i>	79, 111
<i>calcium acetate (phos binder)</i> 82	
CALQUENCE.....	36
CAMILA.....	96
CAMZYOS.....	67
<i>candesartan cilexetil</i>	63
<i>candesartan cilexetil-hctz</i>	67
CAPLYTA.....	44
CAPRELSA.....	36
<i>captopril</i>	63
<i>carbamazepine</i>	24
<i>carbamazepine er</i>	24
<i>carbidopa</i>	42
<i>carbidopa-levodopa</i>	42
<i>carbidopa-levodopa er</i>	42
<i>carbidopa-levodopa-entacapone</i>	42
CARDAMYST.....	65
<i>carglumic acid</i>	80
<i>carisoprodol</i>	120
<i>carteolol hcl</i>	114
CARTIA XT.....	65
<i>carvedilol</i>	64
<i>casprofungin acetate</i>	29
CAYSTON.....	117
<i>cefaclor</i>	16
<i>cefaclor er</i>	16
<i>cefadroxil</i>	16
<i>cefazolin sodium</i>	16
<i>cefdinir</i>	16
<i>cefepime hcl</i>	16
<i>cefepime-dextrose</i>	17
<i>cefixime</i>	17
<i>cefoxitin sodium</i>	17
<i>cefpodoxime proxetil</i>	17
<i>cefprozil</i>	17
<i>ceftaroline fosamil</i>	17
<i>ceftazidime</i>	17
<i>ceftriaxone sodium</i>	17
<i>ceftriaxone sodium in dextrose</i>	17
<i>ceftriaxone sodium-dextrose</i> ..	17
<i>cefuroxime axetil</i>	17
<i>cefuroxime sodium</i>	17
<i>celecoxib</i>	10
<i>cephalexin</i>	17
CERDELGA.....	85

Last Updated 7/2026

<i>cetirizine hcl</i>	115
<i>cevimeline hcl</i>	76
CHARLOTTE 24 FE	91
CHATEAL EQ	91
<i>chlorhexidine gluconate</i>	76
<i>chloroquine phosphate</i>	41
<i>chlorpromazine hcl</i>	28
<i>chlorthalidone</i>	69
<i>chlorzoxazone</i>	120
CHOLBAM	85
<i>cholestyramine</i>	69
<i>cholestyramine light</i>	69
CIBINQO	101
<i>ciclopirox</i>	79
<i>ciclopirox olamine</i>	80
<i>cilostazol</i>	62
CIMDUO	50
<i>cimetidine</i>	84
CIMZIA	105
CIMZIA (1 SYRINGE)	105
CIMZIA (2 SYRINGE)	105
CIMZIA-STARTER	105
<i>cinacalcet hcl</i>	111
CINRYZE	100
<i>ciprofloxacin hcl</i>	20, 113
<i>ciprofloxacin in d5w</i>	20
<i>citalopram hydrobromide</i>	26
<i>cladribine (10 tabs)</i>	74
<i>cladribine (4 tabs)</i>	74
<i>cladribine (5 tabs)</i>	74
<i>cladribine (6 tabs)</i>	74
<i>cladribine (7 tabs)</i>	74
<i>cladribine (8 tabs)</i>	74
<i>cladribine (9 tabs)</i>	74
<i>cladribine 10 mg tablet therapy</i> <i>pack oral</i>	74
CLARAVIS	76
<i>clarithromycin</i>	19
<i>clarithromycin er</i>	19
<i>clemastine fumarate</i>	115
<i>clindamycin hcl</i>	14
<i>clindamycin palmitate hcl</i>	15
<i>clindamycin phos (once-daily)</i>	80
<i>clindamycin phos (twice-daily)</i>	80
<i>clindamycin phos-benzoyl perox</i>	76
<i>clindamycin phosphate</i>	15, 80

H4676_26DCF_C V14

<i>clindamycin phosphate in d5w</i>	15
<i>clindamycin phosphate in nacl</i>	15
CLINISOL SF	82
<i>clobazam</i>	22, 23
<i>clobetasol prop emollient base</i>	77
<i>clobetasol propionate</i>	77
<i>clobetasol propionate e</i>	77
<i>clomipramine hcl</i>	28
<i>clonazepam</i>	52
<i>clonidine</i>	63
<i>clonidine hcl</i>	63
<i>clonidine hcl er</i>	71
<i>clopidogrel bisulfate</i>	62
<i>clorazepate dipotassium</i>	52
<i>clotrimazole</i>	29
<i>clotrimazole-betamethasone</i>	79
<i>clozapine</i>	47
COARTEM	41
COBENFY	73
COBENFY STARTER PACK	73
<i>colchicine</i>	31
<i>colchicine-probenecid</i>	31
<i>colesevelam hcl</i>	69
<i>colestipol hcl</i>	69
<i>colistimethate sodium (cba)</i>	15
COMBIPATCH	91
COMBIVENT RESPIMAT	119
COMETRIQ (100 MG DAILY DOSE)	36
COMETRIQ (140 MG DAILY DOSE)	36
COMETRIQ (60 MG DAILY DOSE)	36
<i>constulose</i>	83
COPIKTRA	36
CORLANOR	67
CORTROPHIN	87
CORTROPHIN GEL	87
COSENTYX	101
COSENTYX (300 MG DOSE)	101
COSENTYX SENSOREADY (300 MG)	101
COSENTYX SENSOREADY PEN	101

Formulary ID 26314, Version 14

Page 124

COSENTYX UNOREADY	101
COTELLIC	36
CREON	85
CRESEMBA	29
<i>cromolyn sodium</i>	113, 117
CRYSELLE	91
CRYSELLE-28	91
CUVRIOR	81
<i>cyclobenzaprine hcl</i>	120
<i>cyclophosphamide</i>	32
<i>cyclosporine</i>	105
<i>cyclosporine (pf)</i>	112
<i>cyclosporine modified</i>	105
<i>cyproheptadine hcl</i>	115
CYRED EQ	91
CYSTAGON	85
CYSTARAN	112
D	
<i>dabigatran etexilate mesylate</i>	60
<i>dalfampridine er</i>	74
<i>danazol</i>	89
<i>dantrolene sodium</i>	48
DANZITEN	34
<i>dapagliflozin</i>	53
<i>dapsone</i>	32
DAPTACEL	107
<i>daptomycin</i>	15
<i>darifenacin hydrobromide er</i>	86
<i>darunavir</i>	51
<i>dasatinib</i>	36
DASETTA 1/35 (28)	91
DASETTA 7/7/7	91
DAURISMO	36
DAYSEE	91
DEBLITANE	96
<i>deferasirox</i>	81
<i>deferasirox granules</i>	81
<i>deferiprone</i>	81
DELSTRIGO	50
DEPO-PROVERA	91
DEPO-SUBQ PROVERA	104
.....	96
DESCOVY	50
<i>desipramine hcl</i>	28
<i>desmopressin ace spray refrig</i>	88
<i>desmopressin acetate</i>	88
<i>desmopressin acetate spray</i>	88
<i>desonide</i>	77

Last Updated 7/2026

desoximetasone.....77
desvenlafaxine succinate er....26
dexamethasone87, 110
DEXAMETHASONE
 INTENSOL 110
dexamethasone sodium
 phosphate 111, 113
dexmethylphenidate hcl 71
dexmethylphenidate hcl er.... 71
dextroamphetamine sulfate 71
dextroamphetamine sulfate er 71
dextrose82
dextrose-sodium chloride82
DIACOMIT.....21
diazepam.....23, 52, 53
DIAZEPAM INTENSOL.....52
diazoxide.....56
diclofenac epolamine.....10
diclofenac potassium10
diclofenac sodium.....10, 113
diclofenac sodium er10
dicloxacillin sodium18
dicyclomine hcl.....84
DIFICID19
diflunisal.....10
difluprednate 113
digoxin.....67
dihydroergotamine mesylate ..31
DILANTIN.....24
diltiazem hcl66
diltiazem hcl er66
diltiazem hcl er beads.....65
diltiazem hcl er coated beads .66
dilt-xr.....66
dimethyl fumarate.....74
dimethyl fumarate starter pack
 74
diphenoxylate-atropine.....83, 84
dipyridamole.....62
disopyramide phosphate.....64
disulfiram13
divalproex sodium21
divalproex sodium er21
dofetilide.....64
donepezil hcl.....25
DOPTELET.....62
DOPTELET SPRINKLE.....62
dorzolamide hcl 114

H4676_26DCF_C V14

dorzolamide hcl-timolol mal 112
dorzolamide hcl-timolol mal pf
 112
DOVATO50
doxazosin mesylate63
doxepin hcl28, 77, 120
doxercalciferol.....111
DOXY 10020
doxycycline hyclate.....20
doxycycline monohydrate21
DRIZALMA SPRINKLE.....74
dronabinol29
*drospirenone-ethinyl estradiol*91
DROXIA33
droxidopa.....63
DUAVEE.....97
duloxetine hcl26, 74
DUPIXENT 119
dutasteride87
E
econazole nitrate29
EDURANT PED49
efavirenz49
efavirenz-emtricitab-tenofo df 50
efavirenz-lamivudine-tenofovir
 50
EGRIFTA SV88
EGRIFTA WR.....88
ELIGARD98
ELINEST.....91
ELIQUIS60
ELIQUIS (1.5 MG PACK).....60
ELIQUIS (2 MG PACK).....60
ELIQUIS DVT/PE STARTER
 PACK60
ELMIRON.....87
eltrombopag olamine.....61
ELURYNG.....91
EMEND.....29
EMGALITY31
EMGALITY (300 MG DOSE)
 31
EMSAM26
emtricitabine.....50
emtricitabine-tenofovir df.....50
emtricitab- rilpivir-tenofov df..50
EMTRIVA.....50
EMZAHH.....91

Formulary ID 26314, Version 14

Page 125

enalapril maleate.....63
*enalapril-hydrochlorothiazide*67
ENBREL.....105
ENBREL MINI105
ENBREL SURECLICK105
ENDOCET12
ENFLONIA107
ENGERIX-B.....107
ENILLORING.....91
ENOBY111
enoxaparin sodium60
ENSACOVE.....36
ENSKYCE.....91
entacapone.....42
entecavir48
ENTRESTO.....67
ENTYVIO PEN.....101
enulose83
ENVARUS XR105
EPIDIOLEX21
epinephrine116
eplerenone68
EPOGEN61
EQUETRO53
ergotamine-caffeine31
ERIVEDGE36
ERLEADA33
erlotinib hcl36
ERRIN96
ertapenem sodium.....19
ERVEBO107
ery80
ERYTHROCIN
 LACTOBIONATE19
erythromycin80, 113
erythromycin base19
erythromycin ethylsuccinate ...20
ERZOFRI44
escitalopram oxalate26, 27
eslicarbazepine acetate24
esomeprazole magnesium .84, 85
ESTARYLLA.....91
estradiol89
estradiol valerate.....89
estradiol-norethindrone acet..91
eszopiclone121
ethambutol hcl32
ethosuximide22

Last Updated 7/2026

<i>etodolac</i>	11	FIRMAGON.....	98	GAMMAKED	100
<i>etodolac er</i>	10	FIRMAGON (240 MG DOSE)		GAMMAPLEX	100
<i>etonogestrel-ethinyl estradiol</i>	91		98	GAMUNEX-C.....	100
<i>etravirine</i>	49	<i>flavoxate hcl</i>	86	GARDASIL 9	108
EUCRISA.....	77	<i>flecainide acetate</i>	64	GATTEX	84
EULEXIN.....	33	<i>fluconazole</i>	29	<i>gauze</i>	56
<i>everolimus</i>	36, 105, 106	<i>fluconazole in sodium chloride</i>		GAUZE.....	56
EVOTAZ.....	50		29	GAVILYTE-C	83
EVRYSDI.....	73	<i>flucytosine</i>	30	GAVILYTE-G.....	83
<i>exemestane</i>	35	<i>fludrocortisone acetate</i>	87	GAVILYTE-N WITH FLAVOR	
EXXUA.....	26	<i>flunisolide</i>	115	PACK	83
EXXUA TITRATION PACK	26	<i>fluocinolone acetonide</i>	77	GAVRETO	36
<i>ezetimibe</i>	70	<i>fluocinonide</i>	77, 78	<i>gefitinib</i>	36
<i>ezetimibe-simvastatin</i>	70	<i>fluocinonide emulsified base</i> ..	77	<i>gemfibrozil</i>	69
F		<i>fluorometholone</i>	113	GEMTESA	86
FABHALTA.....	101	<i>fluorouracil</i>	79	<i>generlac</i>	83
FALMINA.....	91	<i>fluoxetine hcl</i>	27	GENGRAF	106
<i>famciclovir</i>	49	<i>fluphenazine decanoate</i>	43	GENOTROPIN.....	88
<i>famotidine</i>	84	<i>fluphenazine hcl</i>	43	GENOTROPIN MINIQUICK	88
FANAPT	44	<i>flurbiprofen</i>	11	<i>gentamicin in saline</i>	14
FANAPT TITRATION PACK		<i>flurbiprofen sodium</i>	113	<i>gentamicin sulfate</i>	14, 80, 113
A	44	<i>fluticasone propionate</i>	78, 116	GENVOYA	50
FANAPT TITRATION PACK		<i>fluticasone propionate diskus</i>		GILOTRIF	37
B	44		115	GLASSIA	85
FANAPT TITRATION PACK		<i>fluticasone propionate hfa</i>	115	<i>glatiramer acetate</i>	74
C	44	<i>fluticasone-salmeterol</i>	119	GLATOPA	75
FASENRA.....	119	<i>fluvoxamine maleate</i>	27	<i>glimepiride</i>	53
FASENRA PEN	119	<i>fondaparinux sodium</i>	60	<i>glipizide</i>	53
<i>febuxostat</i>	31	<i>formoterol fumarate</i>	116	<i>glipizide er</i>	53
<i>felbamate</i>	21	<i>fosamprenavir calcium</i>	51	<i>glipizide-metformin hcl</i>	53, 54
<i>felodipine er</i>	65	<i>fosfomycin tromethamine</i>	15	<i>glucagon emergency</i>	56
<i>fenofibrate</i>	69	<i>fosinopril sodium</i>	63	<i>glyburide</i>	54
<i>fenofibrate micronized</i>	69	<i>fosinopril sodium-hctz</i>	67	<i>glyburide micronized</i>	54
<i>fenofibric acid</i>	69	FOTIVDA	36	<i>glyburide-metformin</i>	54
<i>fentanyl</i>	11	FRUZAQLA.....	36	<i>glycerol phenylbutyrate</i>	85
<i>fesoterodine fumarate er</i>	86	FULPHILA.....	61	<i>glycopyrrolate</i>	84
FETZIMA.....	27	<i>furosemide</i>	68	GLYXAMBI.....	54
FETZIMA TITRATION	27	FYAVOLV	92	GOCOVRI.....	42
FIASP	56	FYLNETRA	61	GOMEKLI.....	34
FIASP FLEXTOUCH	56	G		<i>granisetron hcl</i>	29
FIASP PENFILL	56	<i>gabapentin</i>	23	<i>griseofulvin microsize</i>	30
<i>fidaxomicin</i>	20	GALAFOLD	85	<i>guanfacine hcl</i>	63
FILSPARI.....	87	<i>galantamine hydrobromide</i>	25	<i>guanfacine hcl er</i>	71
<i>finasteride</i>	87	<i>galantamine hydrobromide er</i>	25	H	
<i>fin golimod hcl</i>	74	GAMMAGARD.....	100	HAEGARDA.....	100
FINTEPLA	21	GAMMAGARD ERC	100	HAILEY 1.5/30	92
FINZALA.....	91	GAMMAGARD S/D LESS IGA		HAILEY 24 FE.....	92
FIRDAPSE.....	73		100	HAILEY FE 1.5/30	92

H4676_26DCF_C V14

Formulary ID 26314, Version 14

Last Updated 7/2026

HAILEY FE 1/20	92	HYFTOR	78	INTELENCE	49
<i>halobetasol propionate</i>	78	HYRNUO	37	INTRALIPID	82
HALOETTE	92	I		INTROVALE	92
<i>haloperidol</i>	43	<i>ibandronate sodium</i>	111	INVEGA HAFYERA	45
<i>haloperidol decanoate</i>	43	IBRANCE	37	INVEGA SUSTENNA	45
<i>haloperidol lactate</i>	43	IBTROZI	37	INVEGA TRINZA	45
HAVRIX	108	IBU	11	IPOL	108
HEATHER	92	<i>ibuprofen</i>	11	<i>ipratropium bromide</i>	116
<i>heparin sodium (porcine)</i>	60	<i>icatibant acetate</i>	100	<i>ipratropium bromide hfa</i>	116
<i>heparin sodium (porcine) pf</i>	60	ICLEVIA	92	<i>ipratropium-albuterol</i>	119
HEPLISAV-B	108	ICLUSIG	37	<i>irbesartan</i>	63
HERNEXEOS	37	<i>icosapent ethyl</i>	70	<i>irbesartan-hydrochlorothiazide</i>	
HETLIOZ LQ	121	IDHIFA	34		67
HIBERIX	108	IDVYNZO	49	ISENTRESS	49
HUMALOG	56	ILARIS	101	ISENTRESS HD	49
HUMALOG JUNIOR		ILUMYA	101	ISIBLOOM	92
KWIKPEN	56	<i>imatinib mesylate</i>	37	ISOLYTE-P IN D5W	82
HUMALOG KWIKPEN	56	IMBRUVICA	37	ISOLYTE-S	80
HUMALOG MIX 50/50		<i>imipenem-cilastatin</i>	19	ISOLYTE-S PH 7.4	80
KWIKPEN	56	<i>imipramine hcl</i>	28	<i>isoniazid</i>	32
HUMALOG MIX 75/25	56	<i>imipramine pamoate</i>	28	<i>isosorb dinitrate-hydralazine</i>	70
HUMALOG MIX 75/25		<i>imiquimod</i>	79	<i>isosorbide dinitrate</i>	70
KWIKPEN	56	<i>imkeldi</i>	37	<i>isosorbide mononitrate</i>	70
HUMALOG TEMPO PEN	56	IMOVAX RABIES	108	<i>isosorbide mononitrate er</i>	70
HUMULIN 70/30	57	IMPAVIDO	41	<i>isotretinoin</i>	76
HUMULIN 70/30 KWIKPEN	57	IMULDOSA	101	<i>isradipine</i>	65
HUMULIN N	57	INCASSIA	96	ITOVEBI	37
HUMULIN N KWIKPEN	57	INCRELEX	88	<i>itraconazole</i>	30
HUMULIN R	57	INCRUSE ELLIPTA	116	<i>ivabradine hcl</i>	67
HUMULIN R U-500		<i>indapamide</i>	69	<i>ivermectin</i>	41
(CONCENTRATED)	57	<i>indomethacin</i>	11	IWILFIN	34
HUMULIN R U-500		<i>indomethacin er</i>	11	IXIARO	108
KWIKPEN	57	INFANRIX	108	J	
<i>hydralazine hcl</i>	70	INGREZZA	73	JAIMESS	92
<i>hydrochlorothiazide</i>	69	INLURIYO	34	JAKAFI	37
<i>hydrocodone-acetaminophen</i>	12	INLYTA	37	JAKAFI XR	37
<i>hydrocodone-ibuprofen</i>	12	INQOVI	33	JANTOVEN	60
<i>hydrocortisone</i>	78, 87, 111	INREBIC	37	JANUMET	54
<i>hydrocortisone (perianal)</i>	78	<i>insulin asp prot & asp flexpen</i>	57	JANUMET XR	54
<i>hydrocortisone butyrate</i>	78	<i>insulin aspart</i>	57	JANUVIA	54
<i>hydrocortisone valerate</i>	78	<i>insulin aspart flexpen</i>	57	JARDIANCE	54
<i>hydrocortisone-acetic acid</i> ...	114	<i>insulin aspart prot & aspart</i> ...	57	JASMIEL	92
<i>hydromorphone hcl</i>	12	<i>insulin lispro</i>	57	JAYPIRCA	37
<i>hydromorphone hcl pf</i>	12	<i>insulin lispro (1 unit dial)</i>	57	JENCYCLA	92
<i>hydroxychloroquine sulfate</i>	41	<i>insulin lispro junior kwikpen</i> ..	57	JENTADUETO	54
<i>hydroxyurea</i>	33	<i>insulin lispro prot & lispro</i>	57	JENTADUETO XR	54
<i>hydroxyzine hcl</i>	115	<i>insulin syringe</i>	57	JINTELI	92
<i>hydroxyzine pamoate</i>	52	INSULIN SYRINGE	58	JOLESSA	92

H4676_26DCF_C V14

Formulary ID 26314, Version 14

Last Updated 7/2026

JUBBONTI.....	111	<i>lamotrigine</i>	21	<i>levobunolol hcl</i>	114
JULEBER.....	92	<i>lamotrigine er</i>	21	<i>levocarnitine</i>	82
JULUCA.....	50	<i>lamotrigine starter kit-blue</i>	21	<i>levocarnitine sf</i>	82
JUNEL 1.5/30.....	92	<i>lamotrigine starter kit-green</i> ..	21	<i>levocetirizine dihydrochloride</i>	115
JUNEL 1/20.....	92	<i>lamotrigine starter kit-orange</i> ..	21	<i>levofloxacin</i>	20
JUNEL FE 1.5/30	92	<i>lansoprazole</i>	85	<i>levofloxacin in d5w</i>	20
JUNEL FE 1/20	92	<i>lanthanum carbonate</i>	82	LEVONEST	93
JUNEL FE 24	92	LANTUS	58	<i>levonorgest-eth estrad 91-day</i> ..	93
JYLAMVO.....	34	LANTUS SOLOSTAR.....	58	<i>levonorgestrel-ethinyl estrad</i> ..	93
JYNNEOS	108	<i>lapatinib ditosylate</i>	37	<i>levonorg-eth estrad triphasic</i> ..	93
K		LARIN 1.5/30.....	93	LEVO-T.....	97
KALETRA	51	LARIN 1/20.....	93	<i>levothyroxine sodium</i>	97
KALLIGA	92	LARIN 24 FE	93	LEVOXYL	97
KALYDECO.....	117	LARIN FE 1.5/30	93	<i>l-glutamine</i>	85
KARIVA	92	LARIN FE 1/20	93	<i>lidocaine</i>	12
<i>kcl in dextrose-nacl</i>	80	<i>latanoprost</i>	114	<i>lidocaine hcl</i>	13
KELNOR 1/35.....	92	LAZCLUZE	34	<i>lidocaine viscous hcl</i>	13
KERENDIA	67	LEDERLE LEUCOVORIN ...	40	<i>lidocaine-prilocaine</i>	13
KESIMPTA.....	75	<i>leflunomide</i>	106	LIFYORLI (125 MG DOSE) ..	34
<i>ketoconazole</i>	30	<i>lenalidomide</i>	33	LIFYORLI (150 MG DOSE) ..	34
<i>ketorolac tromethamine</i> ..	11, 113	LENVIMA (10 MG DAILY DOSE)	38	LILETTA (52 MG).....	93
KEVZARA.....	102	LENVIMA (12 MG DAILY DOSE)	38	<i>linezolid</i>	15
KINERET.....	102	LENVIMA (14 MG DAILY DOSE)	38	LINZESS	83
KINRIX.....	108	LENVIMA (18 MG DAILY DOSE)	38	<i>liothyronine sodium</i>	98
KISQALI (200 MG DOSE) ...	37	LENVIMA (20 MG DAILY DOSE)	38	<i>liraglutide</i>	54
KISQALI (400 MG DOSE) ...	37	LENVIMA (24 MG DAILY DOSE)	38	<i>lisinopril</i>	63
KISQALI (600 MG DOSE) ...	37	LENVIMA (4 MG DAILY DOSE)	38	<i>lisinopril-hydrochlorothiazide</i> ..	67
KLAYESTA.....	30	LENVIMA (8 MG DAILY DOSE)	38	LITFULO	102
KLOR-CON	81	LEQEMBI IQLIK	73	<i>lithium</i>	53
KLOR-CON 10	80	LEQSELVI.....	102	<i>lithium carbonate</i>	53
KLOR-CON M10.....	80	LESSINA.....	93	<i>lithium carbonate er</i>	53
KLOR-CON M15.....	80	<i>letrozole</i>	35	LIVMARLI.....	84
KLOR-CON M20.....	80	<i>leucovorin calcium</i>	40	LIVTENCITY	48
KLOXXADO	13	LEUKERAN	33	LODOCO	67
KOMZIFTI.....	34	LEUKINE.....	61	<i>lofexidine hcl</i>	13
KOSELUGO	37	<i>leuprolide acetate</i>	98	LOJAIMIESS	93
KRAZATI	34	<i>leuprolide acetate (3 month)</i> ..	98	LOKELMA.....	82
KURVELO.....	92	<i>levalbuterol hcl</i>	116	<i>lomustine</i>	33
KYLEENA.....	92	<i>levetiracetam</i>	22	LONSURF.....	34
KYMBEE.....	88	<i>levetiracetam er</i>	22	<i>loperamide hcl</i>	84
L				<i>lopinavir-ritonavir</i>	51
<i>labetalol hcl</i>	64			<i>lorazepam</i>	53
<i>lacosamide</i>	24			LORAZEPAM INTENSOL ...	53
<i>lactulose</i>	83			LORBRENA.....	38
<i>lactulose encephalopathy</i>	83			LORYNA	93
LAGEVRIO	52			<i>losartan potassium</i>	63
<i>lamivudine</i>	48			<i>losartan potassium-hctz</i>	67
<i>lamivudine-zidovudine</i>	50				

H4676_26DCF_C V14

Formulary ID 26314, Version 14

Last Updated 7/2026

<i>lovastatin</i>	69	MEKINIST	38	<i>micafungin sodium</i>	30
LOW-OGESTREL	93	MEKTOVI.....	38	MICROGESTIN 1.5/30.....	93
<i>loxapine succinate</i>	43	MELEYA	97	MICROGESTIN 1/20.....	93
LO-ZUMANDIMINE	93	<i>meloxicam</i>	11	MICROGESTIN FE 1.5/30	94
<i>lubiprostone</i>	83	<i>memantine hcl</i>	25	MICROGESTIN FE 1/20	94
LUIZZA 1.5/30	93	<i>memantine hcl er</i>	25	<i>midazolam</i>	23
LUIZZA 1/20	93	<i>memantine hcl-donepezil hcl er</i>	25	<i>midodrine hcl</i>	63
LUMAKRAS	34		25	<i>mifepristone</i>	56
LUMIGAN	114	MENQUADFI.....	108	<i>miglustat</i>	85
LUPKYNIS	106	MENVEO.....	108	MILI	94
LUPRON DEPOT (1-MONTH)	98	<i>mercaptopurine</i>	33, 34	<i>milnacipran hcl</i>	74
.....	98	<i>meropenem</i>	19	MIMVEY	94
LUPRON DEPOT (3-MONTH)	98	<i>meropenem-sodium chloride</i> ..	19	<i>minocycline hcl</i>	21
.....	98	<i>mesalamine</i>	110	<i>minoxidil</i>	70
LUPRON DEPOT (4-MONTH)	98	<i>mesalamine er</i>	110	MIRENA (52 MG)	94
.....	98	<i>mesna</i>	40	<i>mirtazapine</i>	26
LUPRON DEPOT (6-MONTH)	98	<i>metaxalone</i>	120	<i>misoprostol</i>	84
.....	98	<i>metformin hcl</i>	54	M-M-R II.....	108
<i>lurasidone hcl</i>	45	<i>metformin hcl er</i>	54	<i>modafinil</i>	121
LUTERA	93	<i>methadone hcl</i>	11	MODEYSO	34
LUTRATE DEPOT.....	98	<i>methazolamide</i>	114	<i>moexipril hcl</i>	63
LYBALVI	45	<i>methenamine hippurate</i>	15	<i>molindone hcl</i>	43
LYLEQ.....	93	<i>methimazole</i>	99	<i>mometasone furoate</i>	78, 116
LYNPARZA.....	38	<i>methocarbamol</i>	120	MONO-LINYAH	94
LYSODREN.....	34	<i>methotrexate sodium</i>	106	<i>montelukast sodium</i>	119
LYTGOBI (12 MG DAILY	38	<i>methotrexate sodium (pf)</i>	106	<i>morphine sulfate</i>	12
DOSE)	38	<i>methoxsalen rapid</i>	79	<i>morphine sulfate (concentrate)</i>	12
LYTGOBI (16 MG DAILY	38	<i>methsuximide</i>	22		12
DOSE)	38	<i>methylphenidate hcl</i>	72	<i>morphine sulfate er</i>	11
LYTGOBI (20 MG DAILY	38	<i>methylphenidate hcl er</i>	72	MOUNJARO.....	54
DOSE)	38	<i>methylphenidate hcl er (cd)</i>	72	MOVANTIK	83
LYZA	96	<i>methylphenidate hcl er (la)</i>	72	<i>moxifloxacin hcl</i>	20, 113
M		<i>methylphenidate hcl er (osm)</i> .	72	<i>moxifloxacin hcl in nacl</i>	20
<i>magnesium sulfate</i>	81	<i>methylphenidate hcl er (xr)</i>	72	MRESVIA	108
<i>malathion</i>	79	<i>methylphenidate hcl er(diffus)</i> 72		<i>mupirocin</i>	80
<i>maraviroc</i>	50	<i>methylprednisolone</i>	88	<i>mycophenolate mofetil</i>	106
<i>marlissa</i>	93	<i>methylprednisolone acetate</i> ..	111	<i>mycophenolate sodium</i>	106
MARPLAN	26	<i>methyldosterone</i>	89	MYFEMBREE	99
MATULANE	33	<i>metoclopramide hcl</i>	28	MYQORZO.....	67
MAVYRET	48	<i>metolazone</i>	69	MYRBETRIQ.....	86
MAYZENT	75	<i>metoprolol succinate er</i>	64	N	
MAYZENT STARTER PACK	75	<i>metoprolol tartrate</i>	65	<i>nabumetone</i>	11
.....	75	<i>metoprolol-hydrochlorothiazide</i>	67	<i>nadolol</i>	65
<i>meclizine hcl</i>	28		67	<i>nafacillin sodium</i>	18
<i>meclofenamate sodium</i>	11	<i>metronidazole</i>	15, 80	<i>nalbuphine hcl</i>	10
<i>medroxyprogesterone acetate</i> 97		<i>metyrosine</i>	67	<i>naloxone hcl</i>	13, 14
<i>mefloquine hcl</i>	41	<i>mexiletine hcl</i>	64	<i>naltrexone hcl</i>	13
<i>megestrol acetate</i>	97	MIBELAS 24 FE.....	93	NAMZARIC.....	25

