

Prescription Benefit Medication Prior Authorization Criteria

QuartzBenefits.com

These criteria apply to drugs picked up at the pharmacy.

These medication prior authorization criteria do not apply to drugs picked up at the pharmacy for State and Local Government members or BadgerCare Plus and/or Medicaid SSI members.

State and Local Government members should call **Navitus** at **(866)** 333-2757 or visit <u>navitus.com</u> for information about your prescription drug benefits.

Quartz BadgerCare Plus and/or Medicaid SSI members must call the **Wisconsin Department of Health and Family Services** at **(800)** 362-3002 or visit <u>forwardhealth.wi.gov</u> for information about your prescription drug benefits.



May 1, 2024 Pharmacy Benefit Drug Prior Authorization Criteria

A medication prior authorization request may be started by members, providers, or designated representatives by fax, electronically on Quartz's website, telephone, mail. Or, for medical benefit medications, also by Health Link, Plan Link, MyQuartzTools, or electronic prior authorization (e-PA) within the electronic medical record. Electronic (e-PA) via Surescripts verifies member eligibility and member benefit information. Quartz sends back e-PA criteria questions to the provider staff which can be answered, and medical records can be attached to the request.

Quartz strongly recommends that the health care provider initiate the prior authorization request process on behalf of the member. This is because the health care provider will be able to include the medical history necessary for a timely decision to be made based on all of the relevant information, including any case specific circumstances that can be considered. Once a request and the supporting documentation have been submitted, a pharmacist or appropriate staff review the prior authorization criteria and exception requirements separately to make a coverage decision.

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Actemra (tocilizumab)				
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Prior Authorization Guideline

Guideline ID	GL-144780
Guideline Name	Actemra (tocilizumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Actemra				
Diagnosis		Moderate to Severely Active Rheumatoid Arthritis		
Approval L	Approval Length 12 month(s)			
Guideline Type Prior Authorization - IL and MN Plans				
Product Name	Generic Na	me	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML		6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML		6650007000E520	Brand

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

- **3.1.2** Trial and failure, contraindication or intolerance to TWO of the following:
 - adalimumab
 - certolizumab
 - etanercept
 - golimumab
 - tofacitinib (ER)
 - upadacitinib

OR

**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immunodeficiency syndromes, bone marrow hyperplasia, leukopenia, thrombocytopenia or significant anemia, or hypersensitivity to methotrexate
significant anemia, or hypersensitivity to methotrexate.

Product Name: Actemra	
Diagnosis Moderate to Severely Active Rheumatoid Arthritis	
Approval Length	12/31/2039
Guideline Type Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

- **3.1.2** Trial and failure, contraindication or intolerance to TWO of the following:
 - adalimumab
 - certolizumab
 - etanercept
 - golimumab
 - tofacitinib (ER)
 - upadacitinib

OR

3.2 Continuation of prior therapy with tocilizumab, verified by paid claims or medical records (e.g. chart notes)

**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immunodeficiency syndromes, bone marrow hyp erplasia, leukopenia, thrombocytopenia or
significant anemia, or hypersensitivity to methotrexate.

Product Name: Actemra	
Diagnosis Systemic Juvenile Idiopathic Arthritis (SJIA)	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

Approval Criteria

1 - Member is 2 years of age or older

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication** or intolerance to ONE of the following for 3 months:
 - corticosteroids
 - methotrexate
 - nonsteroidal anti-inflammatories

OR

**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immunodeficiency syndromes, bone marrow hyperplasia, leu kopenia, thrombocytopenia or significant
anemia, or hypersensitivity to methotrexate

Product Name: Actemra	
Diagnosis Systemic Juvenile Idiopathic Arthritis (SJIA)	
Approval Length 12 month(s)	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

1 - Member is 2 years of age or older

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication** or intolerance to ONE of the following for 3 months:
 - corticosteroids
 - methotrexate
 - nonsteroidal anti-inflammatories