H4676_26DCF_C V14

Formulary ID 26314, Version 14

Last Updated 7/2026

<i>naproxen</i>	11	<i>norethin ace-eth estrad-fe</i>	94	NUTRILIPID.....	82
<i>naproxen dr</i>	11	<i>norethindrone</i>	97	NYAMYC	30
<i>naproxen sodium</i>	11	<i>norethindrone acetate</i>	97	NYLIA 1/35.....	94
<i>naratriptan hcl</i>	31	<i>norethindrone acet-ethinyl est</i> 94		NYLIA 7/7/7	94
NATACYN	113	<i>norethindrone-eth estradiol</i> ...	94	<i>nystatin</i>	30
<i>nateglinide</i>	55	<i>norgestimate-eth estradiol</i>	94	<i>nystatin-triamcinolone</i>	79
NAYZILAM.....	23	<i>norgestim-eth estrad triphasic</i> 94		NYSTOP.....	30
<i>nebivolol hcl</i>	65	NORLYROC	97	O	
NECON 0.5/35 (28)	94	NORPACE CR.....	64	OCTAGAM.....	100
<i>nefazodone hcl</i>	27	NORTREL 0.5/35 (28).....	94	<i>octreotide acetate</i>	99
<i>neomycin sulfate</i>	14	NORTREL 1/35 (21).....	94	ODEFSEY	50
<i>neomycin-polymyxin-dexameth</i>		NORTREL 1/35 (28).....	94	ODOMZO.....	38
.....	112	NORTREL 7/7/7	94	OFEV	118
<i>neomycin-polymyxin-gramicidin</i>		<i>nortriptyline hcl</i>	28	<i>ofloxacin</i>	20, 113, 115
.....	112	NORVIR.....	51	OGSIVEO.....	38
<i>neomycin-polymyxin-hc</i> 114, 115		NOVOLIN 70/30.....	58	OJEMDA.....	38, 39
NERLYNX.....	38	NOVOLIN 70/30 FLEXPEN .	58	OJJAARA.....	34
NEULASTA.....	61, 62	NOVOLIN 70/30 FLEXPEN		<i>olanzapine</i>	46
NEULASTA ONPRO	61	RELION	58	<i>olmesartan medoxomil</i>	63
NEUPRO.....	42	NOVOLIN 70/30 RELION ...	58	<i>olmesartan medoxomil-hctz</i> ...	67
<i>nevirapine</i>	49	NOVOLIN N.....	58	<i>olmesartan-amlodipine-hctz</i> ...	67
<i>nevirapine er</i>	49	NOVOLIN N FLEXPEN	58	<i>omega-3-acid ethyl esters</i>	70
NEXLETOL	67	NOVOLIN N FLEXPEN		<i>omeprazole</i>	85
NEXLIZET.....	67	RELION	58	OMNIPOD 5 DEXG7G6	
NEXPLANON	94	NOVOLIN N RELION	58	INTRO GEN 5.....	59
NGENLA	88	NOVOLIN R.....	58	OMNIPOD 5 DEXG7G6 PODS	
<i>niacin er (antihyperlipidemic)</i> 70		NOVOLIN R FLEXPEN.....	58	GEN 5.....	59
NICOTROL NS.....	14	NOVOLIN R FLEXPEN		OMNIPOD 5 G7 INTRO (GEN	
<i>nifedipine</i>	65	RELION	58	5).....	59
<i>nifedipine er</i>	65	NOVOLIN R RELION	58	OMNIPOD 5 G7 PODS (GEN	
<i>nifedipine er osmotic release</i> ..	65	NOVOLOG	59	5).....	59
NIKKI.....	94	NOVOLOG 70/30 FLEXPEN		OMNIPOD 5 LIBRE INTRO.59	
<i>nilotinib d-tartrate</i>	38	RELION	59	OMNIPOD 5 LIBRE PODS...59	
<i>nilotinib hcl</i>	38	NOVOLOG FLEXPEN.....	59	OMNIPOD DASH INTRO	
<i>nilutamide</i>	33	NOVOLOG FLEXPEN		(GEN 4)	59
<i>nimodipine</i>	65	RELION	59	OMNIPOD DASH PODS (GEN	
NINLARO.....	34	NOVOLOG MIX 70/30	59	4).....	59
<i>nintedanib esylate</i>	118	NOVOLOG MIX 70/30		OMNITROPE.....	88, 89
<i>nitazoxanide</i>	41	FLEXPEN	59	<i>ondansetron</i>	29
<i>nitisinone</i>	85	NOVOLOG MIX 70/30		<i>ondansetron hcl</i>	29
NITRO-BID	70	RELION	59	ONGENTYS.....	42
NITRO-DUR.....	70	NOVOLOG PENFILL	59	ONUREG	34
<i>nitrofurantoin macrocrystal</i> ...	15	NOVOLOG RELION.....	59	OPIPZA	46
<i>nitrofurantoin monohyd macro</i>		NUBEQA	33	OPSUMIT.....	118
.....	15	NUCALA	119, 120	OPVEE	14
<i>nitroglycerin</i>	70	NUEDEXTA	73	ORENCIA	102
NORA-BE.....	97	NUPLAZID.....	45	ORENCIA CLICKJECT	102
<i>norelgestromin-eth estradiol</i> ..	94	NURTEC	31	ORFADIN	86

H4676_26DCF_C V14

Formulary ID 26314, Version 14

Last Updated 7/2026

ORGOVYX.....	99
ORIAHNN	99
ORILISSA.....	99
ORKAMBI.....	117
ORLADEYO.....	100
orphenadrine citrate er	120
ORQUIDEA.....	97
ORSERDU	34
oseltamivir phosphate	51
OSENVELT	111
OTEZLA	79
OTEZLA XR.....	79
OTEZLA/OTEZLA XR INITIATION PK.....	79
oxacillin sodium in dextrose...	18
oxcarbazepine.....	24
oxcarbazepine er	24
OXERVATE	112
oxybutynin chloride	86
oxybutynin chloride er.....	86
oxycodone hcl	12
oxycodone-acetaminophen	12
OXYCONTIN	12
OZEMPIC (0.25 OR 0.5 MG/DOSE).....	55
OZEMPIC (1 MG/DOSE).....	55
OZEMPIC (2 MG/DOSE).....	55
P	
paliperidone er	46
PANRETIN	40
pantoprazole sodium	85
paricalcitol	112
paroxetine hcl	27
paroxetine hcl er.....	27
PAXLOVID (150/100).....	52
PAXLOVID (300/100 & 150/100)	52
PAXLOVID (300/100).....	52
pazopanib hcl	39
PEDIARIX	108
PEDVAX HIB.....	108
peg 3350-kcl-na bicarb-nacl ..	83
peg-3350/electrolytes	83
PEGASYS.....	104, 105
PEMAZYRE	39
pen needles	59
PEN NEEDLES.....	59
PENBRAYA	108

H4676_26DCF_C V14

penciclovir	80
penicillamine	81
penicillin g pot in dextrose	18
penicillin g sodium	19
penicillin v potassium	19
penmenvy	108
PENTACEL.....	109
pentamidine isethionate.....	41
pentazocine-naloxone hcl.....	12
pentoxifylline er	68
perampanel.....	22
perindopril erbumine.....	64
permethrin	79
perphenazine	28
perphenazine-amitriptyline	26
PERSERIS.....	46
phenelzine sulfate	26
phenobarbital	23
phenoxybenzamine hcl.....	63
PHENYTEK.....	24
phenytoin	24
PHENYTOIN INFATABS.....	24
phenytoin sodium extended	24
PHILITH	95
PIFELTRO	49
pilocarpine hcl.....	76, 114
pimecrolimus	78
pimozide.....	43
PIMTREA	95
pindolol.....	65
pioglitazone hcl	55
pioglitazone hcl-metformin hcl	55
piperacillin sod-tazobactam so	19
piperacillin-tazobactam-nacl .	19
PIQRAY (200 MG DAILY DOSE)	39
PIQRAY (250 MG DAILY DOSE)	39
PIQRAY (300 MG DAILY DOSE)	39
pirfenidone.....	118
piroxicam.....	11
PLENAMINE.....	82
pnv 27-ca/fe/fa	82
podofilox.....	79
polymyxin b sulfate	15

Formulary ID 26314, Version 14

Page 131

polymyxin b-trimethoprim	113
pomalidomide	33
PONVORY.....	75
PONVORY STARTER PACK	75
PORTIA-28	95
posaconazole	30
potassium chloride.....	81
potassium chloride crys er.....	81
potassium chloride er	81
potassium citrate er	81
pramipexole dihydrochloride .42	
pramipexole dihydrochloride er	42
prasugrel hcl.....	62
pravastatin sodium	69
praziquantel.....	41
prazosin hcl	63
prednisolone	88
prednisolone acetate.....	113
prednisolone sodium phosphate	88, 111, 113
prednisone	111
PREDNISONE INTENSOL.....	111
pregabalin.....	23
PREMARIN	89, 90
PREMPHASE.....	95
PREMPRO	95
prenatal.....	82
pretomanid.....	32
PREVALITE	70
PREVYMIS.....	48
PREZCOBIX.....	50
PREZISTA	51
PRIFTIN	32
primaquine phosphate	41
PRIMAXIN IV	15
primidone.....	23
PRIORIX	109
PRIVIGEN	100
probenecid	31
prochlorperazine	28
prochlorperazine maleate.....	28
PROCRT	62
progesterone	97
PROGRAF.....	106
PROLASTIN-C	86
promethazine hcl	28, 115

Last Updated 7/2026

<i>promethazine-phenylephrine</i>	120
PROMETHEGAN.....	28
<i>propafenone hcl</i>	64
<i>propranolol hcl</i>	65
<i>propranolol hcl er</i>	65
<i>propylthiouracil</i>	99
PROQUAD.....	109
<i>protriptyline hcl</i>	28
PULMOZYME.....	117
<i>pyrazinamide</i>	32
<i>pyridostigmine bromide</i>	32
<i>pyridostigmine bromide er</i>	32
<i>pyrimethamine</i>	41
PYRUKYND.....	62
PYRUKYND TAPER PACK	62
Q	
QINLOCK.....	39
QUADRACEL	109
<i>quetiapine fumarate</i>	46
<i>quetiapine fumarate er</i>	46
<i>quinapril hcl</i>	64
<i>quinapril-hydrochlorothiazide</i>	68
<i>quinidine gluconate er</i>	64
<i>quinidine sulfate</i>	64
<i>quinine sulfate</i>	41
QULIPTA.....	31
R	
RABAVERT	109
RADICAVA ORS.....	73
RADICAVA ORS STARTER	
KIT	73
RALDESY	27
<i>raloxifene hcl</i>	97
<i>ramelteon</i>	121
<i>ramipril</i>	64
<i>ranolazine er</i>	68
<i>rasagiline mesylate</i>	42
REBIF.....	75
REBIF REBIDOSE	75
REBIF REBIDOSE	
TITRATION PACK.....	75
REBIF TITRATION PACK..	75
RECLIPSEN.....	95
RECOMBIVAX HB	109
RECORLEV	99
RELENZA DISKHALER	52
RELGAABI.....	23
RELISTOR.....	83

H4676_26DCF_C V14

<i>repaglinide</i>	55
REPATHA.....	70
REPATHA SURECLICK	70
RETACRIT	62
RETEVMO.....	39
REVCIVI	86
REVLIMID	33
REVUFORJ.....	35
REXTOVY	14
REXULTI.....	46
REYATAZ	51
REZDIFFRA	98
REZENOPY	14
REZLIDHIA.....	35
REZUROCK	106
RHOPRESSA.....	114
<i>ribavirin</i>	48
<i>rifabutin</i>	32
<i>rifampin</i>	32
<i>rilpivirine hcl</i>	49
<i>riluzole</i>	73
<i>rimantadine hcl</i>	52
<i>risedronate sodium</i>	112
<i>risperidone</i>	46
<i>risperidone microspheres er</i> ...	46
<i>ritonavir</i>	51
<i>rivastigmine</i>	25
<i>rivastigmine tartrate</i>	25
<i>rizatriptan benzoate</i>	31
ROCKLATAN	114
<i>roflumilast</i>	117
ROMVIMZA.....	35
<i>ropinirole hcl</i>	42
<i>ropinirole hcl er</i>	42
<i>rosuvastatin calcium</i>	69
ROTARIX	109
ROTATEQ	109
ROZLYTREK	39
RUBRACA.....	39
<i>rufinamide</i>	24
RUKOBIA.....	50
RYBELSUS.....	55
RYDAPT	39
RYKINDO.....	46
RYLAZE	35
S	
SANTYL	79
<i>sapropterin dihydrochloride</i> ...	86

Formulary ID 26314, Version 14

Page 132

SAVELLA TITRATION PACK	
.....	74
SCSEMBLIX.....	39
<i>scopolamine</i>	29
SECUADO	47
SELARSDI.....	102
<i>selegiline hcl</i>	43
<i>selenium sulfide</i>	78
SELZENTRY	50
SEREVENT DISKUS	116
SEROSTIM	89
<i>sertraline hcl</i>	27
SETLAKIN.....	95
<i>sevelamer carbonate</i>	82
SHAROBEL	97
SHINGRIX.....	109
SIGNIFOR.....	99
SIKLOS	34
<i>sildenafil citrate</i>	118
SILIQ.....	102
<i>silver sulfadiazine</i>	79
SIMBRINZA	114
SIMLANDI (1 PEN)	106
SIMLANDI (1 SYRINGE)...	106
SIMLANDI (2 PEN)	106
SIMLANDI (2 SYRINGE)...	106
SIMLIYA	95
SIMPESSE	95
SIMPONI.....	106
<i>simvastatin</i>	69
<i>sirolimus</i>	106
SIRTURO	32
SKYLA.....	95
SKYTROFA.....	89
<i>sodium chloride</i>	79, 81
<i>sodium chloride (pf)</i>	81
<i>sodium fluoride</i>	81
<i>sodium oxybate</i>	121
<i>sodium phenylbutyrate</i>	86
<i>sodium polystyrene sulfonate</i> .	82
<i>sofosbuvir-velpatasvir</i>	48
<i>solifenacin succinate</i>	87
SOLQUA.....	59
SOLTAMOX.....	33
SOMAVERT	99
<i>sorafenib tosylate</i>	39
<i>sotalol hcl</i>	64
<i>sotalol hcl (af)</i>	64

Last Updated 7/2026

SOTYKTU	102
SPIRIVA RESPIMAT.....	116
<i>spironolactone</i>	68
<i>spironolactone-hctz</i>	68
SPRINTEC 28	95
SPRITAM.....	22
SPS (SODIUM POLYSTYRENE SULF) ...	82
STARJEMZA.....	102
STELARA	103
STEQEYMA	103
STIOLTO RESPIMAT	120
STIVARGA.....	39
STOBOCLO.....	112
<i>streptomycin sulfate</i>	14
STRIBILD.....	50
SUBVENITE.....	22
SUCRAID	86
<i>sucrafate</i>	84
<i>sulfacetamide sodium</i>	113
<i>sulfacetamide sodium (acne)</i> ..	20
<i>sulfacetamide-prednisolone</i> .	113
<i>sulfadiazine</i>	20
<i>sulfamethoxazole-trimethoprim</i>	15, 20
<i>sulfasalazine</i>	110
<i>sulindac</i>	11
<i>sumatriptan</i>	31
<i>sumatriptan succinate</i>	32
<i>sunitinib malate</i>	39
SUNLENCA.....	50, 51
SYEDA.....	95
SYMDEKO	117
SYMLINPEN 120.....	55
SYMLINPEN 60.....	55
SYMPAZAN.....	23
SYMTUZA.....	51
SYNAREL	99
SYNJARDY	55
SYNJARDY XR	55
SYNTHROID.....	98
T	
TABLOID	34
TABRECTA.....	39
<i>tacrolimus</i>	78, 107
<i>tacrolimus er</i>	107
<i>tadalafil</i>	87
<i>tadalafil (pah)</i>	118

H4676_26DCF_C V14

TADLIQ	118
TAFINLAR	39
TAGRISO	39
TALTZ	103
TALZENNA.....	39
<i>tamoxifen citrate</i>	33
<i>tamsulosin hcl</i>	87
TARINA 24 FE	95
TARINA FE 1/20 EQ.....	95
TARPEYO.....	99
TASCENSO ODT	75
<i>tasimelteon</i>	121
TAVNEOS	62
<i>tazarotene</i>	76
TAZICEF.....	17
TAZVERIK.....	39
TEFLARO	18
<i>telmisartan</i>	63
<i>telmisartan-amlodipine</i>	68
<i>telmisartan-hctz</i>	68
<i>temazepam</i>	121
TENIVAC	109
<i>tenofovir disoproxil fumarate</i> .	48
TEPEZZA.....	113
TEPMETKO.....	39
<i>terazosin hcl</i>	63
<i>terbinafine hcl</i>	30
<i>terbutaline sulfate</i>	116
<i>terconazole</i>	30
<i>teriflunomide</i>	75
<i>teriparatide</i>	112
<i>testosterone</i>	89
<i>testosterone cypionate</i>	89
<i>testosterone enanthate</i>	89
<i>tetrabenazine</i>	73
<i>tetracycline hcl</i>	21
THALOMID.....	33
<i>theophylline</i>	117
<i>theophylline er</i>	117
<i>thioridazine hcl</i>	43
<i>thiothixene</i>	43
<i>tiagabine hcl</i>	23
TIBSOVO.....	35
<i>ticagrelor</i>	62
TICOVAC	109
<i>tigecycline</i>	15
TILIA FE.....	95
<i>timolol maleate</i>	65, 114

Formulary ID 26314, Version 14

Page 133

<i>tinidazole</i>	16
<i>tiopronin</i>	87
<i>tiotropium bromide</i>	116
TIVICAY.....	49
TIVICAY PD.....	49
<i>tizanidine hcl</i>	48
TOBI PODHALER	117
<i>tobramycin</i>	113, 117
<i>tobramycin sulfate</i>	14
<i>tobramycin-dexamethasone</i> ..	113
<i>tofacitinib citrate</i>	103
<i>tofacitinib citrate er</i>	103
<i>tolterodine tartrate</i>	87
<i>tolterodine tartrate er</i>	87
<i>tolvaptan</i>	81
<i>tolvaptan (hyponatremia)</i>	81
<i>topiramate</i>	22
<i>toremifene citrate</i>	33
<i>torseamide</i>	68
TOUJEO MAX SOLOSTAR.....	60
TOUJEO SOLOSTAR	60
TRADJENTA	55
<i>tramadol hcl</i>	12
<i>tramadol-acetaminophen</i>	12
<i>trandolapril</i>	64
<i>tranexamic acid</i>	62
<i>tranylcypromine sulfate</i>	26
<i>travoprost (bak free)</i>	114
<i>trazodone hcl</i>	27
TRELEGY ELLIPTA.....	120
TRELSTAR MIXJECT	99
TREMFYA	103, 104
TREMFYA ONE-PRESS.....	103
TREMFYA PEN	103
TREMFYA-CD/UC INDUCTION.....	104
<i>tretinoin</i>	40, 76
<i>triamcinolone acetonide</i> ...	76, 78
<i>triamcinolone in absorbase</i> ...	78
<i>triamterene-hctz</i>	68
<i>trientine hcl</i>	81
TRI-ESTARYLLA	95
<i>trifluoperazine hcl</i>	43
<i>trifluridine</i>	49
<i>trihexyphenidyl hcl</i>	41
TRIJARDY XR	55
TRIKAFTA	117
TRI-LEGEST FE.....	95

Last Updated 7/2026

TRI-LINYAH.....	95	<i>valproic acid</i>	22	VOQUEZNA.....	84
TRI-LO-ESTARYLLA	95	<i>valsartan</i>	63	VOQUEZNA DUAL PAK.....	84
TRI-LO-MARZIA.....	95	<i>valsartan-hydrochlorothiazide</i>		VOQUEZNA TRIPLE PAK ..	84
TRI-LO-MILI.....	95		68	VORANIGO.....	35
TRI-LO-SPRINTEC.....	95	VALTOCO 10 MG DOSE.....	23	<i>voriconazole</i>	30
<i>trimethobenzamide hcl</i>	29	VALTOCO 15 MG DOSE.....	23	VOSEVI	48
<i>trimethoprim</i>	16	VALTOCO 20 MG DOSE.....	23	VOWST.....	84
TRI-MILI	96	VALTOCO 5 MG DOSE.....	23	VRAYLAR.....	47
<i>trimipramine maleate</i>	28	<i>vancomycin hcl</i>	16	VYFEMLA.....	96
<i>trinatal rx 1</i>	83	<i>vancomycin hcl in nacl</i>	16	VYLIBRA	96
TRINTELLIX.....	27	VANFLYTA	40	W	
TRI-SPRINTEC	96	VAQTA.....	110	<i>warfarin sodium</i>	60
TRIUMEQ.....	51	<i>varenicline tartrate</i>	14	WEGOVY	68
<i>triumeq pd</i>	51	<i>varenicline tartrate (starter)</i> ..	14	WELIREG	35
TRI-VYLIBRA	96	<i>varenicline tartrate(continue)</i> 14		WERA	96
TRI-VYLIBRA LO	96	VARIVAX.....	110	WINREVAIR	118
<i>tropium chloride</i>	87	VAXCHORA	110	WIXELA INHUB.....	120
<i>tropium chloride er</i>	87	VELIVET	96	WYMZYA FE	96
TRULANCE.....	83	VELTASSA.....	82	WYOST.....	112
TRULICITY.....	55	VEMLIDY.....	48	X	
TRUMENBA	109	VENCLEXTA	40	XALKORI.....	40
TRUQAP.....	39	VENCLEXTA STARTING		XARELTO	61
TUKYSA.....	39	PACK	40	XARELTO STARTER PACK	
TURALIO	40	<i>venlafaxine hcl</i>	27		61
TURQOZ.....	96	<i>venlafaxine hcl er</i>	27	XATMEP.....	35
TWINRIX.....	109	VENTOLIN HFA.....	116	XCOPRI	22
TYBOST	51	VEOZAH.....	73	XCOPRI (250 MG DAILY	
TYMLOS	112	<i>verapamil hcl</i>	66	DOSE)	22
TYPHIM VI.....	109	<i>verapamil hcl er</i>	66	XCOPRI (350 MG DAILY	
TYVASO DPI		VERQUVO	68	DOSE)	22
MAINTENANCE KIT.....	118	VERSACLOZ	47	XDEMVI.....	113
TYVASO DPI TITRATION		VERZENIO	40	XELJANZ.....	104
KIT	118	VESTURA.....	96	XELJANZ XR.....	104
TYZAVAN.....	16	VIENVA.....	96	XERMELO.....	84
U		<i>vigabatrin</i>	23	XIFAXAN	84
UBRELVY	31	VIGAFYDE.....	24	XIGDUO XR.....	55, 56
UNITHROID.....	98	VIJOICE.....	40	XOLAIR.....	120
UPTRAVI.....	118	<i>vilazodone hcl</i>	27	XOLREMDI.....	62
UPTRAVI TITRATION	118	VIMKUNYA.....	110	XOSPATA.....	40
<i>ursodiol</i>	84	<i>viorele</i>	96	XPOVIO (100 MG ONCE	
<i>ustekinumab</i>	104	VIRACEPT	51	WEEKLY).....	35
<i>ustekinumab-aauz</i>	104	VIREAD.....	48	XPOVIO (40 MG ONCE	
<i>ustekinumab-aekn</i>	104	VITRAKVI.....	40	WEEKLY).....	35
UZEDY	47	VIVITROL	13	XPOVIO (40 MG TWICE	
V		VIVOTIF	110	WEEKLY).....	35
<i>valacyclovir hcl</i>	49	VIZIMPRO.....	40	XPOVIO (60 MG ONCE	
VALCHLOR.....	33	VOLNEA.....	96	WEEKLY).....	35
<i>valganciclovir hcl</i>	48	VONJO.....	40		

H4676_26DCF_C V14

Formulary ID 26314, Version 14
Page 134

Last Updated 7/2026

XPOVIO (60 MG TWICE WEEKLY).....	35
XPOVIO (80 MG ONCE WEEKLY).....	35
XPOVIO (80 MG TWICE WEEKLY).....	35
XROMI.....	34
XTANDI.....	33
XTRENBO.....	112
XULANE.....	96
XYWAV.....	121
Y	
YESINTEK.....	104
YF-VAX.....	110
YONSA.....	33
YORVIPATH.....	112
YUTREPIA.....	118

Z	
ZAFEMY.....	96
<i>zaleplon</i>	121
ZARXIO.....	62
ZAVZPRET.....	31
ZEJULA.....	40
ZELBORAF.....	40
ZEMAIRA.....	86
ZENATANE.....	76
ZENPEP.....	86
ZEPBOUND.....	121
ZEPBOUND KWIKPEN.....	121
ZEPOSIA.....	75
ZEPOSIA 7-DAY STARTER PACK.....	75
ZEPOSIA STARTER KIT.....	75
<i>zidovudine</i>	50
ZILBRYSQ.....	104

<i>ziprasidone hcl</i>	47
<i>ziprasidone mesylate</i>	47
ZITHROMAX.....	20
ZOLINZA.....	35
<i>zolmitriptan</i>	32
<i>zolpidem tartrate</i>	121
<i>zolpidem tartrate er</i>	121
ZONISADE.....	25
<i>zonisamide</i>	25
ZOSYN.....	16
ZOVIA 1/35 (28).....	96
ZTALMY.....	24
ZTLIDO.....	13
ZUMANDIMINE.....	96
ZURNAI.....	13
ZURZUVAE.....	26
ZYDELIG.....	40
ZYKADIA.....	40

2026 Troy Medicare
2026 Prior Authorization Criteria
CURRENT AS OF 08/01/2026

ACITRETIN

Products Affected

- acitretin*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a dermatologist or an oncologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For prophylaxis of skin cancer in patients with previously treated skin cancers who have undergone an organ transplantation the request will be approved. For psoriasis: the patient has documented trial of, contraindication to, or medical reason for not using at least 2 of the treatment options listed: topical steroids, tazarotene, methotrexate, and cyclosporine.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

ACL INHIBITORS

Products Affected

- NEXLETOLE
- NEXLIZET

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	18 years or older
Prescriber Restrictions	Prescriber must be a cardiologist or specialist in the treatment of lipid disorders
Coverage Duration	New starts will be authorized for 4 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For new starts ALL of the following must be provided: 1) Documentation of baseline low density lipoprotein cholesterol (LDL-C), 2) Member has tried and failed a high-intensity statin (i.e. atorvastatin 40-80 mg, rosuvastatin 20-40 mg) at maximum tolerated dose for 3 months via claim history or chart notes OR documentation has been provided that the member is not able to tolerate a statin. In addition to the initial criteria above if the new start is for the diagnosis of hypercholesterolemia, the following are required: 1) Member has a diagnosis of heterozygous familial hypercholesterolemia (FH) OR hypercholesterolemia, 2) Member has tried ezetimibe at a maximum tolerated dose and LDL-C is not at goal, or documentation has been provided that the patient is not able to tolerate ezetimibe. In addition to the initial criteria above if the new start is for cardiovascular risk reduction, the following are required: 1) Member has established cardiovascular disease (documented history of coronary artery disease, symptomatic peripheral artery disease, and/or cerebrovascular atherosclerotic disease, 2) Member does not have established cardiovascular disease but is considered high risk (one of the following): Diabetes Mellitus (Type 1 or Type 2) in females over 65 years of age or males over 60 years of age OR a Reynolds Risk score greater than 30% or a SCORE Risk score greater than 7.5% over 10 years OR a coronary artery