OR

Notes	**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant anemia, or hypersensitivity to methotrexate
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Product Name: Actemra	
Diagnosis Polyarticular Juvenile Idiopathic Arthritis (PJIA)	
Approval Length	12/31/2039
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

1 - Diagnosis of polyarticular juvenile idiopathic arthritis (PJIA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

- **3.1.2** Trial and failure, contraindication or intolerance to TWO of the following:
 - adalimumab
 - etanercept
 - tofacitinib

OR

**Absolute contraindications to methotrexate are pregnancy, nursing,
alcoholism, alcoholic liver disease or other
chronic liver disease, immunodeficiency syndromes, bone marrow hyp

erplasia, leukopenia, thrombocytopenia or
significant anemia, or hypersensitivity to methotrexate.

Product Name: Actemra			
Diagnosis Polyarticular Juvenile Idiopathic Arthritis (PJIA)			
Approval Length 12 month(s)			
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

1 - Diagnosis of polyarticular juvenile idiopathic arthritis (PJIA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

- **3.1.2** Trial and failure, contraindication or intolerance to TWO of the following:
 - adalimumab
 - etanercept
 - tofacitinib

OR

3.2 Continuation of prior therapy with tocilizumab, verified by paid claims or medical records (e.g. chart notes)

**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immunodeficiency syndromes, bone marrow hyperplasia, leukopenia, thrombocytopenia or
significant anemia, or hypersensitivity to methotrexate.

Product Name: Actemra			
Diagnosis Giant Cell Arteritis (GCA)			
Approval Length 12/31/2039			
Guideline Type Prior Authorization - All Plans Except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

Approval Criteria

1 - Diagnosis of Giant Cell Arteritis (GCA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** ONE of the following:
- **3.1** One of the following:
 - Symptoms relapsed despite use of corticosteroids or methotrexate
 - Contraindication** to methotrexate
 - Inability to taper corticosteroids

OR

3.2 Continuation of prior therapy with tocilizumab, verified by a paid claims or medical records (e.g., chart notes)

**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immunodeficiency syndromes, bone marrow hyperplasia, leu kopenia, thrombocytopenia or significant anemia, or hypersensitivity to methotrexate
anemia, or hypersensitivity to methotrexate

Product Name: Actemra			
Diagnosis Giant Cell Arteritis (GCA)			
Approval Length 12 month(s)			
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

Approval Criteria

1 - Diagnosis of Giant Cell Arteritis (GCA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** ONE of the following:
- **3.1** One of the following:
 - Symptoms relapsed despite use of corticosteroids or methotrexate
 - Contraindication** to methotrexate
 - Inability to taper corticosteroids

OR

**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant
anemia, or hypersensitivity to methotrexate

Product Name: Actemra			
Diagnosis Systemic Sclerosis - Associated Interstitial Lung Disease (SSc-ILD)			
Approval Length 12/31/2039			
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

1 - Diagnosis of Systemic Sclerosis – Associated Interstitial Lung Disease (SSc-ILD)

AND

- 2 One of the following:
- **2.1** Submission of medical records (e.g., chart notes), documenting one of the following:
- **2.1.1** Decline in pulmonary function despite use of one of the following standard treatments:
 - mycophenolate
 - cyclophosphamide
 - azathioprine

OR

- **2.1.2** Contraindication to one of the following standard agents:
 - mycophenolate
 - cyclophosphamide
 - azathioprine

OR

2.2 Continuation of prior therapy with tocilizumab, verified by a paid claims or medical records (e.g., chart notes)

AND

- **3** Prescribed by or in consultation with one of the following:
 - rheumatologist
 - pulmonologist

Product Name: Actemra	
Diagnosis	Systemic Sclerosis - Associated Interstitial Lung Disease (SSc-ILD)

Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

1 - Diagnosis of Systemic Sclerosis – Associated Interstitial Lung Disease (SSc-ILD)