PA Criteria	Criteria Details
	calcium score greater than 400 Agaston units at any time in the past. For continuation of therapy or reauthorization requests for all indications: Documentation provided that the member has obtained clinical benefit from medication (e.g., LDL-C lowering from baseline)
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ACTEMRA

Products Affected

- ACTEMRA ACTPEN
- ACTEMRA SUBCUTANEOUS
- AVTOZMA SUBCUTANEOUS

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For pJIA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For RA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For sJIA, Giant Cell Arteritis and Systemic Sclerosis-Associated Interstitial Lung Disease: Approve
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ADALIMUMAB

Products Affected

- *adalimumab-fkjp (2 pen)*
- *adalimumab-fkjp (2 syringe) subcutaneous prefilled syringe kit 20 mg/0.4ml, 40 mg/0.8ml*
- SIMLANDI (1 PEN) SUBCUTANEOUS AUTO-INJECTOR KIT 40 MG/0.4ML, 80 MG/0.8ML
- SIMLANDI (1 SYRINGE)
- SIMLANDI (2 PEN)
- SIMLANDI (2 SYRINGE) SUBCUTANEOUS PREFILLED SYRINGE KIT 20 MG/0.2ML, 40 MG/0.4ML

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For ankylosing spondylitis: Trial of, medical reason for not using, or contraindication to naproxen. For pJIA: Trial of, medical reason for not using, or contraindication to 1 of the following DMARDs: methotrexate or leflunomide. For RA: Trial of, medical reason for not using, or contraindication to 1 disease modifying antirheumatic drug (DMARD) (methotrexate, leflunomide, or sulfasalazine). For PsA, psoriasis, Hidradenitis Suppurativa, Crohn's Disease (CD), Ulcerative Colitis (UC) or Uveitis: approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

ADEMPAS

Products Affected

- ADEMPAS

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with Phosphodiesterase Inhibitors used for Pulmonary Hypertension or Other Soluble Guanylate Cyclase Stimulators.
Required Medical Information	Pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group 1)- Initial: [Note: documentation required] (1) PAH was confirmed by right heart catheterization AND (2) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg AND (3) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND (4) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood units. Continuation of therapy: patient has positive clinical response to treatment. Chronic thromboembolic pulmonary hypertension (CTEPH) (WHO Group 4) - Initial: [Note: documentation required] (1) Patient has persistent or recurrent CTEPH after pulmonary endarterectomy (PEA) OR (2) patient has inoperable CTEPH with the diagnosis confirmed by right heart catheterization AND by computed tomography (CT), magnetic resonance imaging (MRI), or pulmonary angiography. Continuation of therapy: patient has positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist or pulmonologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	No

ALPHA-1 PROTEINASE INHIBITORS

Products Affected

- ARALAST NP INTRAVENOUS SOLUTION RECONSTITUTED 1000 MG, 500 MG
- GLASSIA
- PROLASTIN-C INTRAVENOUS SOLUTION
- ZEMAIRA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of hereditary alpha1-antitrypsin deficiency as evident by pretreatment serum AAT levels below 11 micromol/L and progressive FEV1 or FVC decline demonstrating symptomatic lung disease.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	If the medication request is for Glassia or Aralast NP, the patient has a documented medical reason (such as trial, intolerance or contraindication) for not using Prolastin-C or Zemaira to treat their medical condition.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ALYFTREK

Products Affected

- ALYFTREK ORAL TABLET 10-50-125 MG, 4-20-50 MG

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with Trikafta, Kalydeco, Orkambi, or Symdeko. Patients with unknown CFTR gene mutations.
Required Medical Information	Documentation of CFTR gene that is responsive to vanzacaftor-tezacaftor-deutivacaftor treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a pulmonologist or a provider who specializes in treatment of CF.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Cystic Fibrosis (CF) - Initial: patient must have at least one F508del mutation in the CFTR gene or a mutation in the CFTR gene that is responsive to the requested medication. Continuation of therapy: approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

AMBRISENTAN

Products Affected

- ambrisentan*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group 1)- Initial: [Note: documentation required] (1) PAH was confirmed by right heart catheterization AND (2) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg AND (3) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND (4) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood units. Continuation of therapy: patient has positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist or pulmonologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

ANTINEOPLASTIC AGENTS

Products Affected

- *abiraterone acetate*
- ABIRTEGA
- AKEEGA
- ALECENSA
- ALUNBRIG
- AUGTYRO
- AVMAPKI FAKZYNJA CO-PACK
- AYVAKIT
- BALVERSA
- *bexarotene*
- BOSULIF
- *bosutinib*
- BRAFTOVI ORAL CAPSULE 75 MG
- BRUKINSA
- CABOMETYX
- CALQUENCE ORAL TABLET
- CAPRELSA
- COMETRIQ (100 MG DAILY DOSE)
ORAL KIT 80 & 20 MG
- COMETRIQ (140 MG DAILY DOSE)
ORAL KIT 3 X 20 MG & 80 MG
- COMETRIQ (60 MG DAILY DOSE)
- COPIKTRA
- COTELLIC
- DANZITEN
- *dasatinib*
- DAURISMO
- ENSACOVE
- ERIVEDGE
- ERLEADA
- *erlotinib hcl*
- EULEXIN
- *everolimus oral tablet 10 mg, 2.5 mg, 5 mg, 7.5 mg*
- *everolimus oral tablet soluble*
- FOTIVDA
- FRUZAQLA
- GAVRETO
- *gefitinib*
- GILOTTRIF
- GOMEKLI
- HERNEXEOS
- HYRNUO
- IBRANCE
- IBTROZI
- ICLUSIG
- IDHIFA
- *imatinib mesylate oral*
- *imkeldi*
- INLURIYO
- INLYTA
- INQOVI
- INREBIC
- ITOVEBI
- IWILFIN
- JAYPIRCA
- KISQALI (200 MG DOSE)
- KISQALI (400 MG DOSE)
- KISQALI (600 MG DOSE)
- KOMZIFTI
- KOSELUGO
- KRAZATI
- *lapatinib ditosylate*
- LAZCLUZE
- *lenalidomide*
- LENVIMA (10 MG DAILY DOSE)
- LENVIMA (12 MG DAILY DOSE)
- LENVIMA (14 MG DAILY DOSE)
- LENVIMA (18 MG DAILY DOSE)
- LENVIMA (20 MG DAILY DOSE)
- LENVIMA (24 MG DAILY DOSE)
- LENVIMA (4 MG DAILY DOSE)
- LENVIMA (8 MG DAILY DOSE)
- LEUKERAN
- LIFYORLI (125 MG DOSE)
- LIFYORLI (150 MG DOSE)
- LONSURF
- LORBRENA

- LUMAKRAS
- LYNPARZA ORAL TABLET
- LYTGOBI (12 MG DAILY DOSE)
- LYTGOBI (16 MG DAILY DOSE)
- LYTGOBI (20 MG DAILY DOSE)
- MEKINIST
- MEKTOVI
- *mercaptopurine oral suspension*
- MODEYSO
- NERLYNX
- *nilotinib d-tartrate*
- *nilotinib hcl*
- *nilutamide*
- NINLARO
- NUBEQA
- ODOMZO
- OGSIVEO ORAL TABLET 100 MG, 150 MG
- OJEMDA
- OJJAARA
- ONUREG
- ORGOVYX
- ORSERDU
- *pazopanib hcl*
- PEMAZYRE
- PIQRAY (200 MG DAILY DOSE)
- PIQRAY (250 MG DAILY DOSE)
- PIQRAY (300 MG DAILY DOSE)
- *pomalidomide*
- QINLOCK
- RETEVMO ORAL TABLET
- REVLIMID
- REVUFORJ
- REZLIDHIA
- ROMVIMZA
- ROZLYTREK
- RUBRACA
- RYDAPT
- SCEMBLIX
- SOLTAMOX
- *sorafenib tosylate*
- STIVARGA
- *sunitinib malate*
- TABLOID
- TABRECTA
- TAFINLAR
- TAGRISSO
- TALZENNA
- TAZVERIK
- TEPMETKO
- THALOMID ORAL CAPSULE 100 MG, 50 MG
- TIBSOVO
- *toremifene citrate*
- *tretinoin oral*
- TRUQAP ORAL TABLET 200 MG
- TRUQAP ORAL TABLET THERAPY PACK
- TUKYSA
- TURALIO ORAL CAPSULE 125 MG
- VANFLYTA
- VENCLEXTA
- VENCLEXTA STARTING PACK
- VERZENIO
- VITRAKVI
- VIZIMPRO
- VONJO
- VORANIGO
- WELIREG
- XALKORI
- XOSPATA
- XPOVIO (100 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 50 MG
- XPOVIO (40 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 10 MG, 40 MG
- XPOVIO (40 MG TWICE WEEKLY) ORAL TABLET THERAPY PACK 40 MG
- XPOVIO (60 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 60 MG
- XPOVIO (60 MG TWICE WEEKLY)
- XPOVIO (80 MG ONCE WEEKLY) ORAL TABLET THERAPY PACK 40 MG, 80 MG
- XPOVIO (80 MG TWICE WEEKLY)

- XTANDI
- YONSA
- ZEJULA ORAL TABLET
- ZELBORAF

- ZOLINZA
- ZYDELIG
- ZYKADIA ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an oncologist or specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

APOMORPHINE

Products Affected

- apomorphine hcl subcutaneous*

PA Criteria	Criteria Details
Exclusion Criteria	Concomitant use with serotonin 5-HT3 receptor antagonists.
Required Medical Information	Reviewer will verify available patient claim history to confirm patient is not using 5-HT3 receptor antagonists.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	If diagnosis is Parkinson's, the patient must have a documented trial of, contraindication to, or medical reason for not using two alternatives such as entacapone, tolcapone, rasagiline, selegiline, carbidopa/levodopa, bromocriptine, pramipexole or ropinirole.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

AQNEURSA

Products Affected

- AQNEURSA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For new starts: 1) The member has a documented diagnosis of Niemann-Pick disease type C (NPC) AND 2) Documentation of genetic testing identifying disease-causing alleles in NPC1 or NPC2 AND 3) Documentation of disease-related neurological symptoms (e.g., developmental delay/regression, ataxia, cataplexy, seizures, motor-function decline, tremors, dysphagia) For reauthorization: Documentation that member has had positive response to therapy (e.g., stabilization in neurological status, decrease in functional Scale for Assessment and Rating of Ataxia [fSARA] score).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

ARCALYST

Products Affected

- ARCALYST

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For deficiency of interleukin-1 receptor antagonist, documented trial of, contraindication to, or medical reason for not using Kineret. For continuation of therapy or reauthorization: Documentation has been provided that patient has clinically benefited from medication.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ARIKAYCE

Products Affected

- ARIKAYCE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Mycobacterium avium complex (MAC): (1) Documented diagnosis of MAC lung disease as verified by failure to achieve at least 2 negative sputum cultures following 6 consecutive months of a combination antibacterial drug regimen AND (2) Provider attestation that medication is being used as part of a combination antibacterial drug regimen.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a pulmonologist or an infectious disease specialist
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

ARISTADA

Products Affected

- ARISTADA INITIO 441 MG/1.6ML, 662 MG/2.4ML, 882
- ARISTADA INTRAMUSCULAR MG/3.2ML
- PREFILLED SYRINGE 1064 MG/3.9ML,

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	The member has a documented history of receiving oral aripiprazole without any clinically significant side effects.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial and failure of, contraindication, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using at least two of the following: Abilify Maintena, Abilify Asimtufii, or Risperidone Microsphere ER.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

AUVELITY

Products Affected

- AUVELITY
- AUVELITY TITRATION PACK

PA Criteria	Criteria Details
Exclusion Criteria	Seizure disorder
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using to two generic antidepressants.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

AZTREONAM LYSINE

Products Affected

- CAYSTON

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist, infectious disease specialist, or an expert in the treatment of cystic fibrosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

BENLYSTA

Products Affected

- BENLYSTA SUBCUTANEOUS

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a rheumatologist, nephrologist, or specialist in the treatment of autoimmune disorders.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For new starts for systemic lupus erythematosus (SLE): concurrent use of two of the following or medical reason for not using glucocorticoids, azathioprine, methotrexate, mycophenolate, or hydroxychloroquine, chloroquine, and cyclophosphamide. For continuation of therapy or reauthorization for SLE: documentation of clinical response to therapy (i.e. fewer flares that required steroid treatment, lower average daily oral prednisone dose, improved daily function either as measured through a validated functional scale or through improved daily performance documented at clinic visits, etc.) For new starts for lupus nephritis (LN): concurrent use of or medical reason for not using background immunosuppressive therapy regimen. For continuation of therapy or reauthorization for LN: Documentation of improvement in renal function (i.e. reduction in UPCR).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	No

BESREMI

Products Affected

- BESREMI

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a hematologist, oncologist, or specialist for submitted diagnosis.
Coverage Duration	The request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

BOSENTAN

Products Affected

- *bosentan oral tablet*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of pulmonary arterial hypertension (PAH) WHO Group I classification and PAH Functional Class.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or cardiologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

BRINSUPRI

Products Affected

- BRINSUPRI

PA Criteria	Criteria Details
Exclusion Criteria	Patients with bronchiectasis due to cystic fibrosis.
Required Medical Information	N/A
Age Restrictions	12 years of age and older.
Prescriber Restrictions	Prescribed by or in consultation with a pulmonologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For non-cystic fibrosis bronchiectasis - Initial: (1) patient has documented history of bronchiectasis diagnosed by chest computed tomography scan AND (2) prescriber confirmation that bronchiectasis is not primarily driven by chronic obstructive pulmonary disease or asthma. Continuation of therapy: patient has positive clinical response to treatment.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CAMZYOS

Products Affected

- CAMZYOS

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For all new starts, documentation of ALL of the following must be provided: 1) Diagnosis of symptomatic New York Heart Association (NYHA) class II or III obstructive hypertrophic cardiomyopathy (oHCM) AND 2) Patient has a left ventricular ejection fraction (LVEF) greater than or equal to 55% AND 3) Assessment of Valsalva left ventricular outflow tract (LVOT) gradient AND 4) Trial of, medical reason for not using or contraindication to BOTH of the following: Beta blockers (i.e. metoprolol, propranolol, atenolol) AND Non-dihydropyridine calcium channel blockers (i.e. verapamil, diltiazem) AND 5) Prescriber attests that patient is not using moderate to strong CYP2C19 or CYP3A4 inhibitors or inducers. For continuation of therapy or reauthorization, all of the following must be provided: 1) Documentation of clinical benefit as evidenced by an improvement from baseline in oHCM symptoms (i.e., improvement in fatigue, chest pain, shortness of breath, LVOT, peak oxygen consumption, etc.) OR improvement or no worsening of NYHA functional class AND 2) Member must also have a left ventricular ejection fraction (LVEF) greater than or equal to 50%.
Indications	All Medically-accepted Indications.

PA Criteria	Criteria Details
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

CARDAMYST

Products Affected

- CARDAMYST

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist or specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Documented history of acute symptomatic episodes of paroxysmal supraventricular tachycardia.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CARGLUMIC ACID

Products Affected

- carglumic acid oral tablet soluble*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CASPOFUNGIN

Products Affected

- *caspofungin acetate*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Documentation of a consultation with an infectious disease specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CERDELGA

Products Affected

- CERDELGA

PA Criteria	Criteria Details
Exclusion Criteria	Patients with undetermined CYP2D6 metabolizer status.
Required Medical Information	Patient's CYP2D6 metabolizer status, as determined by an FDA approved test. For reauthorization, documentation has been provided that patient has obtained clinical benefit from medication (e.g. increased platelet count, improvement in anemia, PFTs, improvement in radiographic scans, improved quality of life).
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a specialist in treatment of Gaucher's disease.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CGRP ANTAGONISTS

Products Affected

- AIMOVIG
- EMGALITY
- EMGALITY (300 MG DOSE)
- NURTEC
- QULIPTA
- UBRELVY
- ZAVZPRET

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For acute migraine new starts - for Ubrelvy and Nurtec requests, must have trial of, contraindication to or medical reason for not using a triptan. For Zavzpret requests, must have trial of, contraindication to or medical reason for not using Ubrelvy or Nurtec. For migraine prophylaxis new starts - at least 4 migraine days per month or one or more severe migraine attacks lasting for greater than 12 hours despite use of abortive therapy (e.g. triptans or NSAIDs). For Emgality requests for episodic cluster headache new starts - approve. For continuation of therapy or reauthorization - For acute migraine (Nurtec, Ubrelvy, Zavzpret), must show documentation of improvement in migraine symptoms (pain, photophobia, phonophobia). For migraine prevention (Nurtec, Emgality, Qulipta, Aimovig), must show documentation of improvement in migraine symptoms. For episodic cluster headache treatment, must show documentation of reduction in frequency of headaches.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

CHOLBAM

Products Affected

- CHOLBAM

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: Patient has documented diagnosis of either: 1) bile acid synthesis disorder due to a single enzyme defect or 2) peroxisomal disorders. For continuation of therapy or reauthorization: prescriber attests: 1) the patient has clinical improvement with therapy (i.e. liver function tests) AND 2) there is no evidence of biliary obstruction or cholestasis
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a hepatologist, gastroenterologist, or metabolic specialist
Coverage Duration	New starts will be authorized for 3 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CIBINQO

Products Affected

- CIBINQO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For atopic dermatitis: Trial of, contraindication to, or medical reason for not using Rinvoq or Dupixent.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

CIMZIA

Products Affected

- CIMZIA (1 SYRINGE)
- CIMZIA (2 SYRINGE)
- CIMZIA SUBCUTANEOUS KIT 2 X 200 MG
- CIMZIA-STARTER

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For ankylosing spondylitis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: Enbrel, an adalimumab product, Cosentyx, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For Crohns Disease: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an adalimumab product or an ustekinumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For non-radiographic axial spondyloarthritis: approve. For psoriasis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: Tremfya, an ustekinumab product, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For PsA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: Cosentyx, an ustekinumab product, Tremfya, Xeljanz, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For RA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation

PA Criteria	Criteria Details
	of therapy. Polyarticular juvenile idiopathic arthritis (pJIA) - Initial: trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product or Xeljanz or or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

CORLANOR

Products Affected

- CORLANOR ORAL SOLUTION
- ivabradine hcl*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	New starts for chronic heart failure must have all of the following: 1) LVEF of 35% or less 2) Sinus rhythm and have resting heart rate greater than or equal to 70 bpm. For pediatric patients with heart failure due to dilated cardiomyopathy: approve.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a cardiologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not receiving a beta blocker.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

CORTROPHIN

Products Affected

- CORTROPHIN
- CORTROPHIN GEL

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	New starts for MS exacerbation, rheumatic disorders, collagen diseases, dermatologic diseases, serum sickness, edematous state (e.g. nephrotic syndrome without uremia), and respiratory diseases: trial of, contraindication to, or medical reason for not using oral corticosteroids. New starts for ophthalmic disease: trial of, contraindication to, or medical reason for not using oral or ophthalmic corticosteroids. Continuation of therapy or reauthorization for MS exacerbation: documentation of symptom improvement and current use of a multiple sclerosis disease modifying agent for maintenance therapy. Continuation of therapy or reauthorization for all other conditions: documented evidence of response to treatment and symptom improvement.
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	MS exacerbation: 1 month. Other conditions: new start for 3 months and reauth end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

COSENTYX

Products Affected

- COSENTYX (300 MG DOSE)
- COSENTYX INTRAVENOUS
- COSENTYX SENSOREADY (300 MG)
- COSENTYX SENSOREADY PEN
- COSENTYX SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 150 MG/ML, 75 MG/0.5ML
- COSENTYX UNOREADY

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	1) Documented diagnosis of ankylosing spondylitis, non-radiographic axial spondyloarthritis, plaque psoriasis, psoriatic arthritis, enthesitis-related arthritis, rheumatoid arthritis or hidradenitis suppurativa. 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CRESEMBA

Products Affected

- CRESEMBA ORAL

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documented diagnosis of medically accepted indication.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an infectious disease specialist or oncologist
Coverage Duration	Request will be authorized for 3 months.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CYSTAGON

Products Affected

- CYSTAGON

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

CYSTARAN

Products Affected

- CYSTARAN

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of diagnosis for cystinosis with corneal cystine crystal accumulation.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an ophthalmologist or metabolic disease specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

DALFAMPRIDINE ER

Products Affected

- dalfampridine er*

PA Criteria	Criteria Details
Exclusion Criteria	History of seizure or moderate/severe renal impairment (CrCl less than or equal to 50 mL/min).
Required Medical Information	For new starts: 1) Attestation that creatinine clearance (CrCl) greater than 50 mL/min was confirmed prior to initiation of therapy, AND 2) Documentation has been provided that member is ambulatory (able to walk at least 25 feet) and has a documented walking impairment, AND 3) For multiple sclerosis, member is currently being treated with a disease modifying agent (e.g. immunomodulator, interferon, etc.) or has a medical reason why member is unable to use a disease modifying agent for their condition. For continuation of therapy or re-authorization requests: 1) Member must experience improvement in walking from baseline due to use of dalfampridine ER.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

DEFERASIROX

Products Affected

- *deferasirox*
- *deferasirox granules*

PA Criteria	Criteria Details
Exclusion Criteria	Creatinine clearance less than 40 mL/min or platelet counts less than 50,000/mm ³ .
Required Medical Information	For all indications: platelet count greater than or equal to 50,000/mm ³ (within 30 days). For chronic iron overload due to transfusions: serum ferritin concentration greater than 1000 mcg/L (lab result within 30 days). For chronic iron overload in non-transfusion-dependent thalassemia syndromes: serum ferritin concentration greater than 300 mcg/L (lab result within 30 days).
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For deferasirox granules oral packets, the member must have medical reason for not using deferasirox tablets or oral soluble tablets.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

DEFERIPRONE

Products Affected

- *deferiprone*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: 1) serum ferritin level above 1,000 mcg/L and absolute neutrophil count (ANC) greater than $1.5 \times 10^9/L$ within 30 days of request, and 2) Trial of, contraindication to, or medical reason for not using deferasirox tablets. For continuation of therapy or reauthorization, decrease in serum ferritin from baseline.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

DEFLAZACORT

Products Affected

- KYMBEE ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	1) Member has a documented diagnosis of Duchenne Muscular Dystrophy as evidenced by one of the following: a) documented mutation of dystrophin gene OR b) muscle biopsy indicating absence of the dystrophin protein AND 2) Patient has trial and failure with, contraindication to or medical reason for not taking prednisone.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a neurologist or specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

DIACOMIT

Products Affected

- DIACOMIT

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For members 2 years and older: Trial of, contraindication to, or medical reason for not using one generic anticonvulsant for appropriate indications. For members under 2 years old: Approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

DIHYDROERGOTAMINE NASAL

Products Affected

- dihydroergotamine mesylate nasal*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: Member has a diagnosis of migraine headaches with or without aura. Prescriber attestation that it will be used for the acute treatment of migraine. For continuation of therapy or reauthorization: Documentation or provider attestation of positive clinical response (e.g., improvement in pain, photophobia, phonophobia).
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Requests will be authorized for 12 weeks.
Other Criteria	Trial of, contraindication to, or medical reason (e.g. intolerance or hypersensitivity) for not using a triptan (e.g., rizatriptan, sumatriptan).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

DOPTELET

Products Affected

- DOPTELET
- DOPTELET SPRINKLE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts for chronic liver disease and chronic immune thrombocytopenia (chronic ITP): documented baseline platelet count of less than 50,000/mcL.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with hematologist, hepatologist or surgeon.
Coverage Duration	For thrombocytopenia with CLD getting procedure: 5 days. For all others: remainder of contract year
Other Criteria	For chronic ITP: trial of, contraindication to, or medical reason for not using a corticosteroid. For thrombocytopenia with chronic liver disease (CLD): approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

DOXEPIN CREAM

Products Affected

- doxepin hcl external*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 1 month.
Other Criteria	Trial of, contraindication to, or medical reason for not using a topical corticosteroid or topical calcineurin inhibitor.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

DUPIXENT

Products Affected

- DUPIXENT SUBCUTANEOUS SOLUTION AUTO-INJECTOR
- DUPIXENT SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 200 MG/1.14ML, 300 MG/2ML

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Chronic spontaneous urticaria (CSU) - Initial: patient has inadequate symptomatic relief despite trial of two weeks of a second-generation oral antihistamine (unless contraindicated). Continuation of therapy: patient has positive clinical response to treatment. Bullous pemphigoid - Initial: patient has had trial of, contraindication to, or medical reason for not using a topical corticosteroid, oral corticosteroid, or an immunosuppressive agent. Continuation of therapy: patient has positive clinical response to treatment. Allergic Fungal Rhinosinusitis: Initial (1) patient is 6 years of age and older with history of sino-nasal surgery AND (2) patient has a trial of, contraindication to, or medical reason for not using nasal corticosteroids. Continuation of therapy: patient has positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Initial therapy only: prescribed by or in consultation with specialist for submitted diagnosis.
Coverage Duration	PN: Initial 6 mo. Cont of therapy and all others initial: end of contract year.
Other Criteria	Atopic dermatitis (AD) in patients 6 months of age and older - Initial: (1) patient has diagnosis of moderate to severe AD, AND (2) has had trial of, contraindication to, or medical reason for not using either a topical corticosteroid or topical calcineurin inhibitor. Continuation of therapy: patient has positive clinical response to treatment. Asthma with eosinophilic phenotype - Initial: (1) patient has baseline blood eosinophil count greater than or equal to 150 cells per microliter, AND (2) asthma remains inadequately controlled despite current treatment with or medical reasons for not using BOTH (i) medium to high dose inhaled corticosteroid AND (ii) additional controller (i.e. leukotriene modifier, long acting beta-2-

PA Criteria	Criteria Details
	agonist, long acting muscarinic antagonist, and sustained released theophylline). Asthma, oral corticosteroid dependent - Initial: asthma remains inadequately controlled despite current treatment with or medical reasons for not using BOTH (i) high dose inhaled corticosteroid AND (ii) additional controller (i.e. leukotriene modifier, long acting beta-2-agonist, long acting muscarinic antagonist, and sustained released theophylline). Asthma with eosinophilic phenotype or oral corticosteroid dependent - Continuation of therapy: clinical improvement in asthma control (i.e. reduction in frequency and/or severity of exacerbations and symptoms OR reduction in daily maintenance oral corticosteroid dose). Chronic rhinosinusitis with nasal polyps (CRSwNP) - Initial: (1) Dupixent is used as add-on maintenance treatment, AND (2) patient has a trial of, contraindication to, or medical reason for not using nasal corticosteroids. Continuation of therapy: patient has positive clinical response to treatment. Prurigo nodularis (PN): attestation is provided confirming diagnosis. Eosinophilic esophagitis (EoE) - Initial: (1) diagnosis has been confirmed by esophageal biopsy AND (2) patient weighs at least 15 kilograms AND (3) patient experienced an inadequate treatment response, intolerance, or has a contraindication to a topical or oral corticosteroid (i.e. fluticasone propionate or budesonide). Continuation of therapy: patient has positive clinical response to treatment. Chronic Obstructive Pulmonary Disease (COPD) - Initial: (1) documented diagnosis of COPD with an eosinophilic phenotype, AND (2) Dupixent is used as an add-on maintenance treatment, AND (3) documentation that patient's COPD is inadequately controlled. Continuation of therapy: patient has positive clinical response to treatment.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

EGRIFTA

Products Affected

- EGRIFTA SV
- EGRIFTA WR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of active antiretroviral therapy for at least 8 weeks.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

EMSAM

Products Affected

- EMSAM

PA Criteria	Criteria Details
Exclusion Criteria	Concomitant use with SSRIs, SNRIs, clomipramine and imipramine, meperidine, tramadol, methadone, pentazocine, and propoxyphene, and the antitussive agent dextromethorphan or carbamazepine
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: Trial of, contraindication to, or medical reason for not using two generic antidepressants.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ENBREL

Products Affected

- ENBREL MINI
- ENBREL SUBCUTANEOUS SOLUTION 25 MG/0.5ML
- ENBREL SUBCUTANEOUS SOLUTION PREFILLED SYRINGE
- ENBREL SURECLICK SUBCUTANEOUS SOLUTION AUTO-INJECTOR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For RA: Trial of, medical reason for not using, or contraindication to 1 disease modifying antirheumatic drug (DMARD) (methotrexate, leflunomide, or sulfasalazine). For pJIA: Trial of, medical reason for not using, or contraindication to 1 of the following DMARDs: methotrexate or leflunomide. For PsA or psoriasis: approve. For ankylosing spondylitis: Trial of, medical reason for not using, or contraindication to naproxen. Continuation of therapy: patient has been receiving Enbrel for a minimum of 4 months and has positive clinical response to treatment.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

ENDARI

Products Affected

- *l-glutamine oral packet*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation that two or more painful sickle cell crises have occurred in the past 12 months.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a hematologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using hydroxyurea for at least three months.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ENTYVIO

Products Affected

- ENTYVIO PEN

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For ulcerative colitis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

EPIDIOLEX

Products Affected

- EPIDIOLEX

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using one generic anticonvulsant for appropriate indications.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