AND

- 2 One of the following:
- **2.1** Submission of medical records (e.g., chart notes), documenting one of the following:
- **2.1.1** Decline in pulmonary function despite use of one of the following standard treatments:
 - mycophenolate
 - cyclophosphamide
 - azathioprine

OR

- **2.1.2** Contraindication to one of the following standard agents:
 - mycophenolate
 - cyclophosphamide
 - azathioprine

OR

AND

- **3** Prescribed by or in consultation with one of the following:
 - rheumatologist
 - pulmonologist

Product Name: Actemra	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis
Approval Length	12/31/2039
Guideline Type	Quantity Exception - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

Approval Criteria

1 - Failure of an adherent 3-month trial of standard maintenance dosing with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)

OR

2 - There is a confirmed history of an approved quantity limit exception (via prior authorization, historical authorization on file or submission of medical records) for tocilizumab

Product Name: Actemra	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis
Approval Length	12 month(s)
Guideline Type	Quantity Exception - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ACTEMRA ACTPEN	TOCILIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 162 MG/0.9ML	6650007000D520	Brand
ACTEMRA	TOCILIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 162 MG/0.9ML	6650007000E520	Brand

1 - Failure of an adherent 3-month trial of standard maintenance dosing with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)

OR

2 - There is a confirmed history of an approved quantity limit exception (vial prior authorization, historical authorization on file or submission of medical records) for tocilizumab

2. Definitions

Definition	Description
Steroid Dependence:	Demonstrated steroid dependence (defined as equivalent to prednisone 10mg daily for >3 months) with the inability to taper or when tapering of dose leads to loss of symptom control

3. Revision History

Date	Notes
4/16/2024	Update guideline

Actiq (Fentanyl)
The National Annual Control Control State Control Con

Prior Authorization Guideline

Guideline ID	GL-129620
Guideline Name	Actiq (Fentanyl)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Fentanyl	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 200 MCG	65100025108450	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 400 MCG	65100025108455	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 600 MCG	65100025108460	Generic

FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 800 MCG	65100025108465	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 1200 MCG	65100025108475	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 1600 MCG	65100025108485	Generic

1 - All of the followi	ng
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1.1 Prescribed by, or in consultation with, an Oncologist or specialty in Pain Management

AND

1.2 Medication is limited to the treatment of breakthrough cancer pain

AND

- **1.3** Person is already tolerant to opioids, defined as:
- **1.3.1** oral morphine 60mg daily for one week

OR

1.3.2 transdermal fentanyl 25mcg/hr for one week

OR

1.3.3 oxycodone 30mg daily for one week

OR

1.3.4 oral hydromorphone 8mg daily for one week

OR

1.3.5 equianalgesic dose of another opioid for at least one week

AND

- 1.4 Person has failed an adequate trial of one of the following:
- **1.4.1** immediate release oxycodone

OR

1.4.2 immediate release oral hydromorphone

OR

1.4.3 immediate release morphine

OR

2 - (Minnesota plans only) – person has stage four metastatic cancer and the requested drug is being used to treat cancer-related pain

Product Name: Fentanyl	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 200 MCG	65100025108450	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 400 MCG	65100025108455	Generic

FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 600 MCG	65100025108460	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 800 MCG	65100025108465	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 1200 MCG	65100025108475	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 1600 MCG	65100025108485	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Fentanyl	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 200 MCG	65100025108450	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 400 MCG	65100025108455	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 600 MCG	65100025108460	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 800 MCG	65100025108465	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 1200 MCG	65100025108475	Generic
FENTANYL CITRATE ORAL TRANSMUCOSAL	FENTANYL CITRATE LOZENGE ON A HANDLE 1600 MCG	65100025108485	Generic