EPRONTIA

Products Affected

- *topiramate oral solution 25 mg/ml*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	The request will be authorized until the end of the contract year.
Other Criteria	Documented trial of, contraindication to, or medical reason for not using topiramate sprinkle capsule or topiramate oral tablet.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ERYTHROPOIETIN STIMULATING AGENTS

Products Affected

- ARANESP (ALBUMIN FREE) INJECTION SOLUTION 100 MCG/ML, 200 MCG/ML, 25 MCG/ML, 40 MCG/ML, 60 MCG/ML
- ARANESP (ALBUMIN FREE) INJECTION SOLUTION PREFILLED SYRINGE
- EPOGEN INJECTION SOLUTION 10000 UNIT/ML, 2000 UNIT/ML, 20000 UNIT/ML, 3000 UNIT/ML, 4000 UNIT/ML, 40000 UNIT/ML
- PROCRIT
- RETACRIT INJECTION SOLUTION 10000 UNIT/ML, 10000 UNIT/ML(1ML), 2000 UNIT/ML, 20000 UNIT/ML, 3000 UNIT/ML, 4000 UNIT/ML, 40000 UNIT/ML

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts for all indications: Hgb within compendia range for treatment of the requested medical condition. For continuation of therapy or re-authorization: Hgb must not exceed 10 g/dL (anemia related to cancer), 11 g/dL (anemia of CKD), 12 g/dL (zidovudine-related anemia in members with HIV and ribavirin-induced anemia), 13 g/dL (elective, noncardiac, nonvascular surgery needing red blood cell allogeneic transfusion reduction).
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 6 months.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	No

ERZOFRI

Products Affected

- ERZOFRI INTRAMUSCULAR MG/1.5ML, 351 MG/2.25ML, 39
SUSPENSION PREFILLED SYRINGE MG/0.25ML, 78 MG/0.5ML
117 MG/0.75ML, 156 MG/ML, 234

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	The member has a documented history of receiving oral paliperidone or oral risperidone without any clinically significant side effects.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For schizophrenia: Trial and failure of, contraindication, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using at least two of the following: Abilify Maintena, Abilify Asimtufii, or Risperidone Microsphere ER. For schizoaffective disorder: approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

EUCRISA

Products Affected

- EUCRISA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a dermatologist, immunologist or an allergist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For patients 2 years of age and older: Trial of, contraindication to, or medical reason for not using topical tacrolimus or pimecrolimus. For patients less than 2 years of age: approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

EVRYSDI

Products Affected

- EVRYSDI

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts, all of the following must be included: 1) Documentation of genetic testing confirming diagnosis AND 2) Documentation of baseline motor function or motor milestone achievement [e.g. CHOP Infant Test of Neuromuscular Disorders (CHOP-INTEND) or Hammersmith Infant Neurological Examination (HINE) for Type 1 or Hammersmith Functional Motor Scale Expanded Scores (HFMSE), or Total Motor Function Measure 32 (MFM-32) for Type II and Type III, or 6 minute walk test in subjects able to walk]. For continuation of therapy or reauthorization, documentation of clinical response has been submitted (e.g. improvement in motor function/motor milestone achievement scores using CHOP-INTEND or HFMSE, MFM-32, 6 minute walk test or HINE improvement in more categories of motor milestones than worsening).
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

FABHALTA

Products Affected

- FABHALTA

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with another complement inhibitor for the treatment of PNH (i.e. Empaveli, Soliris, or Ultomiris).
Required Medical Information	PNH - Initial: patient has documented diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) confirmed by (1) flow cytometry analysis confirming presence of PNH clones AND (2) patient has signs and symptoms of PNH (i.e. anemia, abdominal pain, dyspnea, kidney disease, pulmonary hypertension, hemolysis/hemoglobinuria, etc.). Continuation of therapy: patient has documented positive clinical response to treatment (i.e. decrease in LDH, increased or stabilization of hemoglobin levels, reduction in transfusions, increased reticulocyte count, etc.). Reduction of proteinuria in adults with immunoglobulin A (IgA) nephropathy - Initial: patient has documented diagnosis of IgA nephropathy AND IgA nephropathy at risk of rapid disease progression (i.e. clinical evidence of rapid disease progression generally a urine protein-to-creatinine ratio or UPCR greater or equal to 1.5g/g OR other clinically relevant tests). Continuation of therapy: patient has documented positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a hematologist, nephrologist or oncologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	No

FASENRA

Products Affected

- FASENRA
- FASENRA PEN

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	New starts for severe asthma with an eosinophilic phenotype: 1)Baseline blood eosinophil count greater than or equal to 150 cells per microliter AND 2) symptoms persist with at least 1 exacerbation in the last 12 months requiring additional treatment (e.g. oral systemic steroids) while on a high dose inhaled corticosteroid with an additional controller medication (ie. long-acting B2 agonist). Continuation of therapy or re-authorization for severe asthma with an eosinophilic phenotype: clinical benefit from use of the drug. New starts for eosinophilic granulomatosis with polyangiitis (EGPA)- Initial: patient has a documented history or the presence of an eosinophil count of more than 1000 cells per microliter or a blood eosinophil level of greater than 10 percent AND trial of, contraindication to, or medical reason for not using one of the following medications: cyclophosphamide or methotrexate. Continuation of therapy: patient has a beneficial response to treatment with the requested drug (i.e. a reduction in the frequency of relapses, decrease in the daily oral corticosteroid dose, or no active vasculitis).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

FILSPARI

Products Affected

- FILSPARI

PA Criteria	Criteria Details
Exclusion Criteria	Coadministration with renin-angiotensin-aldosterone system (RAAS) inhibitors, endothelin receptor antagonists, or aliskiren
Required Medical Information	For new starts: Documentation is provided that the member has diagnosis of primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression. Member has proteinuria. For continuation of therapy or reauthorization: Documentation of positive clinical response (ie. decrease in urine protein-to-creatinine ratio (UPCR)).
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a nephrologist.
Coverage Duration	New starts will be authorized for 9 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

FINTEPLA

Products Affected

- FINTEPLA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using one generic anticonvulsant for appropriate indications.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

FIRDAPSE

Products Affected

- FIRDAPSE

PA Criteria	Criteria Details
Exclusion Criteria	History of seizures.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

FLUCYTOSINE

Products Affected

- *flucytosine oral*

PA Criteria	Criteria Details
Exclusion Criteria	Complete dihydropyrimidine dehydrogenase (DPD) enzyme deficiency
Required Medical Information	Attestation member is taking in combination with amphotericin B.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

GALAFOLD

Products Affected

- GALAFOLD

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed must be a geneticist, cardiologist, nephrologist or specialist experienced in the treatment of Fabry disease.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

GATTEX

Products Affected

- GATTEX

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: attestation of 1) Colonoscopy and upper gastrointestinal endoscopy with removal of polyps within six months prior to starting treatment for adults or 2) Fecal occult blood testing within six months prior to starting treatment for pediatric patients and 3) Baseline laboratory assessments (bilirubin, alkalinephosphatase, lipase, and amylase). For continuation of therapy or reauthorization: Documentation is provided that the member has obtained a clinical benefit (e.g. reduction in parenteral fluid volume, reduction in number of days receiving parenteral nutrition).
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a gastroenterologist.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

GLP-1 AGONISTS

Products Affected

- *liraglutide*
- MOUNJARO SUBCUTANEOUS SOLUTION AUTO-INJECTOR
- OZEMPIC (0.25 OR 0.5 MG/DOSE) SUBCUTANEOUS SOLUTION PEN-INJECTOR 2 MG/3ML
- OZEMPIC (1 MG/DOSE) SUBCUTANEOUS SOLUTION PEN-INJECTOR 4 MG/3ML
- OZEMPIC (2 MG/DOSE)
- RYBELSUS
- TRULICITY SUBCUTANEOUS SOLUTION AUTO-INJECTOR

PA Criteria	Criteria Details
Exclusion Criteria	The member has an indication of weight loss/obesity only or type 1 diabetes. The member has concurrent use of any GLP-1 receptor agonist.
Required Medical Information	The member has a documented diagnosis via chart notes of type 2 diabetes.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Diabetes: patient has diagnosis of type 2 diabetes mellitus. All other indications: patient must have medically accepted indication.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

GNRH AGONISTS

Products Affected

- ELIGARD
- FIRMAGON (240 MG DOSE)
- FIRMAGON SUBCUTANEOUS SOLUTION RECONSTITUTED 80 MG
- *leuprolide acetate (3 month)*
- *leuprolide acetate injection*
- LUPRON DEPOT (1-MONTH)
- LUPRON DEPOT (3-MONTH)
- LUPRON DEPOT (4-MONTH)
- LUPRON DEPOT (6-MONTH)
- LUTRATE DEPOT
- TRELSTAR MIXJECT

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Fibroids and endometriosis: 6 months. All other indications: end of contract year.
Other Criteria	If the medication request is for the treatment of prostate cancer and if the request is for any other GnRH agonist other than Eligard or leuprolide, the patient must have a documented trial of, contraindication to, or medical reason for not using Eligard or leuprolide to treat their prostate cancer.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

GOCOVRI

Products Affected

- GOCOVRI

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	New starts: trial of, contraindication to, or medical reason for not using generic amantadine. Continuation of therapy or reauthorization: Member demonstrates clinical benefit (i.e. improvement in levodopa-induced dyskinesia or decreased off episodes).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

GROWTH HORMONES

Products Affected

- GENOTROPIN MINIQUICK SUBCUTANEOUS PREFILLED SYRINGE
- GENOTROPIN SUBCUTANEOUS CARTRIDGE
- NGENLA
- OMNITROPE SUBCUTANEOUS SOLUTION CARTRIDGE
- OMNITROPE SUBCUTANEOUS SOLUTION RECONSTITUTED
- SKYTROFA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts for growth hormone deficiency: Documentation showing bone age testing, height, weight, and Growth Hormone Stimulation Test results OR Insulin Growth Factor 1 level. For continuation of therapy or reauthorization for growth hormone deficiency: documentation (medical records) showing positive response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an endocrinologist or nephrologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts for growth hormone deficiency: 1) If the request is not for Genotropin, trial of, contraindication to, or medical reason for not using Genotropin. For requests for all other medically accepted indications other than growth hormone deficiency, the request will be approved for products other than Skytrofa.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

HEREDITARY ANGIOEDEMA AGENTS

Products Affected

- CINRYZE
- HAEGARDA
- ORLADEYO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an allergist, immunologist, rheumatologist or hematologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For continuation of therapy or reauthorization: Documentation has been provided that patient has clinically benefited from medication.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

HIGH RISK MEDICATION

Products Affected

- *benztropine mesylate oral*
- *cyproheptadine hcl oral syrup 2 mg/5ml*
- *cyproheptadine hcl oral tablet*
- *diphenoxylate-atropine oral liquid*
- *diphenoxylate-atropine oral tablet 2.5-0.025 mg*
- *dipyridamole oral*
- *ergotamine-caffeine*
- *glyburide micronized oral tablet 3 mg, 6 mg*
- *glyburide oral tablet 1.25 mg, 2.5 mg, 5 mg*
- *glyburide-metformin oral tablet 1.25-250 mg*
- *glyburide-metformin oral tablet 2.5-500 mg, 5-500 mg*
- *guanfacine hcl er oral tablet extended release 24 hour 1 mg, 2 mg, 3 mg, 4 mg*
- *guanfacine hcl oral*
- *hydroxyzine hcl oral syrup 10 mg/5ml*
- *hydroxyzine hcl oral tablet*
- *indomethacin oral capsule 25 mg, 50 mg*
- *ketorolac tromethamine oral*
- *nifedipine oral*
- *promethazine hcl oral solution 6.25 mg/5ml*
- *promethazine hcl oral tablet*
- *promethazine hcl rectal suppository 12.5 mg, 25 mg*
- *promethazine-phenylephrine*
- **PROMETHEGAN RECTAL SUPPOSITORY 50 MG**

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For patients 65 years old and older the prescriber has documented: 1) the benefits of treatment with the drug outweigh the potential risks and 2) the risks and side effects have been discussed and will be monitored.
Age Restrictions	Prior authorization only applies to members 65 years old or older.
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	No

HIGH RISK MEDICATION - PROTECTED CLASS DRUGS

Products Affected

- *amitriptyline hcl oral*
- *amoxapine*
- *clomipramine hcl oral*
- *nortriptyline hcl oral*
- *perphenazine-amitriptyline*
- *phenobarbital oral elixir*
- *phenobarbital oral tablet*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For patients 65 years old and older the prescriber has documented: 1) the benefits of treatment with the drug outweigh the potential risks and 2) the risks and side effects have been discussed and will be monitored.
Age Restrictions	Prior authorization only applies to members 65 years old or older.
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

HIGH RISK MEDICATION, BUTALBITAL

Products Affected

- BAC (BUTALBITAL-ACETAMIN-CAFF)
- *butalbital-acetaminophen oral tablet 50-325 mg*
- *butalbital-apap-caff-cod oral capsule 50-325-40-30 mg*
- *butalbital-apap-caffeine oral capsule 50-325-40 mg*
- *butalbital-apap-caffeine oral solution*
- *butalbital-apap-caffeine oral tablet 50-325-40 mg*
- *butalbital-asa-caff-codeine*
- *butalbital-aspirin-caffeine oral capsule*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For patients 65 years old and older the prescriber has documented: 1) the benefits of treatment with the drug outweigh the potential risks and 2) the risks and side effects have been discussed and will be monitored.
Age Restrictions	Prior authorization only applies to members 65 years old or older.
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using an oral NSAID.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

HIGH RISK MEDICATION, MEGESTROL

Products Affected

- *megestrol acetate oral suspension 40 mg/ml, 400 mg/10ml, 625 mg/5ml*
- *megestrol acetate oral tablet*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For patients 65 years old and older the prescriber has documented: the benefits of treatment with the drug outweigh the potential risks identified for people 65 years old and older.
Age Restrictions	Prior authorization only applies to members 65 years old or older.
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

HIGH RISK MEDICATION, SHORT TERM MUSCLE RELAXANT

Products Affected

- *carisoprodol oral*
- *chlorzoxazone oral tablet 500 mg*
- *metaxalone oral tablet 800 mg*
- *methocarbamol oral tablet 500 mg, 750 mg*
- *orphenadrine citrate er*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For patients 65 years old and older the prescriber has documented: 1) the benefits of treatment with the drug outweigh the potential risks identified for people 65 years old and older, and 2) the risks and side effects have been discussed and will be monitored.
Age Restrictions	Prior authorization only applies to members 65 years old or older.
Prescriber Restrictions	N/A
Coverage Duration	New starts will be authorized for 30 days. Continuation of therapy or reauth will be for 90 days.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

HIGH RISK MEDICATION, SLEEP AGENTS

Products Affected

- *eszopiclone*
- *temazepam*
- *zaleplon oral capsule 10 mg, 5 mg*
- *zolpidem tartrate er*
- *zolpidem tartrate oral tablet 10 mg*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For patients 65 years old and older the prescriber has documented: 1) the benefits of treatment with the drug outweigh the potential risks identified for people 65 years old and older, and 2) the risks and side effects have been discussed and will be monitored. For zolpidem immediate release 10mg and zolpidem ER: trial of or medical reason for not using zolpidem immediate release 5mg.
Age Restrictions	Prior authorization only applies to members 65 years old or older.
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

HYFTOR

Products Affected

- HYFTOR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: documentation of diagnosis of tuberous sclerosis with facial angiofibroma. For continuation of therapy or reauthorization: documentation that the member has experienced a clinical benefit from treatment (e.g. improvement in size and color of angiofibroma).
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a neurologist or provider who specializes in the treatment of genetic or dermatologic disorders.
Coverage Duration	New starts: 3 months. Cont. of therapy or reauthorization: until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

ICATIBANT

Products Affected

- *icatibant acetate subcutaneous solution
prefilled syringe*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an immunologist, allergist, rheumatologist, or hematologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

ILARIS

Products Affected

- ILARIS SUBCUTANEOUS SOLUTION

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation was submitted indicating that the member was evaluated for active or latent TB infection (i.e. tuberculin skin test)
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For sJIA: approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

ILUMYA

Products Affected

- ILUMYA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For psoriasis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

IMBRUVICA

Products Affected

- IMBRUVICA ORAL CAPSULE
- IMBRUVICA ORAL SUSPENSION
- IMBRUVICA ORAL TABLET 140 MG, 280 MG, 420 MG

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an oncologist or specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts for treatment of graft-versus-host disease (GVHD): Trial of, contraindication to, or medical reason for not using a systemic corticosteroid. For continuation of therapy of for treatment of GVHD: documentation of clinical benefit from use of the drug (i.e. symptom improvement, reduction in corticosteroid dose). For all other indications, approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

IMPAVIDO

Products Affected

- IMPAVIDO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of diagnosis with one of the following: (a) Visceral leishmaniasis due to <i>Leishmania donovani</i> , (b) Cutaneous leishmaniasis due to <i>Leishmania braziliensis</i> , <i>Leishmania guyanensis</i> , or <i>Leishmania panamensis</i> , (c) Mucosal leishmaniasis due to <i>Leishmania braziliensis</i> .
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 28 days.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

INCRELEX

Products Affected

- INCRELEX

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an endocrinologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

JAKAFI

Products Affected

- JAKAFI
- JAKAFI XR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an oncologist or specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts for treatment of graft-versus-host disease (GVHD): Trial of, contraindication to, or medical reason for not using a systemic corticosteroid. For continuation of therapy of for treatment of GVHD: documentation of clinical benefit from use of the drug (i.e. symptom improvement, reduction in corticosteroid dose). For all other indications, approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

KALYDECO

Products Affected

- KALYDECO

PA Criteria	Criteria Details
Exclusion Criteria	Combination use with Orkambi, Symdeko, or Trikafta.
Required Medical Information	Documentation of CFTR gene that is responsive to ivacaftor treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or an expert in the treatment of cystic fibrosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

KERENDIA

Products Affected

- KERENDIA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: 1) Documentation of diagnosis of chronic kidney disease (CKD) due to type 2 diabetes mellitus OR heart failure (HF) with left ventricular ejection fraction (LVEF) greater than or equal to 40% AND AND 2) Documentation of serum potassium levels less than or equal to 5 mEq/L AND 3) eGFR greater than or equal to 25ml/min/1.73 m ² AND 4) Documentation that member is taking Kerendia in combination with an angiotensin-converting enzyme inhibitor (ACEi) or angiotensin receptor blocker (ARB) at maximum tolerated doses or documentation has been provided that the member is unable to tolerate ACEi or ARB. For continuation of therapy or reauthorization for patients with CKD due to type 2 diabetes mellitus: 1) Documentation of serum potassium levels less than or equal to 5.5 mEq/L AND 2) Documentation that member is taking Kerendia in combination with an ACEi or ARB at maximum tolerated doses or documentation has been provided that the member is unable to tolerate ACEi or ARB. For continuation of therapy or reauthorization for patients with HF with LVEF greater than or equal to 40%: 1) Documentation of serum potassium levels less than 6 mEq/L AND 2) Documentation that member is taking Kerendia in combination with an ACEi or ARB at maximum tolerated doses or documentation has been provided that the member is unable to tolerate ACEi or ARB.

PA Criteria	Criteria Details
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

KEVZARA

Products Affected

- KEVZARA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For RA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For polymyalgia rheumatica (PMR): Trial of, medical reason for not using, or contraindication to corticosteroids. For pJIA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

KINERET

Products Affected

- KINERET SUBCUTANEOUS
SOLUTION PREFILLED SYRINGE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For RA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For cryopyrin-associated periodic syndromes or deficiency of interleukin-1 receptor antagonist: Approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

LEQEMBI IQLIK

Products Affected

- LEQEMBI IQLIK

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist, geriatric psychiatrist, or geriatrician who specializes in treating dementia
Coverage Duration	Initial: 6 months. Continuation of therapy: end of the contract year.
Other Criteria	Initiation: (1) the patient has documentation of 18 months of treatment with intravenous Leqembi prior to initiation of subcutaneous maintenance therapy & (2) prior to initiation with intravenous Leqembi, the patient has an initial diagnosis of mild cognitive impairment (MCI) caused by Alzheimer's Disease (AD) or mild AD consistent with Stage 3 or Stage 4 Alzheimer's disease & (3) documentation of both of the following: a positive result for the presence of beta-amyloid plaques & baseline Magnetic Resonance Imaging (MRI) scan prior to initiation with intravenous Leqembi. Continuation of therapy: patient has a positive clinical response to treatment.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

H4676_26DCF_C V14 Formulary ID 26314, Version 14 Last Updated 7/2026
Page 240

Please refer to our Evidence of Coverage for more information about this coverage. You can find information on what the symbols and abbreviations on this table mean by going to Pages 7-8.

LEQSELVI

Products Affected

- LEQSELVI

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation was submitted indicating that the member was evaluated for active or latent TB infection (i.e. tuberculin skin test).
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Documentation of confirmed diagnosis and other causes of hair loss have been ruled out.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

LITFULO

Products Affected

- LITFULO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Initial: (1) documented diagnosis via chart notes of severe alopecia areata AND (2) patient is not receiving in combination with either of the following: (i) Targeted immunomodulator (i.e. Olumiant, Enbrel, Cimzia, Simponi, Orencia, adalimumab, Xeljanz, Rinvoq) OR (ii) potent immunosuppressant. Continuation of therapy: documentation of positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Documentation of confirmed diagnosis and other causes of hair loss have been ruled out.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

LIVMARLI

Products Affected

- LIVMARLI

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a gastroenterologist or hepatologist.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For new starts: 1) Trial of, contraindication to, or medical reason for not using both of the following: cholestyramine AND rifampin. 2) Prescriber attests that the member has cholestasis 3) Baseline serum bile acid level is provided. 4) Documentation of patients weight. For continuation of therapy or reauthorization: 1) Documentation submitted indicating the member has had all of the following: an improvement in pruritis (e.g. improved observed scratching, decreased sleep disturbances/nighttime awakenings due to scratching, etc.) AND reduction in serum bile acid level from baseline. 2) Prescriber attests that patient has had no evidence of hepatic decompensation (e.g. variceal hemorrhage, ascites, hepatic encephalopathy, portal hypertension, etc.). 3) Documentation of patients weight.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

LIVTENCITY

Products Affected

- LIVTENCITY

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Cytomegalovirus (CMV): (1) Documented diagnosis of CMV infection AND (2) Member is a recipient of one of the following: (a) hematopoietic stem cell transplant, (b) solid organ transplant AND (3) patient has tried and failed treatment with valganciclovir, ganciclovir, cidofovir, or foscarnet AND (4) patient weighs greater than or equal to 35 kg.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a transplant, oncologist, or infectious disease specialist.
Coverage Duration	Request will be authorized for 8 weeks.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

LODOCO

Products Affected

- LODOCO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be, or in consultation with a specialist in the treatment of cardiovascular disease, such as a cardiologist
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Documentation that patient has established atherosclerotic disease or multiple risk factors for cardiovascular disease AND documentation that patient does not have pre-existing blood dyscrasias (ex. leukopenia, thrombocytopenia) and patient does not have renal failure (CrCl less than 15 ml/min) or severe hepatic impairment
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

LUCEMYRA

Products Affected

- *lofexidine hcl*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 14 days.
Other Criteria	Patient must have trial of, contraindication to, or medical reason for not using clonidine.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

LUPKYNIS

Products Affected

- LUPKYNIS

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with cyclophosphamide.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be rheumatologist, nephrologist, or other specialist in the treatment of autoimmune disorders.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For new starts: 1) Documentation of urine protein/creatinine ratio (UPCR), 2) Documentation that the member has a baseline eGFR greater than 45 mL/min/1.73m ² or that benefit outweighs risk of using this medication at current eGFR, and 3) Concurrent use of or medical reason for not using background immunosuppressive therapy regimen. For continuation of therapy or reauthorization: Documentation of improvement in renal function (i.e. reduction in UPCR or no confirmed decrease from baseline eGFR greater than or equal to 20%).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

LYBALVI

Products Affected

- LYBALVI

PA Criteria	Criteria Details
Exclusion Criteria	Concomitant use with opioids.
Required Medical Information	Attestation from the provider that the member has had an opioid-free period of a minimum of 7 days after last use of shorting-acting opioids and 14 days from last use of long-acting opioids before initiating Lybalvi.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Documented trial of, contraindication to, or medical reason for not using at least two generic antipsychotics, one of which must be generic olanzapine.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

MAVYRET

Products Affected

- MAVYRET

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Detectable HCV RNA viral load prior to treatment within 6 months of request. In addition, documentation of treatment history, and if cirrhotic, documentation of compensated or decompensated cirrhosis.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 8-16 weeks as per AASLD-IDSA guidance.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

METHOTREXATE ORAL SOLUTION

Products Affected

- JYLAMVO
- XATMEP

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an oncologist, a rheumatologist, a dermatologist, or other appropriate specialist
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Polyarticular Juvenile Idiopathic Arthritis (pJIA) - Initial: (1) patient had been diagnosed with pJIA AND (2) patient had tried, intolerant or has medical reason for not using at least one non-steroidal anti-inflammatory agents (NSAIDs) AND methotrexate.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

METHYLTESTOSTERONE

Products Affected

- *methyltestosterone oral*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

METYROSIONE

Products Affected

- *metyrosine*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of one of the following: 1) Concurrent use of alpha adrenergic blockers, 2) Medical reason for being unable to use an alpha adrenergic blocker, OR 3) Patient is not a candidate for surgical resection and requires long term treatment with metyrosine.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

MIFEPRISTONE

Products Affected

- *mifepristone oral tablet 300 mg*

PA Criteria	Criteria Details
Exclusion Criteria	For all members patient must not be currently on simvastatin, lovastatin, cyclosporine, dihydroergotamine, ergotamine, fentanyl, pimozide, quinidine, sirolimus, and tacrolimus.
Required Medical Information	Reviewer will verify available claim history to confirm member is not taking simvastatin, lovastatin, cyclosporine, dihydroergotamine, ergotamine, fentanyl, pimozide, quinidine, sirolimus or tacrolimus concurrently with mifepristone.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an endocrinologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

MIGLUSTAT

Products Affected

- *miglustat*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts, documentation of diagnosis for mild to moderate type 1 Gaucher disease. For continuation of therapy or reauthorization: documentation of clinical benefit from use of the drug (i.e. increased platelet count, improvement in anemia, PFT's, improvement in radiographic scans, improved quality of life).
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a specialist in treatment of Gaucher's disease
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

MULTIPLE SCLEROSIS AGENTS

Products Affected

- BAFIERTAM
- BETASERON SUBCUTANEOUS KIT
- *cladribine (10 tabs)*
- *cladribine (4 tabs)*
- *cladribine (5 tabs)*
- *cladribine (6 tabs)*
- *cladribine (7 tabs)*
- *cladribine (8 tabs)*
- *cladribine (9 tabs)*
- *cladribine 10 mg tablet therapy pack oral*
- *dimethyl fumarate oral capsule delayed release 120 mg, 240 mg*
- *dimethyl fumarate starter pack oral capsule delayed release therapy pack*
- *fingolimod hcl*
- *glatiramer acetate subcutaneous solution prefilled syringe 20 mg/ml, 40 mg/ml*
- GLATOPA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 20 MG/ML, 40 MG/ML
- KESIMPTA
- MAYZENT
- MAYZENT STARTER PACK
- PONVORY
- PONVORY STARTER PACK
- REBIF REBIDOSE SUBCUTANEOUS SOLUTION AUTO-INJECTOR
- REBIF REBIDOSE TITRATION PACK SUBCUTANEOUS SOLUTION AUTO-INJECTOR
- REBIF SUBCUTANEOUS SOLUTION PREFILLED SYRINGE
- REBIF TITRATION PACK SUBCUTANEOUS SOLUTION PREFILLED SYRINGE
- TASCENSO ODT
- *teriflunomide*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	If the medication request is for glatiramer, Glatopa, or dimethyl fumarate, the request will be approved. If the member is over 17 years of age and the

PA Criteria	Criteria Details
	request is not for glatiramer, Glatopa, or dimethyl fumarate for multiple sclerosis, the member must have a documented trial of, contraindication to or a medical reason for not using 2 of the following dimethyl fumarate, glatiramer, Glatopa, teriflunomide, or fingolimod. If the request is for fingolimod and the member is 17 years of age or younger, the request will be approved.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

MYFEMBREE

Products Affected

- MYFEMBREE

PA Criteria	Criteria Details
Exclusion Criteria	Patient has history of osteoporosis or hepatic impairment.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an OB, gynecologist or reproductive endocrinologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts for menorrhagia: Trial of, contraindication to, or medical reason for not using an estrogen-progestin contraceptive therapy. For new starts if one of the following drugs has been tried previously, a trial of estrogen-progestin contraceptive therapy is not required: gonadotropin-releasing hormone (GnRH) agonists or tranexamic acid. New starts for endometriosis: Trial of, contraindication to, or medical reason for not using the following concurrently for endometriosis: analgesic pain reliever (e.g. NSAIDs, COX-2 inhibitors) AND either combined estrogen-progestin oral contraceptive, progestin (e.g. medroxyprogesterone acetate, norethindrone), gonadotropin-releasing hormone (GnRH) agonists (e.g. Lupron Depot), OR danazol. For continuation of therapy or reauthorization both of the following are required: 1) Treatment does not exceed the eligible maximum lifetime treatment duration of 2 years, and 2) Documentation has been provided that the member has obtained clinical benefit from medication (e.g. reduced menstrual bleeding from baseline, pain relief).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