Approval Criteria

1 - Prescribed by, or in consultation with, an Oncologist or specialty in Pain Management
AND
2 - Medication is limited to the treatment of breakthrough cancer pain
AND
3 - Person is already tolerant to opioids, defined as:
3.1 oral morphine 60mg daily for one week
OR
3.2 transdermal fentanyl 25mcg/hr for one week
OR
3.3 oxycodone 30mg daily for one week
OR
3.4 oral hydromorphone 8mg daily for one week
OR
3.5 equianalgesic dose of another opioid for at least one week
AND
4 - Person has failed an adequate trial of one of the following:
4.1 immediate release oxycodone

OR

4.2 immediate release oral hydromorphone

OR

4.3 immediate release morphine

2. Revision History

Date	Notes
11/6/2023	New Program

Actonel (risedronate)
The Section of the Control

Prior Authorization Guideline

Guideline ID	GL-129870	
Guideline Name	Actonel (risedronate)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: risedronate 5 mg	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
RISEDRONATE SODIUM	RISEDRONATE SODIUM TAB 5 MG	30042065100305	Generic

Approval Criteria

1 - One of the following:

1.1 Both of the following:

- Diagnosis of osteoporosis due to corticosteroid use
- Trial and failure, contraindication, or intolerance to alendronate

OR

- **1.2** For diagnoses other than osteoporosis due to corticosteroid use, trial and failure, contraindication, or intolerance to ALL of the following:
 - alendronate
 - ibandronate
 - other strengths of risedronate (i.e., 35 mg, 150 mg)

Product Name: risedronate 5 mg	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
RISEDRONATE SODIUM	RISEDRONATE SODIUM TAB 5 MG	30042065100305	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: risedronate 5 mg	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
RISEDRONATE SODIUM	RISEDRONATE SODIUM TAB 5 MG	30042065100305	Generic

- 1 One of the following:
- **1.1** Both of the following:
 - Diagnosis of osteoporosis due to corticosteroid use
 - Trial and failure, contraindication, or intolerance to alendronate

OR

- **1.2** For diagnoses other than osteoporosis due to corticosteroid use, trial and failure, contraindication, or intolerance to ALL of the following:
 - alendronate
 - ibandronate
 - other strengths of risedronate (i.e., 35 mg, 150 mg)

2. Revision History

Date	Notes
10/12/2023	2024 New Implementation

Acute Migraine Treatments				
The second secon				

Prior Authorization Guideline

Guideline ID	GL-127880	
Guideline Name	Acute Migraine Treatments	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Frovatriptan, Brand Reyvow		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FROVATRIPTAN SUCCINATE	FROVATRIPTAN SUCCINATE TAB 2.5 MG (BASE EQUIVALENT)	67406030100320	Generic
REYVOW	LASMIDITAN SUCCINATE TAB 50 MG	67406540600310	Brand
REYVOW	LASMIDITAN SUCCINATE TAB 100 MG	67406540600320	Brand

Approval Criteria

- 1 Trial and failure of at least 2 of the following:
 - sumatriptan
 - naratriptan
 - rizatriptan
 - eletriptan
 - zolmitriptan
 - almotriptan
 - frovatriptan (not required for request for frovatriptan)

OR

2 - If the member has contraindication to triptan, than trial and failure of 2 non-triptan, prescription strength analgesics that are listed as effective for treatment of migraines by the American Headache Society treatment guidelines is required (ex. nonsteroidal anti-inflammatory drugs (NSAIDs), ergotamine derivatives, etc.)

Product Name: Generic Frovatriptan, Brand Reyvow		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FROVATRIPTAN SUCCINATE	FROVATRIPTAN SUCCINATE TAB 2.5 MG (BASE EQUIVALENT)	67406030100320	Generic
REYVOW	LASMIDITAN SUCCINATE TAB 50 MG	67406540600310	Brand
REYVOW	LASMIDITAN SUCCINATE TAB 100 MG	67406540600320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Generic Frovatriptan, Brand Reyvow		
Approval Length	12/31/2039	
Guideline Type	Step Therapy - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FROVATRIPTAN SUCCINATE	FROVATRIPTAN SUCCINATE TAB 2.5 MG (BASE EQUIVALENT)	67406030100320	Generic
REYVOW	LASMIDITAN SUCCINATE TAB 50 MG	67406540600310	Brand
REYVOW	LASMIDITAN SUCCINATE TAB 100 MG	67406540600320	Brand

- 1 Trial and failure of at least 2 of the following:
 - sumatriptan
 - naratriptan
 - rizatriptan
 - eletriptan
 - zolmitriptan
 - almotriptan
 - frovatriptan (not required for request for frovatriptan)

OR

2 - If the member has contraindication to triptan, than trial and failure of 2 non-triptan, prescription strength analysesics that are listed as effective for treatment of migraines by the American Headache Society treatment guidelines is required (ex. nonsteroidal anti-inflammatory drugs (NSAIDs), ergotamine derivatives, etc.)

Product Name: Generic Frovatriptan, Brand Reyvow		
Approval Length	12/31/2039	
Guideline Type	Quantity Limits - All Plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
FROVATRIPTAN SUCCINATE	FROVATRIPTAN SUCCINATE TAB 2.5 MG (BASE EQUIVALENT)	67406030100320	Generic
REYVOW	LASMIDITAN SUCCINATE TAB 50 MG	67406540600310	Brand
REYVOW	LASMIDITAN SUCCINATE TAB 100 MG	67406540600320	Brand

Approval Criteria

1 - Member has greater than or equal to 2 migraine headaches per week

AND

2 - Member is on migraine headache prophylaxis treatment

Product Name: Generic Frovatriptan, Brand Reyvow		
Approval Length 12 month(s)		
Guideline Type Quantity Limits - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
FROVATRIPTAN SUCCINATE	FROVATRIPTAN SUCCINATE TAB 2.5 MG (BASE EQUIVALENT)	67406030100320	Generic
REYVOW	LASMIDITAN SUCCINATE TAB 50 MG	67406540600310	Brand
REYVOW	LASMIDITAN SUCCINATE TAB 100 MG	67406540600320	Brand

Approval Criteria

1 - Member has greater than or equal to 2 migraine headaches per week

AND

2 - Member is on migraine headache prophylaxis treatment

2. Revision History

Date	Notes
8/25/2023	New Program

Aczone (dapsone)	
(a) The behavioring most bridging in The Gray to be been associated, or about the photo day associate activation.	_

Prior Authorization Guideline

Guideline ID	GL-128132
Guideline Name Aczone (dapsone)	
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Dapsone 5%		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
DAPSONE	DAPSONE GEL 5%	90051015004020	Generic

Approval Criteria

- 1 Trial and failure of two different prior treatments for acne
 - tretinoin 0.01% gel, 0.025% cream/gel, 0.05% cream/gel, 0.1% cream

- adapalene (0.1% gel/cream, 0.3% gel)
- azelaic acid
- tazarotene
- oral minocycline
- oral doxycycline
- clindamycin 1% gel
- clindamycin 1.2%/benzoyl peroxide 5% gel
- erythromycin 2% topical

Product Name: Generic Dapsone 7.5%		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Step Therapy - IL and MN Plans		

Product Name			Brand/Generic
DAPSONE	DAPSONE GEL 7.5%	90051015004030	Generic

- 1 Trial and failure of two different prior treatments for acne
 - tretinoin 0.01% gel, 0.025% cream/gel, 0.05% cream/gel, 0.1% cream
 - adapalene (0.1% gel/cream, 0.3% gel)
 - azelaic acid
 - tazarotene
 - oral minocycline
 - oral doxycycline
 - clindamycin 1% gel
 - clindamycin 1.2%/benzoyl peroxide 5% gel
 - erythromycin 2% topical

AND

2 - Trial and failure of generic dapsone 5%

Product Name: Generic Dapsone 5%, Generic Dapsone 7.5%		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		