MYQORZO

Products Affected

- MYQORZO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist.
Coverage Duration	Initial: 6 months. Continuation of therapy: end of contract year.
Other Criteria	For all new starts, documentation of ALL of the following must be provided: 1) Diagnosis of symptomatic obstructive hypertrophic cardiomyopathy (oHCM) AND 2) Patient has a left ventricular ejection fraction (LVEF) greater than or equal to 55% AND 3) Assessment of Valsalva left ventricular outflow tract (LVOT) gradient AND 4) Trial of, medical reason for not using or contraindication to BOTH of the following: Beta blockers (i.e. metoprolol, propranolol, atenolol) AND Non-dihydropyridine calcium channel blockers (i.e. verapamil, diltiazem). For continuation of therapy or reauthorization, all of the following must be provided: 1) Provider attests that the member has demonstrated a clinically meaningful benefit from therapy AND 2) Member must also have a left ventricular ejection fraction (LVEF) greater than or equal to 40%.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

NASAL ANTISEIZURE AGENTS

Products Affected

- NAYZILAM
- VALTOCO 10 MG DOSE
- VALTOCO 15 MG DOSE NASAL LIQUID THERAPY PACK 2 X 7.5 MG/0.1ML
- VALTOCO 20 MG DOSE NASAL LIQUID THERAPY PACK 2 X 10 MG/0.1ML
- VALTOCO 5 MG DOSE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	Nayzilam: 12 years of age or older. Valtoco: 2 years of age or older.
Prescriber Restrictions	Initial therapy only: prescribed by or in consultation with a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

NITISINONE

Products Affected

- nitisinone*
- ORFADIN ORAL SUSPENSION

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a geneticist, metabolic specialist, hepatologist, or liver transplant specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Hereditary Tyrosinemia Type 1 (HT-1): diagnosis was confirmed by genetic testing confirming a mutation of the FAH gene OR elevated levels of succinylacetone.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

NON-AMPHETAMINE CENTRAL NERVOUS SYSTEM AGENTS

Products Affected

- armodafinil*
- modafinil oral*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

NUCALA

Products Affected

- NUCALA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	New starts for severe asthma: 1) Baseline blood eosinophil count greater than or equal to 150 cells per microliter AND 2) symptoms with equal to or greater than 1 exacerbations in the previous 12 months while on a high-dose inhaled corticosteroid AND 3) the patient has a documented trial of, contraindication to, or medical reason for not using both Dupixent and Fasenra. New starts for eosinophilic granulomatosis with polyangiitis (EGPA): 1) trial of, contraindication to, or medical reason for not using one of the following medications: cyclophosphamide or methotrexate AND 2) the patient has a documented trial of, contraindication to, or medical reason for not using Fasenra. New starts for hypereosinophilic syndrome without an identifiable non-hematologic secondary cause: 1) 2 or more flares within the past 12 months AND 2) trial of, contraindication to, or medical reason for not using oral corticosteroids. New starts for chronic rhinosinusitis with nasal polyps: 1) the patient has a documented trial of, contraindication to, or medical reason for not using both Dupixent and Xolair. Continuation of therapy or re-authorization for all indications: clinical benefit from use of the drug. COPD - initial: (1) documented diagnosis of COPD AND (2) Documentation of blood eosinophil count of greater than or equal to 300 cells/microliter AND (3) Nucala is used as add-on maintenance treatment AND (4) patient has had a trial of, contraindication to, or medical reason

PA Criteria	Criteria Details
	for not using at least two of the following: a) long acting beta2-agonist (LABA), b) long acting muscarinic antagonist (LAMA) or c) inhaled corticosteroid (ICS) AND (5) documentation of COPD exacerbation within the past 12 months. Continuation of therapy: patient has positive clinical response to treatment.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

NUEDEXTA

Products Affected

- NUEDEXTA

PA Criteria	Criteria Details
Exclusion Criteria	Complete atrioventricular (AV) block without implanted pacemaker, or at high risk of complete AV block. History of heart failure. Concomitant use with MAOIs or use of MAOIs within 14 days. Concomitant use with drugs containing quinidine, quinine, or mefloquine. History of quinine-, mefloquine-, dextromethorphan/quinidine-, or quinidine-induced thrombocytopenia, hepatitis, bone marrow depression, or lupus-like syndrome. Non-Part D indications.
Required Medical Information	Confirmation diagnosis is for Part D indication.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a neurologist or psychiatrist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

NUPLAZID

Products Affected

- NUPLAZID ORAL CAPSULE
- NUPLAZID ORAL TABLET 10 MG

PA Criteria	Criteria Details
Exclusion Criteria	Patient has a history of dementia-related psychosis.
Required Medical Information	For hallucinations and delusions associated with Parkinson's disease psychosis: documented diagnosis of Parkinson's disease must be made prior to the onset of psychotic symptoms.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

OCTREOTIDE

Products Affected

- *octreotide acetate injection solution 100 mcg/ml, 1000 mcg/ml, 200 mcg/ml, 50 mcg/ml, 500 mcg/ml*
- *octreotide acetate intramuscular*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts for acromegaly: pt meets one of the following (1) inadequate response to surgery and/or radiotherapy OR (2) pt is not an appropriate candidate for surgery and/or radiotherapy OR (3) pt is experiencing negative effects due to tumor size (ex: optic nerve compression). Continuation of therapy or reauthorization: documentation of clinical improvement with therapy.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Continuation of therapy or reauthorization: documentation of clinical improvement with therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

OFEV

Products Affected

- nintedanib esylate*
- OFEV

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist, rheumatologist, or lung transplant specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For a diagnosis of idiopathic pulmonary fibrosis: 1) Documentation of disease as demonstrated on a high resolution CT scan or through lung biopsy and 2) Documented trial of, contraindication to, or medical reason for not using pirfenidone. For a diagnosis of systemic sclerosis-associated interstitial lung disease (SSc-ILD): documented trial of, contraindication to, or medical reason for not using mycophenolate mofetil or cyclophosphamide. For a diagnosis of chronic fibrosing interstitial lung diseases (ILDs) with a progressive phenotype: documentation is provided confirming diagnosis.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

OPSUMIT

Products Affected

- OPSUMIT

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Pulmonary arterial hypertension (PAH) World Health Organization (WHO) Group I - Initial: [Note: documentation required] (1) PAH was confirmed by right heart catheterization AND (2) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg AND (3) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg AND (4) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood units. Continuation of therapy: patient has positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or cardiologist
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using sildenafil.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ORAL ANTIPSYCHOTICS

Products Affected

- BYSANTI
- BYSANTI TITRATION PACK A
- BYSANTI TITRATION PACK B
- BYSANTI TITRATION PACK C
- CAPLYTA
- COBENFY
- COBENFY STARTER PACK
- FANAPT
- FANAPT TITRATION PACK A
- FANAPT TITRATION PACK B ORAL TABLET
- FANAPT TITRATION PACK C ORAL TABLET
- OPIPZA ORAL FILM 10 MG, 2 MG, 5 MG
- VRAYLAR ORAL CAPSULE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Schizophrenia: 1) request is for Caplyta, Cobenfy, Fanapt, Opienza, Bysanti, or Vraylar AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics. Manic or mixed episodes associated with bipolar I disorder: 1) request is for Fanapt, Opienza, Bysanti, or Vraylar AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics. Major depressive disorder associated with bipolar I or II disorder: 1) request is for Caplyta AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics. Adjunctive treatment of major depressive disorder: 1) request is for Caplyta, Opienza or Vraylar AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics AND 3) if the request is for Vraylar: provider attestation that the member is concurrently using an antidepressant. Irritability associated with autistic disorder: 1) request is for

PA Criteria	Criteria Details
	Opipza AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics. Treatment of Tourette's disorder: 1) request is for Opipza AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics. Bipolar I disorder: 1) request is for Opipza AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics. Depressed bipolar I disorder: 1) request is for Vraylar AND 2) trial of, contraindication to, or medical reason for not using two generic antipsychotics.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ORENCIA

Products Affected

- ORENCIA CLICKJECT
- ORENCIA INTRAVENOUS
- ORENCIA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 125 MG/ML, 50 MG/0.4ML, 87.5 MG/0.7ML

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For pJIA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For PsA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, Xeljanz, Enbrel, Cosentyx, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For RA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For acute graft versus host disease: Attestation member is taking in combination with a calcineurin inhibitor and methotrexate.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ORIAHNN

Products Affected

- ORIAHNN

PA Criteria	Criteria Details
Exclusion Criteria	Patient has history of osteoporosis or hepatic impairment.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an OB, gynecologist or reproductive endocrinologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: Trial of, contraindication to, or medical reason for not using an estrogen-progestin contraceptive therapy. For new starts if one of the following drugs has been tried previously, a trial of estrogen-progestin contraceptive therapy is not required: gonadotropin-releasing hormone (GnRH) agonists or tranexamic acid. For continuation of therapy or reauthorization both of the following are required: 1) Treatment does not exceed the eligible maximum lifetime treatment duration of 2 years, and 2) Documentation has been provided that the member has obtained clinical benefit from medication (e.g. reduced menstrual bleeding from baseline).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ORILISSA

Products Affected

- ORILISSA

PA Criteria	Criteria Details
Exclusion Criteria	Patient has osteoporosis or severe hepatic impairment.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an OB or gynecologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using the following concurrently for endometriosis: analgesic pain reliever (e.g. NSAIDs, COX-2 inhibitors) AND either combined estrogen-progestin oral contraceptive, progestin (e.g. medroxyprogesterone acetate, norethindrone), gonadotropin-releasing hormone (GnRH) agonists (e.g. Lupron Depot), OR danazol. For continuation of therapy or reauthorization both of the following are required: 1) Treatment does not exceed the eligible maximum lifetime treatment duration of 2 years for 150mg tablet or 6 months for 200mg tablet, and 2) Documentation has been provided that the member has obtained clinical benefit from the medication.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

H4676_26DCF_C V14 Formulary ID 26314, Version 14 Last Updated 7/2026
Page 277

Please refer to our Evidence of Coverage for more information about this coverage. You can find information on what the symbols and abbreviations on this table mean by going to Pages 7-8.

ORKAMBI

Products Affected

- ORKAMBI

PA Criteria	Criteria Details
Exclusion Criteria	Combination use with Kalydeco, Symdeko, or Trikafta.
Required Medical Information	Documentation of CFTR gene that is responsive to lumacaftor-ivacaftor treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or an expert in the treatment of cystic fibrosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

OTEZLA

Products Affected

- OTEZLA
- OTEZLA XR
- OTEZLA/OTEZLA XR INITIATION PK

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For moderate to severe psoriasis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For PsA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, Xeljanz, Enbrel, Cosentyx, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For Behcet's Syndrome or mild psoriasis: Approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

OXERVATE

Products Affected

- OXERVATE

PA Criteria	Criteria Details
Exclusion Criteria	Treatment duration greater than 16 weeks per affected eye(s).
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by an ophthalmologist or an optometrist.
Coverage Duration	Initial: 8 weeks. Continuation of therapy: additional 8 weeks.
Other Criteria	Initial: confirmed diagnosis. Continuation of therapy: patient has previously received less than or equal to 8 weeks of treatment per affected eye(s) and the patient has a recurrence of neurotrophic keratitis.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

OXYCODONE ER

Products Affected

- OXYCONTIN ORAL TABLET ER 12 MG, 20 MG, 30 MG, 40 MG, 60 MG, 80 MG
HOUR ABUSE-DETERRENT 10 MG, 15 MG

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Members being treated for active cancer diagnoses, sickle cell diagnoses, those in hospice care, or receiving palliative care will be excluded from the concurrent benzodiazepine and muscle relaxant therapy requirement. For new starts, ALL of the following are required: (1) Member has documented history of receiving an immediate-release opioid, (2) Member has a documented trial of, contraindication to, or medical reason for not using long-acting morphine sulfate, (3) If member is on concurrent benzodiazepines and/or muscle relaxant therapy, the prescriber attests that concurrent therapy is medically necessary, (4) Member is not being treated for substance abuse with buprenorphine-containing products. For continuing therapy, ALL of the following are required: (1) Member's pain has been assessed within the last 6 months, (2) Member has demonstrated clinical improvement in pain and function on current medication regimen, (3) If member is on concurrent benzodiazepines and/or muscle relaxant therapy, the prescriber attests that concurrent therapy is medically necessary, (4) Member is not being treated for substance abuse with buprenorphine-containing products.
Indications	All Medically-accepted Indications.

PA Criteria	Criteria Details
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PALIPERIDONE INJECTABLE

Products Affected

- INVEGA HAFYERA 156 MG/ML, 234 MG/1.5ML, 39 MG/0.25ML, 78 MG/0.5ML
- INTRAMUSCULAR SUSPENSION
- PREFILLED SYRINGE 1092 MG/3.5ML, 1560 MG/5ML
- INVEGA TRINZA INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 273 MG/0.88ML, 410 MG/1.32ML, 546 MG/1.75ML, 819 MG/2.63ML
- INVEGA SUSTENNA
- INTRAMUSCULAR SUSPENSION
- PREFILLED SYRINGE 117 MG/0.75ML,

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	The member has a documented history of receiving oral risperidone or oral paliperidone without any clinically significant side effects. For requests for Invega Trinza, the member has documented treatment with Invega Sustenna for at least 4 months. For requests for Invega Hafyera, the member has documented treatment with Invega Sustenna for at least 4 months OR the member has documented treatment with Invega Trinza for at least one three-month cycle.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For requests for Invega Sustenna, Invega Trinza, or Invega Hafyera for schizophrenia: Trial and failure of, contraindication, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using at least two of the following: Abilify Maintena, Abilify Asimtufii, or Risperidone Microsphere ER. For requests for Invega Sustenna for schizoaffective disorder: approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PCSK9 INHIBITORS

Products Affected

- REPATHA
- REPATHA SURECLICK

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For all diagnoses (besides for risk reduction of major cardiovascular events): documented diagnosis of medically accepted indication. For risk reduction of major cardiovascular (CV) events (CV death, myocardial infarction, stroke, unstable angina requiring hospitalization, or coronary revascularization): 1) Patient has an increased risk of major CV events such as death from CV disease, heart attack, stroke, certain types of chest pain conditions (including but not limited to, unstable angina) requiring hospitalization or certain types of heart surgery and (2) tried or has contraindication to high intensity statin.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PEGINTERFERON

Products Affected

- PEGASYS SUBCUTANEOUS SOLUTION 180 MCG/ML
- PEGASYS SUBCUTANEOUS SOLUTION PREFILLED SYRINGE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For Hepatitis C: 1) Labs within 3 months of request: liver function tests and detectable HCV RNA viral load. 2) Documentation of genotype, treatment history, and if cirrhotic, documentation of compensated or decompensated cirrhosis. For Hepatitis B: 1) Labs within 3 months of request: ALT/AST, and 2) HBeAg status. For polycythemia vera, approve.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a gastroenterologist, hepatologist, infectious disease doctor or transplant specialist.
Coverage Duration	Request will be authorized for 24 to 48 weeks as defined by compendia.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

PENICILLAMINE

Products Affected

- *penicillamine oral tablet*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For RA: Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz. For other indications, approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PERSERIS

Products Affected

- PERSERIS

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	The member has a documented history of receiving oral risperidone without any clinically significant side effects.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial and failure of, contraindication, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using at least two of the following: Abilify Maintena, Abilify Asimtufii, or Risperidone Microsphere ER.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PHENOXYBENZAMINE

Products Affected

- *phenoxybenzamine hcl oral*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using doxazosin.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PIRFENIDONE

Products Affected

- *pirfenidone*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For idiopathic pulmonary fibrosis, documentation of all of the following: 1) confirmation of diagnosis on high resolution CT scan or through lung biopsy AND 2) FVC greater than or equal to 50% of the predicted value.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or lung transplant specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

POSACONAZOLE

Products Affected

- *posaconazole oral*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Documentation of a consultation with an infectious disease specialist, a transplant specialist, or an oncologist.
Coverage Duration	28 days for oropharyngeal candidiasis, end of contract year for other indications
Other Criteria	For treatment of oropharyngeal candidiasis: trial of, contraindication to, or medical reason for not using fluconazole or itraconazole. For prophylaxis of invasive aspergillus infections due to being severely immunocompromised: trial of, contraindication to, or medical reason for not using voriconazole.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PRETOMANID

Products Affected

- pretomanid*

PA Criteria	Criteria Details
Exclusion Criteria	MDR-TB that is not treatment-intolerant or nonresponsive to standard therapy
Required Medical Information	Documentation of use in combination with bedaquiline and linezolid.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an infectious disease specialist.
Coverage Duration	Request will be authorized for 26 weeks.
Other Criteria	Documentation of prior trial of or medical reason for not using first-line TB regimen containing isoniazid and rifampin.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PREVYMIS

Products Affected

- PREVYMIS ORAL

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a hematologist, oncologist, infectious disease, or transplant specialist.
Coverage Duration	Request will be authorized for 6 months.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

PROMACTA

Products Affected

- *eltrombopag olamine oral packet 12.5 mg, 25 mg*
- *eltrombopag olamine oral tablet 12.5 mg, 25 mg, 50 mg, 75 mg*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For persistent or chronic immune thrombocytopenia (ITP): Documented baseline platelet count less than 30,000 cells/ microL. For severe aplastic anemia: Documentation of baseline platelet count less than 20,000 cells/microL OR platelet count less than 30,000 cells/microL with bleeding OR reticulocyte count less than 20,000 cells/microL OR absolute neutrophil count less than 500 cells/microL. For thrombocytopenia in patients with Hepatitis C infection: documented baseline platelet count less than 75,000 cells/microL.
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For persistent or chronic immune thrombocytopenia (ITP): Trial of, contraindication to, or medical reason for not using glucocorticosteroids. For severe aplastic anemia: Trial of, contraindication to, or medical reason for not using at least one immunosuppressive agent.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

PYRUKYND

Products Affected

- PYRUKYND
- PYRUKYND TAPER PACK

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: 1) documentation of diagnosis and 2) baseline hemoglobin level. For continuation of therapy or reauthorization: documentation of clinical improvement (e.g. reduction in number of blood transfusions, or increase or stabilization in hemoglobin level). If the criteria are not met, may authorize up to 14 days of a Pyrukynd Taper Pack to allow for tapering.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a hematologist.
Coverage Duration	New starts: 6 mo. Cont of therapy or reauth: end of contract yr. Denial: 14 days for dose tapering.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

RADICAVA

Products Affected

- RADICAVA ORS
- RADICAVA ORS STARTER KIT

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	New starts: 6 months. Cont. of therapy or reauthorization: until end of contract year.
Other Criteria	For new starts: 1) documentation of ALS functional rating scale (ALSFRS-R) score and 2) documentation that the member has been on riluzole, is beginning therapy as an adjunct to treatment with Radicava, or provider has provided a medical reason why patient is unable to use riluzole. For continuation of therapy or reauthorization: documentation from provider of clinical stabilization in symptoms (e.g. stabilization of ALS functional rating scale (ALSFRS-R) score).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

RAVICTI

Products Affected

- *glycerol phenylbutyrate*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Provider is a geneticist, metabolic specialist, gastroenterologist, hepatologist, or liver transplant specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using sodium phenylbutyrate.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

RECORLEV

Products Affected

- RECORLEV

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using ketoconazole tablets.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

RELISTOR

Products Affected

- RELISTOR ORAL
- RELISTOR SUBCUTANEOUS SOLUTION
- RELISTOR SUBCUTANEOUS SOLUTION PREFILLED SYRINGE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For opioid induced constipation, chronic non-cancer pain: Patient must have documented trial of or medical reason for not using at least two the following: 1) lubiprostone, OR 2) lactulose OR 3) Movantik. Additionally, patient must have a medical reason for not being able to use oral Relistor in order to receive Relistor injection. For opioid-induced constipation, in patients with advanced illness or pain caused by active cancer requiring opioid dosage escalation for palliative care: Approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

REVCovi

Products Affected

- REVCovi

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Patient has diagnosis of adenosine deaminase severe combined immune deficiency (ADA-SCID) confirmed by one of the following: 1) documentation of deficiency or absence of adenosine deaminase OR 2) genetic testing revealing mutations in both alleles of the ADA1 gene
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an immunologist, geneticist, hematologist, oncologist, or provider who specializes in the treatment of ADA-SCID or related disorders.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

REXULTI

Products Affected

- REXULTI

PA Criteria	Criteria Details
Exclusion Criteria	Dementia-related psychosis
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For schizophrenia: trial of, contraindication to, or medical reason for not using two generic antipsychotics. For major depressive disorder: trial of, contraindication to, or medical reason for not using two generic antidepressants. For agitation due to dementia: 1) patient has documented diagnosis of Alzheimer's disease AND 2) medication will not be used on an "as needed" basis
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

REZDIFFRA

Products Affected

- REZDIFFRA

PA Criteria	Criteria Details
Exclusion Criteria	Patients with decompensated cirrhosis.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a hepatologist, gastroenterologist, endocrinologist or specialist in the treatment of liver disease.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Initial therapy: (1) documented diagnosis of noncirrhotic nonalcoholic steatohepatitis (NASH) with moderate to advanced liver fibrosis, (2) documentation of stage F2 to F3 fibrosis as confirmed by a biopsy or a non-invasive test (NIT), (3) the drug is being prescribed at an FDA approved dose according to the member's weight and (4) prescriber attestation to providing lifestyle counseling on nutrition, exercise and avoiding excessive alcohol intake. For reauthorization: (1) the member has shown clinical benefit from the medication (e.g., the resolution of steatohepatitis and no worsening of liver fibrosis, or at least one stage improvement in liver fibrosis and no worsening of steatohepatitis), (2) the member continues to have a fibrosis stage of 3 or less and (3) the drug continues to be prescribed at an FDA approved dose according to the member's weight.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	No

REZUROCK

Products Affected

- REZUROCK

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a hematologist, oncologist, or transplant specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: documented trial of, contraindication to, or medical reason for not using at least two lines of systemic immunosuppressive therapy (e.g. corticosteroids, tacrolimus, mycophenolate mofetil, Imbruvica, or Jakafi), one of which must be a systemic corticosteroid. For continuation of therapy or re-authorization: documentation of clinical benefit from use of the drug (i.e. symptom improvement, reduction in corticosteroid dose).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

RUFINAMIDE

Products Affected

- *rufinamide oral suspension 40 mg/ml*
- *rufinamide oral tablet*

PA Criteria	Criteria Details
Exclusion Criteria	History of familial Short QT syndrome
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using one alternative generic anticonvulsant for appropriate indications.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

RYKINDO

Products Affected

- RYKINDO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	The member has a documented history of receiving oral risperidone without any clinically significant side effects.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial and failure of, contraindication, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using at least two of the following: Abilify Maintena, Abilify Asimtufii, or Risperidone Microsphere ER.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

RYLAZE

Products Affected

- RYLAZE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an oncologist, hematologist, or specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SAPROPTERIN

Products Affected

- *sapropterin dihydrochloride oral packet*
- *sapropterin dihydrochloride oral tablet*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: documentation of elevated baseline phenylalanine levels. Continuation of therapy or reauthorization: prescriber attests the member has improvement in phenylalanine levels from baseline.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	New starts will be authorized for 3 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SECUADO

Products Affected

- SECUADO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using to one generic antipsychotics.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SEROSTIM

Products Affected

- SEROSTIM SUBCUTANEOUS
SOLUTION RECONSTITUTED 4 MG, 5
MG, 6 MG

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a HIV specialist, gastroenterologist, nutritional support specialist or ID specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For initial starts for HIV wasting/cachexia: 1) Member must be on anti-retroviral therapy and 2) Alternative causes of wasting have been ruled out (diarrhea, malignancies, inadequate caloric intake, etc)
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SIGNIFOR

Products Affected

- SIGNIFOR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Member is not a candidate for surgery or surgery was not curative.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an endocrinologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SILDENAFIL ORAL

Products Affected

- *sildenafil citrate oral suspension reconstituted*
- *sildenafil citrate oral tablet 20 mg*

PA Criteria	Criteria Details
Exclusion Criteria	Documentation of concurrent nitrate or Adempas use.
Required Medical Information	Pulmonary Arterial Hypertension: Documentation of pulmonary arterial hypertension (PAH) WHO Group I and PAH Functional Class. Reviewer will verify available patient claim history to confirm patient is not using nitrates or Adempas. Secondary Raynaud's phenomenon- Initial: [Note: documentation required] (1) diagnosis of secondary Raynaud's phenomenon AND (2) diagnosis of primary condition which Raynaud's phenomenon is secondary to (e.g., lupus, scleroderma, rheumatoid arthritis, Sjogren's syndrome, thyroid disease). Continuation of therapy: patient has positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist, cardiologist, or rheumatologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For sildenafil suspension: Documentation of trial of, contraindication to, or medical reason for not using sildenafil tablet.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SILIQ

Products Affected

- SILIQ

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For psoriasis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SIMPONI

Products Affected

- SIMPONI SUBCUTANEOUS SOLUTION AUTO-INJECTOR
- SIMPONI SUBCUTANEOUS SOLUTION PREFILLED SYRINGE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For ankylosing spondylitis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: Enbrel, an adalimumab product, Cosentyx, or Xeljanz, or 2) If utilized within the past 120 days, approve for continuation of therapy. For PsA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Cosentyx, Tremfya, Xeljanz, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For RA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: Enbrel, an adalimumab product, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For adults with UC: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following: an adalimumab product, an ustekinumab product, Tremfya, or Xeljanz or 2) If utilized within the past 120 days, approve for continuation of therapy. For pediatric patients with UC: approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SIRTURO

Products Affected

- SIRTURO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation (consistent with pharmacy claims data, OR for new members to the health plan consistent with medical chart history) that the member is currently taking Sirturo as part of a combination regimen with other antimycobacterial drugs to treat MDR-TB.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an infectious disease specialist.
Coverage Duration	Request will be authorized for 24 weeks.
Other Criteria	Documentation of prior trial of or medical reason for not using first-line TB regimen containing isoniazid and rifampin.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SODIUM OXYBATE

Products Affected

- *sodium oxybate*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a sleep specialist, pulmonologist, or neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For somnolence associated with narcolepsy in adults: trial of, contraindication to, or medical reason for not using a CNS stimulant (e.g. methylphenidate, modafinil, armodafinil, etc.). For somnolence associated with narcolepsy in pediatric patients: approve. For cataplexy associated with narcolepsy, approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SODIUM PHENYLBUTYRATE

Products Affected

- *sodium phenylbutyrate oral powder 3 gm/tsp*
- *sodium phenylbutyrate oral tablet*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Provider is a geneticist, metabolic specialist, gastroenterologist, hepatologist, or liver transplant specialist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SOFOSBUVIR/VELPATASVIR

Products Affected

- *sofosbuvir-velpatasvir*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Detectable HCV RNA viral load prior to treatment within 6 months of request. In addition, documentation of treatment history, and if cirrhotic, documentation of compensated or decompensated cirrhosis.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 12-24 weeks based on AASLD-IDSA guidelines
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SOMAVERT

Products Affected

- SOMAVERT

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts for acromegaly: pt meets one of the following (1) inadequate response to surgery and/or radiotherapy OR (2) pt is not an appropriate candidate for surgery and/or radiotherapy OR (3) pt is experiencing negative effects due to tumor size (ex: optic nerve compression). Continuation of therapy or reauthorization: documentation of clinical improvement with therapy.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an endocrinologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SOTYKTU

Products Affected

- SOTYKTU

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For moderate to severe psoriasis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. Active Psoriatic Arthritis- Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Tremfya, Xeljanz, Enbrel, Cosentyx, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

SUCRAID

Products Affected

- SUCRAID

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: documentation of diagnosis of congenital sucrase-isomaltase deficiency. For continuation of therapy or reauthorization: Prescriber attests that member has obtained a clinical benefit (e.g. fewer total stools, greater number of hard and formed stools, fewer watery and soft stools, decrease in breath hydrogen output)
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SYMDEKO

Products Affected

- SYMDEKO

PA Criteria	Criteria Details
Exclusion Criteria	Combination use with Kalydeco, Orkambi, or Trikafta.
Required Medical Information	Documentation of CFTR gene that is responsive to tezacaftor-ivacaftor treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or an expert in the treatment of cystic fibrosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

SYNAREL

Products Affected

- SYNAREL

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using the following concurrently for endometriosis: analgesic pain reliever (e.g. NSAIDs, COX-2 inhibitors) AND either combined estrogen-progestin oral contraceptive, progestin (e.g. medroxyprogesterone acetate, norethindrone), OR gonadotropin-releasing hormone (GnRH) agonists (e.g. Lupron Depot)
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TADALAFIL

Products Affected

- *tadalafil (pah)*
- TADLIQ

PA Criteria	Criteria Details
Exclusion Criteria	Documentation of concurrent nitrate or Adempas use.
Required Medical Information	Documentation of pulmonary arterial hypertension (PAH) WHO Group I and PAH Functional Class. Reviewer will verify available patient claim history to confirm patient is not using nitrates or Adempas.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or cardiologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For Tadliq: Documentation of trial of, contraindication to, or medical reason for not using tadalafil tablets.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TADALAFIL, BPH

Products Affected

- *tadalafil oral tablet 5 mg*

PA Criteria	Criteria Details
Exclusion Criteria	Diagnosis of erectile dysfunction
Required Medical Information	Diagnosis of Benign prostatic hyperplasia (BPH) required AND trial of, contraindication to, or medical reason for not using an alpha blocker (e.g. tamsulosin, terazosin).
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TALTZ

Products Affected

- TALTZ SUBCUTANEOUS SOLUTION AUTO-INJECTOR
- TALTZ SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 20 MG/0.25ML, 40 MG/0.5ML, 80 MG/ML

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For ankylosing spondylitis: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: Enbrel, an adalimumab product, Cosentyx, or Xeljanz, or 2) If utilized within the past 120 days, approve for continuation of therapy. For non-radiographic axial spondyloarthritis: approve. For psoriasis: Either 1) Trial of, medical reason for not using, or contraindication (e.g., safety concerns, not indicated for patient's age) to 2 of the following therapies: an ustekinumab product, Tremfya, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy. For PsA: Either 1) Trial of, medical reason for not using, or contraindication to 2 of the following therapies: an ustekinumab product, Cosentyx, Tremfya, Xeljanz, Enbrel, or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TARPEYO

Products Affected

- TARPEYO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: documentation is required that the member has 1) Diagnosis of primary immunoglobulin A nephropathy (IgAN) and 2) at risk of disease progression. Member has proteinuria greater than or equal to 0.5 g/day. For continuation of therapy: documentation that member has been on Tarpeyo for less than 9 months. For reauthorizations: Requests will not be allowed as the safety and efficacy of subsequent courses of Tarpeyo have not been established.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a nephrologist.
Coverage Duration	Request will be authorized for 9 months.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TASIMELTEON

Products Affected

- HETLIOZ LQ
- tasimelteon*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts of non-24 hour sleep-wake cycle: 1) Member is totally blind with no perception of light, 2) diagnosis of non-24 confirmed by a physiologic circadian phase marker (ex: dim light melatonin onset, assessment of core body temp or measurement of urinary melatonin levels) OR actigraphy with evaluation of sleep logs. For continuation of therapy or reauthorization: documentation of clinical benefit from use of the drug. For night-time sleep disturbances in Smith-Magenis Syndrome (SMS): approve
Age Restrictions	N/A
Prescriber Restrictions	Provider is a sleep specialist or neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TAVNEOS

Products Affected

- TAVNEOS

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a rheumatologist or hematologist.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For new starts: 1) Prescriber attests that Tavneos will be prescribed in combination with corticosteroids AND cyclophosphamide unless there is documented trial of, contraindication to, or medical reason for not using these therapies. 2) Documentation of baseline Birmingham Vasculitis Activity Score (BVAS) score 3) Prescriber attestation that the patient will have liver function tests before treatment (ALT, AST, alkaline phosphate, and total bilirubin) and every 4 weeks after start of therapy for the first 6 months of treatment 4) Prescriber attestation that the patient has been screened for and does not have active hepatitis B virus (HBV) infection at baseline. For continuation of therapy or reauthorization: 1) Documentation of remission (BVAS score of 0) OR improvement in BVAS score 2) Prescriber attestation that patient has no abnormality in liver function tests (abnormality: ALT or AST greater than 3 times the upper limit of normal and bilirubin greater than 2 times the upper limit of normal) 3) Prescriber attestation that patient has no active HBV infection.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TEPEZZA

Products Affected

- TEPEZZA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of moderate to severe active thyroid eye disease (TED) as evidenced by one or more of the following: a) eyelid retraction greater than or equal to 2 mm, b) moderate or severe soft tissue involvement, c) exophthalmos greater than or equal to 3 mm above normal for race and sex or d) periodic or constant diplopia OR Documentation of chronic TED with one or more of the following: a) a 3 mm or greater increase in exophthalmos from before diagnosis of TED, b) exophthalmos greater than or equal to 3 mm above normal for race and sex.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an ophthalmologist or endocrinologist
Coverage Duration	Request will be authorized for 6 months.
Other Criteria	For active TED: member has had a trial and failure, contraindication to, or medical reason for not using oral or IV glucocorticoids.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TERIPARATIDE

Products Affected

- BONSITY
- teriparatide subcutaneous solution pen-injector 560 mcg/2.24ml*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation showing patient falls into one of the following categories: a bone mineral density (BMD) value consistent with osteoporosis (i.e., T-scores equal to or less than -2.5) or patient has had an osteoporotic fracture or patient has T-scores from -1.5 to -2.5 at the femoral neck or spine, and a 10-year probability of hip fracture greater than or equal to 3% or a 10-year probability of any major osteoporosis-related fracture greater than or equal to 20% based on the United States-adapted FRAX model.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	In addition, the following criteria is also applicable: 1) Trial of, medical reason for not using, or contraindication to an oral bisphosphonate and a denosumab product.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

THIOLA

Products Affected

- *tiopronin oral*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TIGECYCLINE

Products Affected

- *tigecycline*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	(1) Patient must have documented diagnosis of one of the following infections: (a) complicated skin and skin structure infection, (b) complicated intraabdominal infection, (c) community-acquired pneumonia AND (2) Trial of, contraindication to, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using at least 2 preferred first-line antibiotics.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an infectious disease specialist.
Coverage Duration	Request will be authorized for 14 days.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TOBI PODHALER

Products Affected

- TOBI PODHALER

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Patient has documented diagnosis of both: 1) cystic fibrosis AND 2) pseudomonas aeruginosa
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TOLVAPTAN

Products Affected

- *tolvaptan*
- *tolvaptan (hyponatremia)*

PA Criteria	Criteria Details
Exclusion Criteria	Concomitant use with strong CYP3A4 inhibitors (i.e. clarithromycin, ketoconazole, itraconazole, ritonavir, lopinavir-ritonavir, indinavir-ritonavir, indinavir, nelfinavir, saquinavir, nefazodone, conivaptan, and telithromycin).
Required Medical Information	Reviewer will verify available patient claim history to confirm patient is not using a strong CYP3A4 inhibitor (i.e. clarithromycin, ketoconazole, itraconazole, ritonavir, lopinavir-ritonavir, indinavir-ritonavir, indinavir, nelfinavir, saquinavir, nefazodone, conivaptan, and telithromycin).
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist, endocrinologist, hepatologist, or nephrologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TOPICAL ANTINEOPLASTIC RETINOIDS

Products Affected

- *bexarotene*
- PANRETIN

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a dermatologist, oncologist or specialist for submitted diagnosis
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TOPICAL TESTOSTERONE

Products Affected

- testosterone transdermal gel 1.62 %, 12.5 mg/act (1%), 20.25 mg/1.25gm (1.62%), 20.25 mg/act (1.62%), 25 mg/2.5gm (1%), 40.5 mg/2.5gm (1.62%), 50 mg/5gm (1%)
- testosterone transdermal solution

PA Criteria	Criteria Details
Exclusion Criteria	Patient has history of prostate cancer or breast cancer.
Required Medical Information	New starts of topical testosterone therapy for hypogonadism must have both of the following characteristics of hypogonadism: 1) symptoms associated with hypogonadism (e.g. unexplained mild anemia, low libido, decreased energy, etc.) 2) Two separate instances of low serum total or free testosterone taken in the morning, as defined by the lab reference range.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TRANSDERMAL LIDOCAINE

Products Affected

- *lidocaine external patch 5 %*
- ZTLIDO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documented diagnosis of a medically-accepted indication.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	If the request is for the product ZTlido, must provide medical reason for not being able to use generic lidocaine 5% patch
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TREMFYA

Products Affected

- TREMFYA ONE-PRESS SUBCUTANEOUS SOLUTION PEN-INJECTOR
- TREMFYA PEN SUBCUTANEOUS SOLUTION AUTO-INJECTOR 100 MG/ML, 200 MG/2ML
- TREMFYA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 100 MG/ML, 200 MG/2ML
- TREMFYA-CD/UC INDUCTION

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For UC: Approve. Crohn's Disease (CD) - Initial: Trial of, contraindication to, or medical reason (i.e. patient has a diagnosis of severe Crohn's disease) for not using (i.e. severe Crohns disease), or contraindication to 1 of the following: mercaptopurine, azathioprine, methotrexate, sulfasalazine, or corticosteroid (e.g., prednisone, methylprednisolone). Continuation of therapy: patient has been receiving Tremfya for a minimum of 4 months and has a positive clinical response to therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

TRIENTINE

Products Affected

- CUVRIOR
- trientine hcl oral capsule 250 mg*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	If the request is for Cuvrior for new starts, member must have trial of, contraindication to, or medical reason for not using trientine hydrochloride.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TRIKAFTA

Products Affected

- TRIKAFTA

PA Criteria	Criteria Details
Exclusion Criteria	Combination use with Kalydeco, Orkambi, or Symdeko.
Required Medical Information	Documentation of CFTR gene that is responsive to elexacaftor-tezacaftor-ivacaftor treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or an expert in the treatment of cystic fibrosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

TYMLOS

Products Affected

- TYMLOS

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation showing patient falls into one of the following categories: a bone mineral density (BMD) value consistent with osteoporosis (i.e., T-scores equal to or less than -2.5) or patient has had an osteoporotic fracture or patient has T-scores from -1.5 to -2.5 at the femoral neck or spine, and a 10-year probability of hip fracture greater than or equal to 3% or a 10-year probability of any major osteoporosis-related fracture greater than or equal to 20% based on the United States-adapted FRAX model.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	The following criteria is also applicable: 1) trial of, contraindication to, or medical reason for not using an oral bisphosphonate and a denosumab product, and 2) therapy does not exceed 2 years.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

TYVASO

Products Affected

- TYVASO DPI MAINTENANCE KIT
INHALATION POWDER 112 X 32MCG
& 112 X64MCG, 112 X 48MCG & 112
X64MCG, 112 X 48MCG & 112
X80MCG, 16 MCG, 32 MCG, 48 MCG,
64 MCG, 80 MCG
- TYVASO DPI TITRATION KIT
INHALATION POWDER 16 & 32 & 48
MCG

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or cardiologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For the treatment of pulmonary arterial hypertension (PAH): 1) documentation of PAH WHO Group I classification and PAH Functional Class and 2) trial of, contraindication to, or medical reason for not using a generic phosphodiesterase inhibitor and a generic endothelin receptor antagonist. For the treatment of pulmonary hypertension associated with interstitial lung disease (PH-ILD, WHO Group 3): documentation of PH-ILD and PAH Functional Class.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	Yes

UPTRAVI

Products Affected

- UPTRAVI ORAL
- UPTRAVI TITRATION

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Documentation of pulmonary arterial hypertension (PAH) WHO Group I and PAH Functional Class.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist or cardiologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not using a generic phosphodiesterase inhibitor and a generic endothelin receptor antagonist.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

USTEKINUMAB

Products Affected

- IMULDOSA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML, 90 MG/ML
- SELARSDI INTRAVENOUS
- SELARSDI SUBCUTANEOUS SOLUTION
- SELARSDI SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML, 90 MG/ML
- STARJEMZA INTRAVENOUS
- STARJEMZA SUBCUTANEOUS SOLUTION
- STARJEMZA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML, 90 MG/ML
- STELARA INTRAVENOUS
- STELARA SUBCUTANEOUS SOLUTION 45 MG/0.5ML
- STELARA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML, 90 MG/ML
- STEQEYMA INTRAVENOUS
- STEQEYMA SUBCUTANEOUS SOLUTION
- STEQEYMA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML, 90 MG/ML
- *ustekinumab subcutaneous solution*
- *ustekinumab subcutaneous solution prefilled syringe 45 mg/0.5ml, 90 mg/ml*
- *ustekinumab-aauz subcutaneous solution prefilled syringe 45 mg/0.5ml, 90 mg/ml*
- *ustekinumab-aekn subcutaneous solution prefilled syringe 45 mg/0.5ml, 90 mg/ml*
- YESINTEK INTRAVENOUS
- YESINTEK SUBCUTANEOUS SOLUTION
- YESINTEK SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 45 MG/0.5ML, 90 MG/ML

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A

PA Criteria	Criteria Details
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

UZEDY

Products Affected

- UZEDY SUBCUTANEOUS MG/0.42ML, 200 MG/0.56ML, 250
SUSPENSION PREFILLED SYRINGE MG/0.7ML, 50 MG/0.14ML, 75
100 MG/0.28ML, 125 MG/0.35ML, 150 MG/0.21ML

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	The member has a documented history of receiving oral risperidone without any clinically significant side effects.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial and failure of, contraindication, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using at least two of the following: Abilify Maintena, Abilify Asimtufii, or Risperidone Microsphere ER.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

VALCHLOR

Products Affected

- VALCHLOR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an oncologist or dermatologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Trial of, contraindication to, or medical reason for not being able to use one of the following: a topical corticosteroids or a topical retinoids.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

VEMLIDY

Products Affected

- VEMLIDY

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts: attestation that member has been tested for HIV infection. If member is HIV-positive, Vemlidy is not used alone.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

VEOZAH

Products Affected

- VEOZAH

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Initial: (1) Documented diagnosis of moderate to severe vasomotor symptoms due to menopause AND (2) Trial of, contraindication to, or medical reason (e.g. intolerance, hypersensitivity or contraindication) for not using a hormonal therapy (e.g., estradiol, oral Premarin, Prempro). Reauthorization: (1) Documentation of positive clinical response to therapy (e.g., decrease in frequency or severity of vasomotor symptoms from baseline)
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

VIGABATRIN

Products Affected

- *vigabatrin*
- VIGAFYDE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For infantile spasms or West syndrome, the request will be approved. For diagnosis of refractory complex partial seizures: 1) documentation of diagnosis, and 2) attestation the member is currently receiving another antiepileptic drug, and 3) attestation the member has experienced treatment failure from two alternative antiepileptic agents.
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

VIJOICE

Products Affected

- VIJOICE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts, all of the following must be included: 1) Documentation of genetic testing confirming diagnosis AND 2) Member has at least one target lesion identified on imaging AND 3) Prescriber attests the patient's condition is severe or life-threatening and necessitates systemic treatment. For continuation of therapy or reauthorization, attestation of a positive clinical response (i.e. reduction in the sum of measurable target lesion volume, absence of progression of non-target lesions, absence of any new lesions, etc.).
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a geneticist, dermatologist, vascular surgeon, hematologist/oncologist, or other specialist in the treatment of PIK3CA-Related Overgrowth Spectrum(PROs).
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

VMAT-2 INHIBITORS

Products Affected

- AUSTEDO
- AUSTEDO XR
- AUSTEDO XR PATIENT TITRATION ORAL TABLET EXTENDED RELEASE THERAPY PACK 12 & 18 & 24 & 30 MG
- INGREZZA ORAL CAPSULE
- INGREZZA ORAL CAPSULE SPRINKLE
- INGREZZA ORAL CAPSULE THERAPY PACK
- *tetrabenazine*

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist, clinical geneticist, or psychiatrist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	If the request is for tetrabenazine, Ingrezza or Ingrezza Sprinkle, request will be approved. If the request is for Austedo or Austedo XR, the member must have trial of or medical reason for not using tetrabenazine. Reauthorization: Confirmation of improvement in tardive dyskinesia symptoms or chorea associated with Huntington disease symptoms.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

VORICONAZOLE

Products Affected

- *voriconazole intravenous*

PA Criteria	Criteria Details
Exclusion Criteria	Non-Part D indications.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 6 months.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

VOSEVI

Products Affected

- VOSEVI

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Detectable HCV RNA viral load prior to treatment within 6 months of request. In addition, documentation of treatment history, and if cirrhotic, documentation of compensated or decompensated cirrhosis.
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized for 12 weeks as per AASLD-IDSA guidance.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

VOWST

Products Affected

- VOWST

PA Criteria	Criteria Details
Exclusion Criteria	Treatment of Clostridioides difficile infection (CDI)
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	If all the criteria are met, the request will be approved for 1 month
Other Criteria	Diagnosis of at least 1 recurrent episode of CDI
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

WEGOVY

Products Affected

- WEGOVY SUBCUTANEOUS SOLUTION AUTO-INJECTOR 0.25 MG/0.5ML, 0.5 MG/0.5ML, 1 MG/0.5ML, 1.7 MG/0.75ML, 2.4 MG/0.75ML

PA Criteria	Criteria Details
Exclusion Criteria	The member has an indication of only weight reduction or maintenance for overweight or obesity. The member has concurrent use of any GLP-1 receptor agonist. The member has a personal history of medullary thyroid carcinoma. The member has Multiple Endocrine Neoplasia syndrome type 2.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Reduction of the risk of major adverse cardiovascular events in adults with established cardiovascular disease and either obesity or overweight- For new starts: The member has an indication for reducing the risk of adverse cardiovascular events (cardiovascular death, non-fatal myocardial infarction, or non-fatal stroke) in adults with established cardiovascular disease. Documentation demonstrates patient has established cardiovascular disease (e.g., prior myocardial infarction, prior stroke, symptomatic peripheral arterial disease). Documentation is provided that the patient is overweight or obese (defined as a BMI of greater than or equal to 27 kg/m ²). Treatment of noncirrhotic metabolic dysfunction-associated steatohepatitis (MASH) - For new starts: The member has a documented diagnosis of MASH with documentation of moderate to advanced liver fibrosis (fibrosis stage 2 or 3) as evidenced by liver biopsy, biomarker testing (AST to platelet ratio index score, fibrosis 4 score, NAFLD fibrosis score, etc.), transient elastography or ultrasonography. For

PA Criteria	Criteria Details
	continuation of therapy or reauthorization: Documentation is provided that the patient has a positive clinical response to treatment.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

WEGOVY TABLETS

Products Affected

- WEGOVY ORAL

PA Criteria	Criteria Details
Exclusion Criteria	The member has an indication of only weight reduction or maintenance for overweight or obesity. The member has concurrent use of any GLP-1 receptor agonist. The member has a personal history of medullary thyroid carcinoma. The member has Multiple Endocrine Neoplasia syndrome type 2.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Reduction of the risk of major adverse cardiovascular events in adults with established cardiovascular disease and either obesity or overweight - For new starts: The member has an indication for reducing the risk of adverse cardiovascular events (cardiovascular death, non-fatal myocardial infarction, or non-fatal stroke) in adults with established cardiovascular disease. Documentation demonstrates patient has established cardiovascular disease (i.e., prior myocardial infarction, prior stroke, symptomatic peripheral arterial disease). Documentation is provided that the patient is overweight or obese (defined as a BMI of greater than or equal to 27 kg/m ²). The member must have documentation or a medical reason for not being able to use the Wegovy injection. For continuation of therapy or reauthorization: Documentation is provided that the patient has a positive clinical response to treatment.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

WHITE BLOOD CELL STIMULATORS

Products Affected

- FULPHILA
- FYLNETRA
- LEUKINE INJECTION SOLUTION RECONSTITUTED
- NEULASTA ONPRO SUBCUTANEOUS SOLUTION PREFILLED SYRINGE
- NEULASTA SUBCUTANEOUS SOLUTION 4 MG/0.4ML
- NEULASTA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE
- ZARXIO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	For new starts for Neulasta and Fylnetra: documentation of trial of, contraindication to, or medical reason for not using Fulphila. Continuation of therapy or re-authorization criteria: diagnosis of chronic neutropenia or a medical reason for continued need for GCSF.
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	New starts will be authorized for 4 months. Cont of therapy or reauth until end of contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

WINREVAIR

Products Affected

- WINREVAIR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	Pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group 1)- Initial: [Note: documentation required] (1) PAH was confirmed by right heart catheterization AND (2) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg AND (3) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND (4) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood units. Continuation of therapy: patient has positive clinical response to treatment.
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist or pulmonologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Documented trial and failure of, or contraindication to combination therapy including one PDE-5 inhibitor AND one endothelin receptor antagonist. Documentation of platelet count of greater than 50,000/mm3.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

XDEM VY

Products Affected

- XDEM VY

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	N/A
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

XELJANZ

Products Affected

- *tofacitinib citrate er*
- *tofacitinib citrate oral solution*
- *tofacitinib citrate oral tablet*
- XELJANZ ORAL SOLUTION
- XELJANZ ORAL TABLET
- XELJANZ XR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For ankylosing spondylitis: Trial of, medical reason for not using, or contraindication to naproxen and 1 of the following: an adalimumab product, Enbrel, or Cosentyx. For pJIA: Trial of, medical reason for not using, or contraindication to 1 of the following DMARDs: methotrexate or leflunomide and 1 of the following: an adalimumab product or Enbrel. For PsA: Trial of, medical reason for not using, or contraindication to 1 of the following: Enbrel, an adalimumab product, or Cosentyx. For RA: Trial of, medical reason for not using, or contraindication to 1 disease modifying antirheumatic drug (DMARD) (methotrexate, leflunomide, or sulfasalazine) and 1 of the following: Enbrel or an adalimumab product. For UC: Approve. Continuation of therapy: patient has been receiving Xeljanz for a minimum of 4 months and has a positive clinical response.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

XERMELO

Products Affected

- XERMELO

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a gastroenterologist or an oncologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: 1) Attestation that diarrhea is inadequately controlled by stable dose of SSA therapy for at least three months. For continuation of therapy or reauthorization: 1) documentation of positive clinical response to xermelo and 2) Attestation to continue to be used in combination with SSA.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

XIFAXAN

Products Affected

- XIFAXAN

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	For HE: gastroenterologist or hepatologist. For IBS-D: gastroenterologist.
Coverage Duration	For HE: contract year. For IBSD: 14 days (cannot exceed 3 courses of 14 days each). For TD: 3 days.
Other Criteria	For diagnosis of hepatic encephalopathy (HE): trial of, contraindication to, or medical reason for not using lactulose. For diagnosis of irritable bowel syndrome with diarrhea (IBSD): No more than 3 courses of 14 days each. For travelers diarrhea (TD) caused by noninvasive strains of E. Coli (with no bloody stools or fever): patient must be intolerant to or must have had a trial of at least 3 days of one of the following agents: ciprofloxacin, ofloxacin, levofloxacin or azithromycin.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

XOLAIR

Products Affected

- XOLAIR

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a pulmonologist, allergist, immunologist, dermatologist, or otolaryngologist.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	New starts for moderate to severe persistent allergic asthma: 1) Evidence of specific allergic sensitivity confirmed by positive skin test (i.e. prick/puncture test) or blood test (i.e. radioallergosorbent test) for a specific IgE or in vitro reactivity to a perennial aeroallergen, AND 2) Pretreatment serum IgE levels greater than 30 IU/mL, AND 3) Symptoms are not adequately controlled with high-dose inhaled corticosteroid (ICS) plus additional controller medication (ie. long-acting B2 agonist) for at least 3 months, or there is a medical reason for not using these drugs. Continuation of therapy or reauthorization criteria for moderate to severe persistent allergic asthma: 1) Reduction in asthma exacerbation resulting in systemic steroid use and/or hospitalization, OR 2) Reduction of rescue inhaler use, OR 3) Documentation of improvement in pulmonary function tests since baseline (prior to initiation of Xolair). New starts for chronic idiopathic urticaria: 1) inadequate symptomatic relief despite trial of two weeks of at least one 2nd generation antihistamine (unless contraindicated), AND 2) disease must be severe enough to warrant short term systemic corticosteroid therapy for management of urticaria. Continuation of therapy or reauthorization criteria for chronic idiopathic urticaria: 1) improvement from baseline of symptoms associated with urticaria within 6 months of Xolair use. New starts for nasal polyps: 1) currently using an intranasal

PA Criteria	Criteria Details
	corticosteroid, will be prescribed an intranasal corticosteroid with request, or has a medical reason for not using an intranasal corticosteroid. Continuation of therapy or reauthorization criteria for nasal polyps: 1) Documentation has been provided that demonstrates a clinical benefit (e.g. improvements in symptom severity, nasal polyp score [NPS], sino-nasal outcome test-22 [SNOT-22], nasal congestion score [NCS]) AND 2) continued use of intranasal corticosteroid, or has a medical reason for not using one. New starts for food allergy: 1) diagnosis of IgE-mediated food allergy 2) Xolair will be used in conjunction with food allergen avoidance. Continuation of therapy or reauthorization criteria for food allergy: 1) Documentation of clinical benefit
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

XOLREMDI

Products Affected

- XOLREMDI

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be an immunologist or a hematologist
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: 1) A documented diagnosis of WHIM (warts, hypogammaglobulinemia, infections and myelokathexis) syndrome confirmed by genotype variant of chemokine receptor 4 (CXCR4) and absolute neutrophil count (ANC) of less than or equal to 400 cells/microliter or white blood cells (WBC) less than or equal to 400 cells/microliter and 2) Documentation of baseline ANC and absolute lymphocyte count (ALC). For renewal 1) Documentation or provider attestation of positive clinical response (i.e. improvement from baseline in ANC, WBC and/or ALC or reduced frequency, duration, or severity of infections, fewer warts, or improved or stabilized clinical signs and/or symptoms of WHIM).
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No

PA Criteria	Criteria Details
Prerequisite Therapy Required	No

XYWAV

Products Affected

- XYWAV

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a sleep specialist, pulmonologist or a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For treatment of somnolence associated with narcolepsy, patient must have documentation of either trial of or a medical reason for being unable to use a CNS stimulant (e.g. methylphenidate, modafinil, armodafinil, etc.). For the treatment of cataplexy associated with narcolepsy or idiopathic hypersomnia, approve.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

YORVIPATH

Products Affected

- YORVIPATH

PA Criteria	Criteria Details
Exclusion Criteria	Diagnosis of acute post-surgical hypoparathyroidism.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with an endocrinologist.
Coverage Duration	New starts will be authorized for 6 months. Cont of therapy or reauth until end of contract year.
Other Criteria	For new starts: 1) Documented diagnosis of chronic hypoparathyroidism AND 2) Provider attestation that patient is currently receiving or has medical reason for not receiving calcium supplementation and active vitamin D treatment AND 3) An albumin-corrected serum calcium level of 7.8 mg/dL or greater. For reauthorization: Documentation of improvement in albumin-corrected serum calcium from baseline.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

YUTREPIA

Products Affected

- YUTREPIA

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a cardiologist or a pulmonologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For the treatment of pulmonary arterial hypertension (PAH): 1) documentation of PAH WHO Group I classification and PAH Functional Class and 2) trial of, contraindication to, or medical reason for not using a generic phosphodiesterase inhibitor and a generic endothelin receptor antagonist. For the treatment of pulmonary hypertension associated with interstitial lung disease (PH-ILD, WHO Group 3): documentation of PHILD and PAH Functional Class.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ZEPBOUND

Products Affected

- ZEPBOUND KWIKPEN
- ZEPBOUND SUBCUTANEOUS SOLUTION AUTO-INJECTOR

PA Criteria	Criteria Details
Exclusion Criteria	The member has an indication of only weight reduction or maintenance for overweight or obesity. The member has concurrent use of any GLP-1 receptor agonist. The member has a personal history of medullary thyroid carcinoma. The member has Multiple Endocrine Neoplasia syndrome type 2.
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a sleep disorder specialist, pulmonologist, ENT, or other provider specializing in obstructive sleep apnea.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For new starts: The member has an indication for moderate to severe obstructive sleep apnea (OSA) in adults with obesity. Documentation of diagnosis of OSA through polysomnography (sleep study) with an apnea-hypopnea index of 15 or more events per hour, or five or more events per hour in the presence of symptoms (e.g., cognitive impairment, fatigue, insomnia, loud snoring) or cardiovascular comorbidities (e.g., hypertension, ischemic heart disease, previous stroke). Documentation is provided that the patient is obese (defined as a BMI of greater than or equal to 30 kg/m ²). For continuation of therapy: Documentation of positive response to treatment. Documentation member has achieved and/or maintained a decrease in weight since baseline.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A

PA Criteria	Criteria Details
Part B Prerequisite	No
Prerequisite Therapy Required	No

ZEPOSIA

Products Affected

- ZEPOSIA
- ZEPOSIA 7-DAY STARTER PACK
- ZEPOSIA STARTER KIT ORAL CAPSULE THERAPY PACK 0.23MG & 0.46MG 0.92MG(21)

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Specialist for submitted diagnosis.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	For multiple sclerosis: Trial of, contraindication to, or medical reason for not using two of the following: dalfampridine ER, dimethyl fumarate, fingolimod, glatiramer, glatopa, or teriflunomide. For ulcerative colitis: Either 1) Trial of, medical reason for not using, or contraindication to 1 of the following: an ustekinumab product or an adalimumab product or 2) If utilized within the past 120 days, approve for continuation of therapy.
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ZILBRYSQ

Products Affected

- ZILBRYSQ

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescriber must be a neurologist, rheumatologist, or other appropriate specialist
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	Patient has tried and failed, a medical reason for not using, or has a contraindication to two (2) or more conventional therapies (i.e. pyridostigmine, corticosteroids, or non-steroidal immunosuppressive therapies)
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	Yes

ZTALMY

Products Affected

- ZTALMY

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	N/A
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a neurologist.
Coverage Duration	Request will be authorized until the end of the contract year.
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

ZURZUVAE

Products Affected

- ZURZUVAE

PA Criteria	Criteria Details
Exclusion Criteria	N/A
Required Medical Information	The member has a documented diagnosis of postpartum depression
Age Restrictions	N/A
Prescriber Restrictions	Prescribed by or in consultation with a psychiatrist or obstetrician/gynecologist
Coverage Duration	Request will be authorized until the end of the contract year
Other Criteria	N/A
Indications	All Medically-accepted Indications.
Off-Label Uses	N/A
Part B Prerequisite	No
Prerequisite Therapy Required	No

PART B VERSUS PART D

Products Affected

- *acetylcysteine inhalation solution 10 %, 20 %*
- *acyclovir sodium intravenous solution 50 mg/ml*
- *albuterol sulfate inhalation nebulization solution (2.5 mg/3ml) 0.083%, (5 mg/ml) 0.5%, 0.63 mg/3ml, 1.25 mg/3ml, 2.5 mg/0.5ml*
- *amphotericin b intravenous solution reconstituted 50 mg*
- *amphotericin b liposome intravenous suspension reconstituted 50 mg*
- *aprepitant oral capsule 125 mg, 40 mg, 80 mg*
- *aprepitant oral capsule therapy pack 80 & 125 mg*
- **ASTAGRAF XL ORAL CAPSULE EXTENDED RELEASE 24 HOUR 0.5 MG, 1 MG, 5 MG**
- *azathioprine oral tablet 50 mg*
- *budesonide inhalation suspension 0.25 mg/2ml, 0.5 mg/2ml, 1 mg/2ml*
- **CLINISOL SF INTRAVENOUS SOLUTION 15 %**
- *cromolyn sodium inhalation nebulization solution 20 mg/2ml*
- *cyclophosphamide oral capsule 25 mg, 50 mg*
- *cyclophosphamide oral tablet 25 mg, 50 mg*
- *cyclosporine modified oral capsule 100 mg, 25 mg, 50 mg*
- *cyclosporine modified oral solution 100 mg/ml*
- *cyclosporine oral capsule 100 mg, 25 mg*
- *dronabinol oral capsule 10 mg, 2.5 mg, 5 mg*
- **EMEND ORAL SUSPENSION RECONSTITUTED 125 MG/5ML**
- **ENERGIX-B INJECTION SUSPENSION 20 MCG/ML**
- **ENERGIX-B INJECTION SUSPENSION PREFILLED SYRINGE 10 MCG/0.5ML, 20 MCG/ML**
- **ENVARBUS XR ORAL TABLET EXTENDED RELEASE 24 HOUR 0.75 MG, 1 MG, 4 MG**
- *everolimus oral tablet 0.25 mg, 0.5 mg, 0.75 mg, 1 mg*
- *formoterol fumarate inhalation nebulization solution 20 mcg/2ml*
- **GAMMAGARD ERC INJECTION SOLUTION 10 GM/100ML, 5 GM/50ML**
- **GAMMAGARD INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 2.5 GM/25ML, 20 GM/200ML, 30 GM/300ML, 5 GM/50ML**
- **GAMMAGARD S/D LESS IGA INTRAVENOUS SOLUTION RECONSTITUTED 10 GM, 5 GM**
- **GAMMAKED INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 20 GM/200ML, 5 GM/50ML**
- **GAMMAPLEX INTRAVENOUS SOLUTION 10 GM/100ML, 10 GM/200ML, 20 GM/200ML, 20 GM/400ML, 5 GM/100ML, 5 GM/50ML**
- **GAMUNEX-C INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 2.5 GM/25ML, 20 GM/200ML, 40 GM/400ML, 5 GM/50ML**
- **GENGRAF ORAL CAPSULE 100 MG, 25 MG**
- *granisetron hcl oral tablet 1 mg*
- **HEPLISAV-B INTRAMUSCULAR SOLUTION PREFILLED SYRINGE 20 MCG/0.5ML**

- IMOVAX RABIES INTRAMUSCULAR SUSPENSION RECONSTITUTED 2.5 UNIT/ML
- INTRALIPID INTRAVENOUS EMULSION 20 %, 30 %
- *ipratropium bromide inhalation solution* 0.02 %
- *ipratropium-albuterol inhalation solution* 0.5-2.5 (3) mg/3ml
- *levalbuterol hcl inhalation nebulization solution* 0.31 mg/3ml, 0.63 mg/3ml, 1.25 mg/3ml
- *mycophenolate mofetil oral capsule* 250 mg
- *mycophenolate mofetil oral suspension reconstituted* 200 mg/ml
- *mycophenolate mofetil oral tablet* 500 mg
- *mycophenolate sodium oral tablet delayed release* 180 mg, 360 mg
- NUTRILIPID INTRAVENOUS EMULSION 20 %
- OCTAGAM INTRAVENOUS SOLUTION 1 GM/20ML, 10 GM/100ML, 10 GM/200ML, 2 GM/20ML, 2.5 GM/50ML, 20 GM/200ML, 30 GM/300ML, 5 GM/100ML, 5 GM/50ML
- *ondansetron hcl oral solution* 4 mg/5ml
- *ondansetron hcl oral tablet* 24 mg, 4 mg, 8 mg
- *ondansetron oral tablet dispersible* 4 mg, 8 mg
- *pentamidine isethionate inhalation solution reconstituted* 300 mg
- PLENAMINE INTRAVENOUS SOLUTION 15 %
- PRIVIGEN INTRAVENOUS SOLUTION 10 GM/100ML, 20 GM/200ML, 40 GM/400ML, 5 GM/50ML
- PROGRAF INTRAVENOUS SOLUTION 5 MG/ML
- PROGRAF ORAL PACKET 0.2 MG, 1 MG
- PULMOZYME INHALATION SOLUTION 2.5 MG/2.5ML
- RABAVERT INTRAMUSCULAR SUSPENSION RECONSTITUTED
- RECOMBIVAX HB INJECTION SUSPENSION 10 MCG/ML, 40 MCG/ML, 5 MCG/0.5ML
- RECOMBIVAX HB INJECTION SUSPENSION PREFILLED SYRINGE 10 MCG/ML, 5 MCG/0.5ML
- *sirolimus oral solution* 1 mg/ml
- *sirolimus oral tablet* 0.5 mg, 1 mg, 2 mg
- *tacrolimus er oral capsule extended release 24 hour* 0.5 mg, 1 mg, 5 mg
- *tacrolimus intravenous solution* 5 mg/ml
- *tacrolimus oral capsule* 0.5 mg, 1 mg, 5 mg
- TENIVAC INTRAMUSCULAR INJECTABLE 5-2 LFU (INJECTION)
- TENIVAC INTRAMUSCULAR SUSPENSION 5-2 LF/0.5ML
- *tobramycin inhalation nebulization solution* 300 mg/4ml, 300 mg/5ml

Details

This drug may be covered under Medicare Part B or D depending upon the circumstances. Information may need to be submitted describing the use and setting of the drug to make the determination.

Index

A

abiraterone acetate 148, 150
 ABIRTEGA 148, 150
 acetylcysteine inhalation solution 10 %, 20
 % 386
 acitretin 136, 137
 ACTEMRA ACTPEN 140
 ACTEMRA SUBCUTANEOUS 140
 acyclovir sodium intravenous solution 50
 mg/ml 386
 adalimumab-fkjp (2 pen)..... 141, 142
 adalimumab-fkjp (2 syringe) subcutaneous
 prefilled syringe kit 20 mg/0.4ml, 40
 mg/0.8ml 141, 142
 ADEMPAS 143, 144
 AIMOVIG..... 169, 170
 AKEEGA 148, 150
 albuterol sulfate inhalation nebulization
 solution (2.5 mg/3ml) 0.083%, (5 mg/ml)
 0.5%, 0.63 mg/3ml, 1.25 mg/3ml, 2.5
 mg/0.5ml 386
 ALECENSA..... 148, 150
 ALUNBRIG..... 148, 150
 ALYFTREK ORAL TABLET 10-50-125
 MG, 4-20-50 MG 146
 ambrisentan 147
 amitriptyline hcl oral 222
 amoxapine 222
 amphotericin b intravenous solution
 reconstituted 50 mg 386
 amphotericin b liposome intravenous
 suspension reconstituted 50 mg 386
 apomorphine hcl subcutaneous 151
 aprepitant oral capsule 125 mg, 40 mg, 80
 mg 386
 aprepitant oral capsule therapy pack 80 &
 125 mg 386
 AQNEURSA..... 152
 ARALAST NP INTRAVENOUS
 SOLUTION RECONSTITUTED 1000
 MG, 500 MG..... 145

ARANESP (ALBUMIN FREE)
 INJECTION SOLUTION 100 MCG/ML,
 200 MCG/ML, 25 MCG/ML, 40
 MCG/ML, 60 MCG/ML 199, 200
 ARANESP (ALBUMIN FREE)
 INJECTION SOLUTION PREFILLED
 SYRINGE 199, 200
 ARCALYST 153
 ARIKAYCE..... 154
 ARISTADA INITIO 155
 ARISTADA INTRAMUSCULAR
 PREFILLED SYRINGE 1064
 MG/3.9ML, 441 MG/1.6ML, 662
 MG/2.4ML, 882 MG/3.2ML 155
 armodafinil 264
 ASTAGRAF XL ORAL CAPSULE
 EXTENDED RELEASE 24 HOUR 0.5
 MG, 1 MG, 5 MG 386
 AUGTYRO 148, 150
 AUSTEDO..... 358
 AUSTEDO XR 358
 AUSTEDO XR PATIENT TITRATION
 ORAL TABLET EXTENDED
 RELEASE THERAPY PACK 12 & 18 &
 24 & 30 MG 358
 AUVELITY 156
 AUVELITY TITRATION PACK 156
 AVMAPKI FAKZYNJA CO-PACK..... 148,
 150
 AVTOZMA SUBCUTANEOUS..... 140
 AYVAKIT 148, 150
 azathioprine oral tablet 50 mg..... 386
B
 BAC (BUTALBITAL-ACETAMIN-CAFF)
 223
 BAFIERTAM 256, 257
 BALVERSA 148, 150
 BENLYSTA SUBCUTANEOUS... 158, 159
 benztropine mesylate oral 220, 221
 BESREMI 160

BETASERON SUBCUTANEOUS KIT 256, 257	
bexarotene	148, 150, 339
BONSITY	334
bosentan oral tablet	161
BOSULIF	148, 150
bosutinib.....	148, 150
BRAFTOVI ORAL CAPSULE 75 MG	148, 150
BRINSUPRI.....	162
BRUKINSA	148, 150
budesonide inhalation suspension 0.25 mg/2ml, 0.5 mg/2ml, 1 mg/2ml	386
butalbital-acetaminophen oral tablet 50-325 mg	223
butalbital-apap-caff-cod oral capsule 50-325-40-30 mg.....	223
butalbital-apap-caffeine oral capsule 50-325-40 mg	223
butalbital-apap-caffeine oral solution	223
butalbital-apap-caffeine oral tablet 50-325-40 mg	223
butalbital-asa-caff-codeine.....	223
butalbital-aspirin-caffeine oral capsule...	223
BYSANTI	272, 273
BYSANTI TITRATION PACK A .	272, 273
BYSANTI TITRATION PACK B..	272, 273
BYSANTI TITRATION PACK C..	272, 273
C	
CABOMETYX	148, 150
CALQUENCE ORAL TABLET	148, 150
CAMZYOS	163, 164
CAPLYTA	272, 273
CAPRELSA	148, 150
CARDAMYST	165
carglumic acid oral tablet soluble	166
carisoprodol oral	225
caspofungin acetate.....	167
CAYSTON.....	157
CERDELGA	168
chlorzoxazone oral tablet 500 mg	225
CHOLBAM.....	171
CIBINQO	172
CIMZIA (1 SYRINGE)	173, 174
CIMZIA (2 SYRINGE)	173, 174
CIMZIA SUBCUTANEOUS KIT 2 X 200 MG	173, 174
CIMZIA-STARTER	173, 174
CINRYZE	219
cladribine (10 tabs)	256, 257
cladribine (4 tabs)	256, 257
cladribine (5 tabs)	256, 257
cladribine (6 tabs)	256, 257
cladribine (7 tabs)	256, 257
cladribine (8 tabs)	256, 257
cladribine (9 tabs)	256, 257
cladribine 10 mg tablet therapy pack oral	256, 257
CLINISOL SF INTRAVENOUS SOLUTION 15 %	386
clomipramine hcl oral	222
COBENFY	272, 273
COBENFY STARTER PACK	272, 273
COMETRIQ (100 MG DAILY DOSE) ORAL KIT 80 & 20 MG	148, 150
COMETRIQ (140 MG DAILY DOSE) ORAL KIT 3 X 20 MG & 80 MG	148, 150
COMETRIQ (60 MG DAILY DOSE)...	148, 150
COPIKTRA.....	148, 150
CORLANOR ORAL SOLUTION.....	175
CORTROPHIN	176
CORTROPHIN GEL	176
COSENTYX (300 MG DOSE).....	177
COSENTYX INTRAVENOUS.....	177
COSENTYX SENSOREADY (300 MG).....	177
COSENTYX SENSOREADY PEN	177
COSENTYX SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 150 MG/ML, 75 MG/0.5ML	177
COSENTYX UNOREADY	177
COTELLIC	148, 150
CRESEMBA ORAL	178
cromolyn sodium inhalation nebulization solution 20 mg/2ml	386
CUVRIOR.....	344

cyclophosphamide oral capsule 25 mg, 50 mg 386
cyclophosphamide oral tablet 25 mg, 50 mg 386
cyclosporine modified oral capsule 100 mg, 25 mg, 50 mg 386
cyclosporine modified oral solution 100 mg/ml 386
cyclosporine oral capsule 100 mg, 25 mg 386
cyproheptadine hcl oral syrup 2 mg/5ml 220, 221
cyproheptadine hcl oral tablet 220, 221
CYSTAGON 179
CYSTARAN 180
D
dalfampridine er 181
DANZITEN 148, 150
dasatinib 148, 150
DAURISMO 148, 150
deferasirox 182
deferasirox granules 182
deferiprone 183
DIACOMIT 185
dihydroergotamine mesylate nasal 186
dimethyl fumarate oral capsule delayed release 120 mg, 240 mg 256, 257
dimethyl fumarate starter pack oral capsule delayed release therapy pack 256, 257
diphenoxylate-atropine oral liquid.. 220, 221
diphenoxylate-atropine oral tablet 2.5-0.025 mg 220, 221
dipyridamole oral 220, 221
DOPTELET 187
DOPTELET SPRINKLE 187
doxepin hcl external 188
dronabinol oral capsule 10 mg, 2.5 mg, 5 mg 386
DUPIXENT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 189, 190
DUPIXENT SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 200 MG/1.14ML, 300 MG/2ML 189, 190
E
EGRIFTA SV 191

EGRIFTA WR 191
ELIGARD 215
eltrombopag olamine oral packet 12.5 mg, 25 mg 294
eltrombopag olamine oral tablet 12.5 mg, 25 mg, 50 mg, 75 mg 294
EMEND ORAL SUSPENSION RECONSTITUTED 125 MG/5ML 386
EMGALITY 169, 170
EMGALITY (300 MG DOSE) 169, 170
EMSAM 192
ENBREL MINI 193, 194
ENBREL SUBCUTANEOUS SOLUTION 25 MG/0.5ML 193, 194
ENBREL SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 193, 194
ENBREL SURECLICK SUBCUTANEOUS SOLUTION AUTO-INJECTOR 193, 194
ENGERIX-B INJECTION SUSPENSION 20 MCG/ML 386
ENGERIX-B INJECTION SUSPENSION PREFILLED SYRINGE 10 MCG/0.5ML, 20 MCG/ML 386
ENSACOVE 148, 150
ENTYVIO PEN 196
ENVARISUS XR ORAL TABLET EXTENDED RELEASE 24 HOUR 0.75 MG, 1 MG, 4 MG 386
EPIDIOLEX 197
EPOGEN INJECTION SOLUTION 10000 UNIT/ML, 2000 UNIT/ML, 20000 UNIT/ML, 3000 UNIT/ML, 4000 UNIT/ML 199, 200
ergotamine-caffeine 220, 221
ERIVEDGE 148, 150
ERLEADA 148, 150
erlotinib hcl 148, 150
ERZOFRI INTRAMUSCULAR SUSPENSION PREFILLED SYRINGE 117 MG/0.75ML, 156 MG/ML, 234 MG/1.5ML, 351 MG/2.25ML, 39 MG/0.25ML, 78 MG/0.5ML 201
eszopiclone 226
EUCRISA 202

EULEXIN 148, 150
everolimus oral tablet 0.25 mg, 0.5 mg, 0.75 mg, 1 mg 386
everolimus oral tablet 10 mg, 2.5 mg, 5 mg, 7.5 mg 148, 150
everolimus oral tablet soluble 148, 150
EVRYSDI 203
F
FABHALTA 204, 205
FANAPT 272, 273
FANAPT TITRATION PACK A ... 272, 273
FANAPT TITRATION PACK B ORAL TABLET 272, 273
FANAPT TITRATION PACK C ORAL TABLET 272, 273
FASENRA 206, 207
FASENRA PEN..... 206, 207
FILSPARI 208
fingolimod hcl 256, 257
FINTEPLA..... 209
FIRDAPSE..... 210
FIRMAGON (240 MG DOSE)..... 215
FIRMAGON SUBCUTANEOUS SOLUTION RECONSTITUTED 80 MG 215
flucytosine oral..... 211
formoterol fumarate inhalation nebulization solution 20 mcg/2ml 386
FOTIVDA 148, 150
FRUZAQLA 148, 150
FULPHILA 366
FYLNETRA..... 366
G
GALAFOLD 212
GAMMAGARD ERC INJECTION SOLUTION 10 GM/100ML, 5 GM/50ML 386
GAMMAGARD INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 2.5 GM/25ML, 20 GM/200ML, 30 GM/300ML, 5 GM/50ML 386
GAMMAGARD S/D LESS IGA INTRAVENOUS SOLUTION RECONSTITUTED 10 GM, 5 GM 386

GAMMAKED INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 20 GM/200ML, 5 GM/50ML 386
GAMMAPLEX INTRAVENOUS SOLUTION 10 GM/100ML, 10 GM/200ML, 20 GM/200ML, 20 GM/400ML, 5 GM/100ML, 5 GM/50ML 386
GAMUNEX-C INJECTION SOLUTION 1 GM/10ML, 10 GM/100ML, 2.5 GM/25ML, 20 GM/200ML, 40 GM/400ML, 5 GM/50ML 386
GATTEX..... 213
GAVRETO 148, 150
gefitinib 148, 150
GENGRAF ORAL CAPSULE 100 MG, 25 MG 386
GENOTROPIN MINIQUEL SUBCUTANEOUS PREFILLED SYRINGE 217, 218
GENOTROPIN SUBCUTANEOUS CARTRIDGE..... 217, 218
GILOTRIF 148, 150
GLASSIA..... 145
glatiramer acetate subcutaneous solution prefilled syringe 20 mg/ml, 40 mg/ml 256, 257
GLATOPA SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 20 MG/ML, 40 MG/ML 256, 257
glyburide micronized oral tablet 3 mg, 6 mg 220, 221
glyburide oral tablet 1.25 mg, 2.5 mg, 5 mg 220, 221
glyburide-metformin oral tablet 1.25-250 mg 220, 221
glyburide-metformin oral tablet 2.5-500 mg, 5-500 mg 220, 221
glycerol phenylbutyrate 297
GOCOVRI 216
GOMEKLI 148, 150
granisetron hcl oral tablet 1 mg 386

guanfacine hcl er oral tablet extended
 release 24 hour 1 mg, 2 mg, 3 mg, 4 mg
 220, 221
 guanfacine hcl oral 220, 221
H
 HAEGARDA 219
 HEPLISAV-B INTRAMUSCULAR
 SOLUTION PREFILLED SYRINGE 20
 MCG/0.5ML 386
 HERNEXEOS 148, 150
 HETLIOZ LQ 330
 hydroxyzine hcl oral syrup 10 mg/5ml .. 220,
 221
 hydroxyzine hcl oral tablet..... 220, 221
 HYFTOR..... 227
 HYRNUO 148, 150
I
 IBRANCE 148, 150
 IBTROZI..... 148, 150
 icatibant acetate subcutaneous solution
 prefilled syringe 228
 ICLUSIG 148, 150
 IDHIFA 148, 150
 ILARIS SUBCUTANEOUS SOLUTION
 229
 ILUMYA..... 230
 imatinib mesylate oral 148, 150
 IMBRUVICA ORAL CAPSULE 231
 IMBRUVICA ORAL SUSPENSION..... 231
 IMBRUVICA ORAL TABLET 140 MG,
 280 MG, 420 MG..... 231
 imkeldi 148, 150
 IMOVAX RABIES INTRAMUSCULAR
 SUSPENSION RECONSTITUTED 2.5
 UNIT/ML 387
 IMPAVIDO..... 232
 IMULDOSA SUBCUTANEOUS
 SOLUTION PREFILLED SYRINGE 45
 MG/0.5ML, 90 MG/ML 350, 351
 INCRELEX..... 233
 indomethacin oral capsule 25 mg, 50 mg
 220, 221
 INGREZZA ORAL CAPSULE..... 358

INGREZZA ORAL CAPSULE SPRINKLE
 358
 INGREZZA ORAL CAPSULE THERAPY
 PACK..... 358
 INLURIYO 148, 150
 INLYTA..... 148, 150
 INQOVI 148, 150
 INREBIC..... 148, 150
 INTRALIPID INTRAVENOUS
 EMULSION 20 %, 30 % 387
 INVEGA HAFYERA INTRAMUSCULAR
 SUSPENSION PREFILLED SYRINGE
 1092 MG/3.5ML, 1560 MG/5ML 283,
 284
 INVEGA SUSTENNA
 INTRAMUSCULAR SUSPENSION
 PREFILLED SYRINGE 117
 MG/0.75ML, 156 MG/ML, 234
 MG/1.5ML, 39 MG/0.25ML, 78
 MG/0.5ML 283, 284
 INVEGA TRINZA INTRAMUSCULAR
 SUSPENSION PREFILLED SYRINGE
 273 MG/0.88ML, 410 MG/1.32ML, 546
 MG/1.75ML, 819 MG/2.63ML .. 283, 284
 ipratropium bromide inhalation solution
 0.02 % 387
 ipratropium-albuterol inhalation solution
 0.5-2.5 (3) mg/3ml 387
 ITOVEBI..... 148, 150
 ivabradine hcl..... 175
 IWILFIN 148, 150
J
 JAKAFI..... 234
 JAKAFI XR 234
 JAYPIRCA 148, 150
 JYLAMVO 251
K
 KALYDECO..... 235
 KERENDIA 236, 237
 KESIMPTA..... 256, 257
 ketorolac tromethamine oral 220, 221
 KEVZARA 238
 KINERET SUBCUTANEOUS SOLUTION
 PREFILLED SYRINGE 239

KISQALI (200 MG DOSE)..... 148, 150
 KISQALI (400 MG DOSE)..... 148, 150
 KISQALI (600 MG DOSE)..... 148, 150
 KOMZIFTI 148, 150
 KOSELUGO 148, 150
 KRAZATI 148, 150
 KYMBEE ORAL TABLET 184

L

lapatinib ditosylate 148, 150
 LAZCLUZE 148, 150
 lenalidomide..... 148, 150
 LENVIMA (10 MG DAILY DOSE) 148, 150
 LENVIMA (12 MG DAILY DOSE) 148, 150
 LENVIMA (14 MG DAILY DOSE) 148, 150
 LENVIMA (18 MG DAILY DOSE) 148, 150
 LENVIMA (20 MG DAILY DOSE) 148, 150
 LENVIMA (24 MG DAILY DOSE) 148, 150
 LENVIMA (4 MG DAILY DOSE) 148, 150
 LENVIMA (8 MG DAILY DOSE) 148, 150
 LEQEMBI IQLIK 240
 LEQSELVI 241
 LEUKERAN 148, 150
 LEUKINE INJECTION SOLUTION
 RECONSTITUTED 366
 leuprolide acetate (3 month) 215
 leuprolide acetate injection 215
 levalbuterol hcl inhalation nebulization
 solution 0.31 mg/3ml, 0.63 mg/3ml, 1.25
 mg/3ml 387
 l-glutamine oral packet 195
 lidocaine external patch 5 % 341
 LIFYORLI (125 MG DOSE)..... 148, 150
 LIFYORLI (150 MG DOSE)..... 148, 150
 liraglutide 214
 LITFULO 242
 LIVMARLI 243, 244
 LIVTENCITY 245
 LODOCO 246

lofexidine hcl 247
 LONSURF 148, 150
 LORBRENA 148, 150
 LUMAKRAS 149, 150
 LUPKYNIS..... 248
 LUPRON DEPOT (1-MONTH) 215
 LUPRON DEPOT (3-MONTH) 215
 LUPRON DEPOT (4-MONTH) 215
 LUPRON DEPOT (6-MONTH) 215
 LUTRATE DEPOT 215
 LYBALVI 249
 LYNPARZA ORAL TABLET 149, 150
 LYTGObI (12 MG DAILY DOSE) 149, 150
 LYTGObI (16 MG DAILY DOSE) 149, 150
 LYTGObI (20 MG DAILY DOSE) 149, 150

M

MAVYRET 250
 MAYZENT 256, 257
 MAYZENT STARTER PACK 256, 257
 megestrol acetate oral suspension 40 mg/ml,
 400 mg/10ml, 625 mg/5ml 224
 megestrol acetate oral tablet 224
 MEKINIST 149, 150
 MEKTOVI 149, 150
 mercaptopurine oral suspension 149, 150
 metaxalone oral tablet 800 mg 225
 methocarbamol oral tablet 500 mg, 750 mg
 225
 methyltestosterone oral 252
 metyrosine 253
 mifepristone oral tablet 300 mg 254
 miglustat 255
 modafinil oral 264
 MODEYSO 149, 150
 MOUNJARO SUBCUTANEOUS
 SOLUTION AUTO-INJECTOR 214
 mycophenolate mofetil oral capsule 250 mg
 387
 mycophenolate mofetil oral suspension
 reconstituted 200 mg/ml 387
 mycophenolate mofetil oral tablet 500 mg
 387
 mycophenolate sodium oral tablet delayed
 release 180 mg, 360 mg 387

MYFEMBREE..... 258, 259
 MYQORZO 260, 261
N
 NAYZILAM 262
 NERLYNX 149, 150
 NEULASTA ONPRO SUBCUTANEOUS
 SOLUTION PREFILLED SYRINGE 366
 NEULASTA SUBCUTANEOUS
 SOLUTION 4 MG/0.4ML 366
 NEULASTA SUBCUTANEOUS
 SOLUTION PREFILLED SYRINGE 366
 NEXLETOL..... 138, 139
 NEXLIZET 138, 139
 NGENLA 217, 218
 nifedipine oral 220, 221
 nilotinib d-tartrate 149, 150
 nilotinib hcl 149, 150
 nilutamide 149, 150
 NINLARO..... 149, 150
 nintedanib esylate..... 270
 nitisinone 263
 nortriptyline hcl oral 222
 NUBEQA..... 149, 150
 NUCALA..... 265, 266
 NUEDEXTA..... 267
 NUPLAZID ORAL CAPSULE..... 268
 NUPLAZID ORAL TABLET 10 MG.... 268
 NURTEC..... 169, 170
 NUTRILIPID INTRAVENOUS
 EMULSION 20 %..... 387
O
 OCTAGAM INTRAVENOUS SOLUTION
 1 GM/20ML, 10 GM/100ML, 10
 GM/200ML, 2 GM/20ML, 2.5
 GM/50ML, 20 GM/200ML, 30
 GM/300ML, 5 GM/100ML, 5 GM/50ML
 387
 octreotide acetate injection solution 100
 mcg/ml, 1000 mcg/ml, 200 mcg/ml, 50
 mcg/ml, 500 mcg/ml 269
 octreotide acetate intramuscular 269
 ODOMZO 149, 150
 OFEV 270

OGSIVEO ORAL TABLET 100 MG, 150
 MG 149, 150
 OJEMDA 149, 150
 OJJAARA 149, 150
 OMNITROPE SUBCUTANEOUS
 SOLUTION CARTRIDGE..... 217, 218
 OMNITROPE SUBCUTANEOUS
 SOLUTION RECONSTITUTED217, 218
 ondansetron hcl oral solution 4 mg/5ml.. 387
 ondansetron hcl oral tablet 24 mg, 4 mg, 8
 mg 387
 ondansetron oral tablet dispersible 4 mg, 8
 mg 387
 ONUREG..... 149, 150
 OPIPZA ORAL FILM 10 MG, 2 MG, 5 MG
 272, 273
 OPSUMIT 271
 ORENCIA CLICKJECT..... 274, 275
 ORENCIA INTRAVENOUS 274, 275
 ORENCIA SUBCUTANEOUS SOLUTION
 PREFILLED SYRINGE 125 MG/ML, 50
 MG/0.4ML, 87.5 MG/0.7ML 274, 275
 ORFADIN ORAL SUSPENSION..... 263
 ORGOVYX..... 149, 150
 ORIAHNN 276
 ORILISSA..... 277
 ORKAMBI..... 278
 ORLADEYO..... 219
 orphenadrine citrate er 225
 ORSERDU 149, 150
 OTEZLA 279
 OTEZLA XR 279
 OTEZLA/OTEZLA XR INITIATION PK
 279
 OXERVATE..... 280
 OXYCONTIN ORAL TABLET ER 12
 HOUR ABUSE-DETERRENT 10 MG,
 15 MG, 20 MG, 30 MG, 40 MG, 60 MG,
 80 MG 281, 282
 OZEMPIC (0.25 OR 0.5 MG/DOSE)
 SUBCUTANEOUS SOLUTION PEN-
 INJECTOR 2 MG/3ML 214

OZEMPIC (1 MG/DOSE) SUBCUTANEOUS SOLUTION PEN- INJECTOR 4 MG/3ML	214
OZEMPIC (2 MG/DOSE)	214
P	
PANRETIN	339
pazopanib hcl	149, 150
PEGASYS SUBCUTANEOUS SOLUTION 180 MCG/ML	286
PEGASYS SUBCUTANEOUS SOLUTION PREFILLED SYRINGE	286
PEMAZYRE	149, 150
penicillamine oral tablet	287
pentamidine isethionate inhalation solution reconstituted 300 mg	387
perphenazine-amitriptyline	222
PERSERIS	288
phenobarbital oral elixir	222
phenobarbital oral tablet	222
phenoxybenzamine hcl oral	289
PIQRAY (200 MG DAILY DOSE) 149, 150	
PIQRAY (250 MG DAILY DOSE) 149, 150	
PIQRAY (300 MG DAILY DOSE) 149, 150	
pirfenidone	290
PLENAMINE INTRAVENOUS SOLUTION 15 %	387
pomalidomide	149, 150
PONVORY	256, 257
PONVORY STARTER PACK	256, 257
posaconazole oral	291
pretomanid	292
PREVYMIS ORAL	293
PRIVIGEN INTRAVENOUS SOLUTION 10 GM/100ML, 20 GM/200ML, 40 GM/400ML, 5 GM/50ML	387
PROCRIT	199, 200
PROGRAF INTRAVENOUS SOLUTION 5 MG/ML	387
PROGRAF ORAL PACKET 0.2 MG, 1 MG	387
PROLASTIN-C INTRAVENOUS SOLUTION	145
promethazine hcl oral solution 6.25 mg/5ml	220, 221
promethazine hcl oral tablet	220, 221
promethazine hcl rectal suppository 12.5 mg, 25 mg	220, 221
promethazine-phenylephrine	220, 221
PROMETHEGAN RECTAL SUPPOSITORY 50 MG	220, 221
PULMOZYME INHALATION SOLUTION 2.5 MG/2.5ML	387
PYRUKYND	295
PYRUKYND TAPER PACK	295
Q	
QINLOCK	149, 150
QULIPTA	169, 170
R	
RABAVERT INTRAMUSCULAR SUSPENSION RECONSTITUTED... ..	387
RADICAVA ORS	296
RADICAVA ORS STARTER KIT	296
REBIF REBIDOSE SUBCUTANEOUS SOLUTION AUTO-INJECTOR	256, 257
REBIF REBIDOSE TITRATION PACK SUBCUTANEOUS SOLUTION AUTO- INJECTOR	256, 257
REBIF SUBCUTANEOUS SOLUTION PREFILLED SYRINGE	256, 257
REBIF TITRATION PACK SUBCUTANEOUS SOLUTION PREFILLED SYRINGE	256, 257
RECOMBIVAX HB INJECTION SUSPENSION 10 MCG/ML, 40 MCG/ML, 5 MCG/0.5ML	387
RECOMBIVAX HB INJECTION SUSPENSION PREFILLED SYRINGE 10 MCG/ML, 5 MCG/0.5ML	387
RECORLEV	298
RELISTOR ORAL	299
RELISTOR SUBCUTANEOUS SOLUTION	299
RELISTOR SUBCUTANEOUS SOLUTION PREFILLED SYRINGE	299
REPATHA	285
REPATHA SURECLICK	285
RETACRIT INJECTION SOLUTION 10000 UNIT/ML, 10000	

UNIT/ML(1ML), 2000 UNIT/ML, 20000
UNIT/ML, 3000 UNIT/ML, 4000
UNIT/ML, 40000 UNIT/ML 199, 200
RETEVMO ORAL TABLET 149, 150
REVCOV 300
REVLIMID 149, 150
REVUFORJ 149, 150
REXULTI 301
REZDIFFRA 302, 303
REZLIDHIA 149, 150
REZUROCK 304
ROMVIMZA 149, 150
ROZLYTREK 149, 150
RUBRACA 149, 150
rufinamide oral suspension 40 mg/ml 305
rufinamide oral tablet 305
RYBELSUS 214
RYDAPT 149, 150
RYKINDO 306
RYLAZE 307
S
sapropterin dihydrochloride oral packet . 308
sapropterin dihydrochloride oral tablet... 308
SCEMBLIX 149, 150
SECUADO 309
SELARSDI INTRAVENOUS 350, 351
SELARSDI SUBCUTANEOUS
SOLUTION 350, 351
SELARSDI SUBCUTANEOUS
SOLUTION PREFILLED SYRINGE 45
MG/0.5ML, 90 MG/ML 350, 351
SEROSTIM SUBCUTANEOUS
SOLUTION RECONSTITUTED 4 MG,
5 MG, 6 MG 310
SIGNIFOR 311
sildenafil citrate oral suspension
reconstituted 312
sildenafil citrate oral tablet 20 mg 312
SILIQ 313
SIMLANDI (1 PEN) SUBCUTANEOUS
AUTO-INJECTOR KIT 40 MG/0.4ML,
80 MG/0.8ML 141, 142
SIMLANDI (1 SYRINGE) 141, 142
SIMLANDI (2 PEN) 141, 142

SIMLANDI (2 SYRINGE)
SUBCUTANEOUS PREFILLED
SYRINGE KIT 20 MG/0.2ML, 40
MG/0.4ML 141, 142
SIMPONI SUBCUTANEOUS SOLUTION
AUTO-INJECTOR 314, 315
SIMPONI SUBCUTANEOUS SOLUTION
PREFILLED SYRINGE 314, 315
sirolimus oral solution 1 mg/ml 387
sirolimus oral tablet 0.5 mg, 1 mg, 2 mg 387
SIRTURO 316
SKYTROFA 217, 218
sodium oxybate 317
sodium phenylbutyrate oral powder 3 gm/tsp
..... 318
sodium phenylbutyrate oral tablet 318
sofosbuvir-velpatasvir 319
SOLTAMOX 149, 150
SOMAVERT 320
sorafenib tosylate 149, 150
SOTYKTU 321
STARJEMZA INTRAVENOUS 350, 351
STARJEMZA SUBCUTANEOUS
SOLUTION 350, 351
STARJEMZA SUBCUTANEOUS
SOLUTION PREFILLED SYRINGE 45
MG/0.5ML, 90 MG/ML 350, 351
STELARA INTRAVENOUS 350, 351
STELARA SUBCUTANEOUS
SOLUTION 45 MG/0.5ML 350, 351
STELARA SUBCUTANEOUS
SOLUTION PREFILLED SYRINGE 45
MG/0.5ML, 90 MG/ML 350, 351
STEQEYMA INTRAVENOUS 350, 351
STEQEYMA SUBCUTANEOUS
SOLUTION 350, 351
STEQEYMA SUBCUTANEOUS
SOLUTION PREFILLED SYRINGE 45
MG/0.5ML, 90 MG/ML 350, 351
STIVARGA 149, 150
SUCRAID 322
sunitinib malate 149, 150
SYMDEKO 323
SYNAREL 324

T

TABLOID 149, 150
 TABRECTA 149, 150
 tacrolimus er oral capsule extended release
 24 hour 0.5 mg, 1 mg, 5 mg..... 387
 tacrolimus intravenous solution 5 mg/ml 387
 tacrolimus oral capsule 0.5 mg, 1 mg, 5 mg
 387
 tadalafil (pah) 325
 tadalafil oral tablet 5 mg 326
 TADLIQ..... 325
 TAFINLAR..... 149, 150
 TAGRISSO 149, 150
 TALTZ SUBCUTANEOUS SOLUTION
 AUTO-INJECTOR..... 327, 328
 TALTZ SUBCUTANEOUS SOLUTION
 PREFILLED SYRINGE 20 MG/0.25ML,
 40 MG/0.5ML, 80 MG/ML 327, 328
 TALZENNA 149, 150
 TARPEYO 329
 TASCENSO ODT..... 256, 257
 tasimelteon 330
 TAVNEOS..... 331, 332
 TAZVERIK..... 149, 150
 temazepam 226
 TENIVAC INTRAMUSCULAR
 INJECTABLE 5-2 LFU (INJECTION)
 387
 TENIVAC INTRAMUSCULAR
 SUSPENSION 5-2 LF/0.5ML 387
 TEPEZZA 333
 TEPMETKO 149, 150
 teriflunomide 256, 257
 teriparatide subcutaneous solution pen-
 injector 560 mcg/2.24ml 334
 testosterone transdermal gel 1.62 %, 12.5
 mg/act (1%), 20.25 mg/1.25gm (1.62%),
 20.25 mg/act (1.62%), 25 mg/2.5gm
 (1%), 40.5 mg/2.5gm (1.62%), 50
 mg/5gm (1%) 340
 testosterone transdermal solution..... 340
 tetrabenazine 358
 THALOMID ORAL CAPSULE 100 MG,
 50 MG 149, 150

TIBSOVO 149, 150
 tigecycline 336
 tiopronin oral..... 335
 TOBI PODHALER..... 337
 tobramycin inhalation nebulization solution
 300 mg/4ml, 300 mg/5ml..... 387
 tofacitinib citrate er 369, 370
 tofacitinib citrate oral solution 369, 370
 tofacitinib citrate oral tablet 369, 370
 tolvaptan..... 338
 tolvaptan (hyponatremia) 338
 topiramate oral solution 25 mg/ml 198
 toremifene citrate 149, 150
 TRELSTAR MIXJECT 215
 TREMFYA ONE-PRESS
 SUBCUTANEOUS SOLUTION PEN-
 INJECTOR..... 342, 343
 TREMFYA PEN SUBCUTANEOUS
 SOLUTION AUTO-INJECTOR 100
 MG/ML, 200 MG/2ML 342, 343
 TREMFYA SUBCUTANEOUS
 SOLUTION PREFILLED SYRINGE 100
 MG/ML, 200 MG/2ML 342, 343
 TREMFYA-CD/UC INDUCTION. 342, 343
 tretinoin oral..... 149, 150
 trientine hcl oral capsule 250 mg 344
 TRIKAFTA..... 345
 TRULICITY SUBCUTANEOUS
 SOLUTION AUTO-INJECTOR 214
 TRUQAP ORAL TABLET 200 MG..... 149,
 150
 TRUQAP ORAL TABLET THERAPY
 PACK..... 149, 150
 TUKYSA 149, 150
 TURALIO ORAL CAPSULE 125 MG. 149,
 150
 TYMLOS 346
 TYVASO DPI MAINTENANCE KIT
 INHALATION POWDER 112 X 32MCG
 & 112 X64MCG, 112 X 48MCG & 112
 X64MCG, 112 X 48MCG & 112
 X80MCG, 16 MCG, 32 MCG, 48 MCG,
 64 MCG, 80 MCG 347, 348

TYVASO DPI TITRATION KIT
 INHALATION POWDER 16 & 32 & 48
 MCG 347, 348

U

UBRELVY 169, 170
 UPTRAVI ORAL 349
 UPTRAVI TITRATION 349
 ustekinumab subcutaneous solution 350, 351
 ustekinumab subcutaneous solution prefilled
 syringe 45 mg/0.5ml, 90 mg/ml.. 350, 351
 ustekinumab-aauz subcutaneous solution
 prefilled syringe 45 mg/0.5ml, 90 mg/ml
 350, 351
 ustekinumab-aekn subcutaneous solution
 prefilled syringe 45 mg/0.5ml, 90 mg/ml
 350, 351

UZEDY SUBCUTANEOUS SUSPENSION
 PREFILLED SYRINGE 100
 MG/0.28ML, 125 MG/0.35ML, 150
 MG/0.42ML, 200 MG/0.56ML, 250
 MG/0.7ML, 50 MG/0.14ML, 75
 MG/0.21ML 352

V

VALCHLOR 353
 VALTOCO 10 MG DOSE 262
 VALTOCO 15 MG DOSE NASAL
 LIQUID THERAPY PACK 2 X 7.5
 MG/0.1ML 262
 VALTOCO 20 MG DOSE NASAL
 LIQUID THERAPY PACK 2 X 10
 MG/0.1ML 262
 VALTOCO 5 MG DOSE 262
 VANFLYTA 149, 150
 VEMLIDY 354
 VENCLEXTA 149, 150
 VENCLEXTA STARTING PACK 149, 150
 VEOZAH 355
 VERZENIO 149, 150
 vigabatrin 356
 VIGAFYDE 356
 VIJOICE 357
 VITRAKVI 149, 150
 VIZIMPRO 149, 150
 VONJO 149, 150

VORANIGO 149, 150
 voriconazole intravenous 359
 VOSEVI 360
 VOWST 361
 VRAYLAR ORAL CAPSULE 272, 273

W

WEGOVY ORAL 364, 365
 WEGOVY SUBCUTANEOUS SOLUTION
 AUTO-INJECTOR 0.25 MG/0.5ML, 0.5
 MG/0.5ML, 1 MG/0.5ML, 1.7
 MG/0.75ML, 2.4 MG/0.75ML ... 362, 363
 WELIREG 149, 150
 WINREVAIR 367

X

XALKORI 149, 150
 XATMEP 251
 XDEMVY 368
 XELJANZ ORAL SOLUTION 369, 370
 XELJANZ ORAL TABLET 369, 370
 XELJANZ XR 369, 370
 XERMELO 371
 XIFAXAN 372
 XOLAIR 373, 374
 XOLREMDI 375, 376
 XOSPATA 149, 150
 XPOVIO (100 MG ONCE WEEKLY)
 ORAL TABLET THERAPY PACK 50
 MG 149, 150
 XPOVIO (40 MG ONCE WEEKLY) ORAL
 TABLET THERAPY PACK 10 MG, 40
 MG 149, 150
 XPOVIO (40 MG TWICE WEEKLY)
 ORAL TABLET THERAPY PACK 40
 MG 149, 150
 XPOVIO (60 MG ONCE WEEKLY) ORAL
 TABLET THERAPY PACK 60 MG. 149,
 150
 XPOVIO (60 MG TWICE WEEKLY).. 149,
 150
 XPOVIO (80 MG ONCE WEEKLY) ORAL
 TABLET THERAPY PACK 40 MG, 80
 MG 149, 150
 XPOVIO (80 MG TWICE WEEKLY).. 149,
 150

XTANDI	150	ZEMAIRA	145
XYWAV	377	ZEPCOUND KWIKPEN.....	380, 381
Y		ZEPCOUND SUBCUTANEOUS	
YESINTEK INTRAVENOUS.....	350, 351	SOLUTION AUTO-INJECTOR	380, 381
YESINTEK SUBCUTANEOUS		ZEPOSIA	382
SOLUTION.....	350, 351	ZEPOSIA 7-DAY STARTER PACK.....	382
YESINTEK SUBCUTANEOUS		ZEPOSIA STARTER KIT ORAL	
SOLUTION PREFILLED SYRINGE 45		CAPSULE THERAPY PACK 0.23MG	
MG/0.5ML, 90 MG/ML	350, 351	&0.46MG 0.92MG(21).....	382
YONSA.....	150	ZILBRYSQ.....	383
YORVIPATH	378	ZOLINZA	150
YUTREPIA.....	379	zolpidem tartrate er	226
Z		zolpidem tartrate oral tablet 10 mg	226
zaleplon oral capsule 10 mg, 5 mg.....	226	ZTALMY	384
ZARXIO	366	ZTLIDO	341
ZAVZPRET	169, 170	ZURZUVAE	385
ZEJULA ORAL TABLET.....	150	ZYDELIG	150
ZELBORAF	150	ZYKADIA ORAL TABLET	150

2026 Troy Medicare

2026 Step Therapy Criteria

CURRENT AS OF 08/01/2026

anticonvulsant step therapy

Products Affected

- *levetiracetam tablet disintegrating soluble 250 mg oral*
- *levetiracetam tablet disintegrating soluble 500 mg oral*
- *perampanel suspension 0.5 mg/ml oral*
- *perampanel tablet 10 mg oral*
- *perampanel tablet 12 mg oral*
- *perampanel tablet 2 mg oral*
- *perampanel tablet 4 mg oral*
- *perampanel tablet 6 mg oral*
- *perampanel tablet 8 mg oral*
- RELGAABI CAPSULE 200 MG ORAL
- RELGAABI CAPSULE 300 MG ORAL
- RELGAABI CAPSULE 400 MG ORAL
- SPRITAM TABLET DISINTEGRATING SOLUBLE 250 MG ORAL
- SPRITAM TABLET DISINTEGRATING SOLUBLE 500 MG ORAL
- SYMPAZAN FILM 10 MG ORAL
- SYMPAZAN FILM 20 MG ORAL
- SYMPAZAN FILM 5 MG ORAL
- XCOPRI (250 MG DAILY DOSE) TABLET THERAPY PACK 100 & 150 MG ORAL
- XCOPRI (350 MG DAILY DOSE) TABLET THERAPY PACK 150 & 200 MG ORAL
- XCOPRI TABLET 100 MG ORAL
- XCOPRI TABLET 150 MG ORAL
- XCOPRI TABLET 200 MG ORAL
- XCOPRI TABLET 25 MG ORAL
- XCOPRI TABLET 50 MG ORAL
- XCOPRI TABLET THERAPY PACK 14 X 12.5 MG & 14 X 25 MG ORAL
- XCOPRI TABLET THERAPY PACK 14 X 150 MG & 14 X 200 MG ORAL
- XCOPRI TABLET THERAPY PACK 14 X 50 MG & 14 X 100 MG ORAL
- ZONISADE SUSPENSION 100 MG/5ML ORAL

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of two generic anticonvulsants. Step 2: Once two generic anticonvulsants have been tried, failed, or contraindicated patients can receive therapy with Spritam, Sympazan, Xcopri, generic perampanel, generic levetiracetam tablet for oral suspension, Relgaabi, or Zonisade oral solution.
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antidepressant step therapy

Products Affected

- EXXUA TABLET EXTENDED RELEASE 24 HOUR 18.2 MG ORAL
- EXXUA TABLET EXTENDED RELEASE 24 HOUR 36.3 MG ORAL
- EXXUA TABLET EXTENDED RELEASE 24 HOUR 54.5 MG ORAL
- EXXUA TABLET EXTENDED RELEASE 24 HOUR 72.6 MG ORAL
- EXXUA TITRATION PACK TABLET EXTENDED RELEASE 24 HOUR 18.2 MG ORAL
- FETZIMA CAPSULE EXTENDED RELEASE 24 HOUR 120 MG ORAL
- FETZIMA CAPSULE EXTENDED RELEASE 24 HOUR 20 MG ORAL
- FETZIMA CAPSULE EXTENDED RELEASE 24 HOUR 40 MG ORAL
- FETZIMA CAPSULE EXTENDED RELEASE 24 HOUR 80 MG ORAL
- FETZIMA TITRATION CAPSULE ER 24 HOUR THERAPY PACK 20 & 40 MG ORAL

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of two generic antidepressants. Step 2: Once two generic antidepressants have been tried, failed, or contraindicated patient can receive therapy with Fetzima or Exxua.
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brinzolamide and dorzolamide-timolol PF step therapy

Products Affected

- *brinzolamide suspension 1 % ophthalmic*
- *dorzolamide hcl-timolol mal pf solution 2-0.5 % ophthalmic*

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of formulary dorzolamide or dorzolamide/timolol ophthalmic solution. Step 2: Once dorzolamide or dorzolamide/timolol ophthalmic solution has been tried, failed, or contraindicated the patient can receive therapy with brinzolamide or dorzolamide-timolol PF ophthalmic Solution.
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drizalma step therapy

Products Affected

- DRIZALMA SPRINKLE CAPSULE
DELAYED RELEASE SPRINKLE 20
MG ORAL
- DRIZALMA SPRINKLE CAPSULE
DELAYED RELEASE SPRINKLE 30
MG ORAL
- DRIZALMA SPRINKLE CAPSULE
DELAYED RELEASE SPRINKLE 40
MG ORAL
- DRIZALMA SPRINKLE CAPSULE
DELAYED RELEASE SPRINKLE 60
MG ORAL

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of generic formulary duloxetine. Step 2: Once generic formulary duloxetine has been tried, failed, or contraindicated the patient can receive therapy with drizalma.
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febuxostat step therapy

Products Affected

- febuxostat tablet 40 mg oral*
- febuxostat tablet 80 mg oral*

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of allopurinol tablet. Step 2: Once allopurinol tablet has been tried, failed, or contraindicated patients can receive therapy with Febuxostat.
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netarsudil step therapy

Products Affected

- RHOPRESSA SOLUTION 0.02 %
OPHTHALMIC
- ROCKLATAN SOLUTION 0.02-0.005 %
OPHTHALMIC

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of latanoprost or travoprost. Step 2: Once latanoprost or travoprost has been tried, failed, or contraindicated patients can receive therapy with Rhopressa or Rocklatan.
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ongentys step therapy

Products Affected

- ONGENTYS CAPSULE 25 MG ORAL
- ONGENTYS CAPSULE 50 MG ORAL

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of entacapone or carbidopa-levodopa-entacapone. Step 2: Once entacapone or carbidopa-levodopa-entacapone has been tried, failed, or contraindicated patients can receive therapy with Ongentys.
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savella step therapy

Products Affected

- SAVELLA TITRATION PACK 12.5 & 25 & 50 MG ORAL

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication to duloxetine, milnacipran or pregabalin. Step 2: Once duloxetine, milnacipran or pregabalin has been tried, failed or contraindicated patients can receive therapy with Savella.
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topical immunomodulators step therapy

Products Affected

- *pimecrolimus cream 1 % external*
- *tacrolimus ointment 0.03 % external*
- *tacrolimus ointment 0.1 % external*

Details

Criteria	Step 1: First line therapy should be a documented trial, failure, or contraindication of two topical corticosteroids. Step 2: Once two topical corticosteroids have been tried, failed, or contraindicated patients can receive therapy with generic pimecrolimus or generic topical tacrolimus.
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urinary incontinence agents step therapy

Products Affected

- darifenacin hydrobromide er tablet extended release 24 hour 15 mg oral*
- darifenacin hydrobromide er tablet extended release 24 hour 7.5 mg oral*
- trospium chloride er capsule extended release 24 hour 60 mg oral*

Details

Criteria	Step 1: First line therapy should be a documented trial, failure or contraindication of 2 of the following: oxybutynin, oxybutynin ER, trospium, tolterodine, tolterodine ER, fesoterodine ER, or solifenacin. Step 2: Once two of the medications listed in Step 1 have been tried, failed, or contraindicated, patients can receive therapy with trospium ER or darifenacin ER
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Index

B

brinzolamide suspension 1 % ophthalmic 402

D

darifenacin hydrobromide er tablet extended
release 24 hour 15 mg oral 409

darifenacin hydrobromide er tablet extended
release 24 hour 7.5 mg oral 409

dorzolamide hcl-timolol mal pf solution 2-
0.5 % ophthalmic 402

DRIZALMA SPRINKLE CAPSULE

DELAYED RELEASE SPRINKLE 20
MG ORAL 403

DRIZALMA SPRINKLE CAPSULE

DELAYED RELEASE SPRINKLE 30
MG ORAL 403

DRIZALMA SPRINKLE CAPSULE

DELAYED RELEASE SPRINKLE 40
MG ORAL 403

DRIZALMA SPRINKLE CAPSULE

DELAYED RELEASE SPRINKLE 60
MG ORAL 403

E

EXXUA TABLET EXTENDED RELEASE

24 HOUR 18.2 MG ORAL 401

EXXUA TABLET EXTENDED RELEASE

24 HOUR 36.3 MG ORAL 401

EXXUA TABLET EXTENDED RELEASE

24 HOUR 54.5 MG ORAL 401

EXXUA TABLET EXTENDED RELEASE

24 HOUR 72.6 MG ORAL 401

EXXUA TITRATION PACK TABLET

EXTENDED RELEASE 24 HOUR 18.2
MG ORAL 401

F

febuxostat tablet 40 mg oral 404

febuxostat tablet 80 mg oral 404

FETZIMA CAPSULE EXTENDED

RELEASE 24 HOUR 120 MG ORAL 401

FETZIMA CAPSULE EXTENDED

RELEASE 24 HOUR 20 MG ORAL . 401

FETZIMA CAPSULE EXTENDED

RELEASE 24 HOUR 40 MG ORAL . 401

FETZIMA CAPSULE EXTENDED

RELEASE 24 HOUR 80 MG ORAL . 401

FETZIMA TITRATION CAPSULE ER 24

HOUR THERAPY PACK 20 & 40 MG

ORAL 401

L

levetiracetam tablet disintegrating soluble

250 mg oral 400

levetiracetam tablet disintegrating soluble

500 mg oral 400

O

ONGENTYS CAPSULE 25 MG ORAL 406

ONGENTYS CAPSULE 50 MG ORAL 406

P

perampanel suspension 0.5 mg/ml oral... 400

perampanel tablet 10 mg oral 400

perampanel tablet 12 mg oral 400

perampanel tablet 2 mg oral 400

perampanel tablet 4 mg oral 400

perampanel tablet 6 mg oral 400

perampanel tablet 8 mg oral 400

pimecrolimus cream 1 % external 408

R

RELGAABI CAPSULE 200 MG ORAL 400

RELGAABI CAPSULE 300 MG ORAL 400

RELGAABI CAPSULE 400 MG ORAL 400

RHOPRESSA SOLUTION 0.02 %

OPHTHALMIC 405

ROCKLATAN SOLUTION 0.02-0.005 %

OPHTHALMIC 405

S

SAVELLA TITRATION PACK 12.5 & 25

& 50 MG ORAL 407

SPRITAM TABLET DISINTEGRATING

SOLUBLE 250 MG ORAL 400

SPRITAM TABLET DISINTEGRATING

SOLUBLE 500 MG ORAL 400

SYMPAZAN FILM 10 MG ORAL 400

SYMPAZAN FILM 20 MG ORAL 400

SYMPAZAN FILM 5 MG ORAL 400

T

tacrolimus ointment 0.03 % external 408

tacrolimus ointment 0.1 % external 408
trospium chloride er capsule extended
release 24 hour 60 mg oral..... 409

X

XCOPRI (250 MG DAILY DOSE)
TABLET THERAPY PACK 100 & 150
MG ORAL 400
XCOPRI (350 MG DAILY DOSE)
TABLET THERAPY PACK 150 & 200
MG ORAL 400
XCOPRI TABLET 100 MG ORAL 400
XCOPRI TABLET 150 MG ORAL 400

XCOPRI TABLET 200 MG ORAL 400
XCOPRI TABLET 25 MG ORAL 400
XCOPRI TABLET 50 MG ORAL 400
XCOPRI TABLET THERAPY PACK 14 X
12.5 MG & 14 X 25 MG ORAL..... 400
XCOPRI TABLET THERAPY PACK 14 X
150 MG & 14 X200 MG ORAL..... 400
XCOPRI TABLET THERAPY PACK 14 X
50 MG & 14 X100 MG ORAL..... 400

Z

ZONISADE SUSPENSION 100 MG/5ML
ORAL..... 400



**Troy Medicare for Dual-eligible Beneficiaries (HMO D-SNP)/Troy
Medicare (HMO)**

2026 Formulary

List of Covered Drugs

Formulary ID#: 26314

PLEASE READ: THIS DOCUMENT CONTAINS INFORMATION
ABOUT THE DRUGS WE COVER IN THIS PLAN

This formulary was updated on 7/31/2026. For more recent information or other questions, please contact Pharmacy Member Service at 1-866-423-8065 (TTY users should call 711), Monday through Sunday, 24 hours a day, or visit <http://www.troymedicare.com>.

Important Message About What You Pay for Vaccines - Our plan covers most Part D vaccines at no cost to you. Call Member Services for more information.

Important Message About What You Pay for Insulin - You won't pay more than \$35 for a one-month supply of each insulin product covered by our plan, no matter what cost-sharing tier it's on. You won't pay more than \$10 for a one-month supply of generic insulin products covered by our plan on Tier 1.