Guideline Type		Step Therapy - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
DAPSONE	DAPSONE GEL 5%		90051015004020	Generic
DAPSONE	DAPSONE G	EL 7.5%	90051015004030	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Generic Dapsone 5%				
Approval Length		12/31/2039		
Guideline Type		Step Therapy - All plans except IL and MN Plans		
Product Generic Na Name		me	GPI	Brand/Generic

Approval Criteria

DAPSONE

- 1 Trial and failure of two different prior treatments for acne
 - tretinoin 0.01% gel, 0.025% cream/gel, 0.05% cream/gel, 0.1% cream
 - adapalene (0.1% gel/cream, 0.3% gel)

DAPSONE GEL 5%

- azelaic acid
- tazarotene
- oral minocycline
- oral doxycycline
- clindamycin 1% gel
- clindamycin 1.2%/benzoyl peroxide 5% gel
- erythromycin 2% topical

Product Name: Generic Dapsone 7.5%		
Approval Length 12/31/2039		
Guideline Type Step Therapy - All plans except IL and MN Plans		

Generic

90051015004020

Product Name	Generic Name	GPI	Brand/Generic
DAPSONE	DAPSONE GEL 7.5%	90051015004030	Generic

- 1 Trial and failure of two different prior treatments for acne
 - tretinoin 0.01% gel, 0.025% cream/gel, 0.05% cream/gel, 0.1% cream
 - adapalene (0.1% gel/cream, 0.3% gel)
 - azelaic acid
 - tazarotene
 - oral minocycline
 - oral doxycycline
 - clindamycin 1% gel
 - clindamycin 1.2%/benzoyl peroxide 5% gel
 - erythromycin 2% topical

AND

2 - Trial and failure of generic dapsone 5%

2. Revision History

Date	Notes
8/25/2023	New Program

Adalimumab biosimilars			
(g) below required to depths to below to secure a state out that the depths could define the secure of the secure			

Prior Authorization Guideline

Guideline ID	GL-145182
Guideline Name	Adalimumab biosimilars
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz				
Diagnosis		Plaque Psoriasis		
Approval Length		12 Month(s)*		
Guideline Type		Prior Authorization – IL and MN Plans		
Product Name	Gen	eric Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADAL MG/0	IMUMAB PEN-INJECTOR KIT 40 .8ML	6627001500F420	Brand
HUMIRA PEN ADAL MG/0		IMUMAB PEN-INJECTOR KIT 40 .8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER ADAL MG/0		IMUMAB PEN-INJECTOR KIT 40 .8ML	6627001500F420	Brand

HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand

HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g. nails, scalp, genitals, or intertriginous areas

AND

3.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g., topical corticosteroids, calcipotriene, retinoids)

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.11 (3 every 28 days) for
	30 days Second PA: Approve at GPI 8 with Ignore Drug Status of I for 12 mont hs

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis Plaque Psoriasis		
Approval Length 12/31/2039*		
Guideline Type Prior Authorization – All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand

HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40	00070045005400	Brand
I IOWINA FEN	MG/0.4ML	6627001500F430	
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand

HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g. nails, scalp, genitals, or intertriginous areas

AND

3.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g., topical corticosteroids, calcipotriene, retinoids)

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve with a MDD of 0.11 (3 every 28 days) for 30 days Second PA: Approve at GPI 8 with Ignore Drug Status of I to 12/31/20
	Second PA: Approve at GPI 8 with Ignore Drug Status of I to 12/31/20

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis Hidradenitis Suppurativa (HS)		
Approval Length 12 Month(s)*		
Guideline Type Prior Authorization – IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand

HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40	00070045005400	Brand
I IOWINA FEN	MG/0.4ML	6627001500F430	
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand

HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of moderate to severe and/or refractory hidradenitis suppurativa (HS) (Hurley II; Hurley III stage)

AND

2 - Prescribed by or in consultation with a dermatologist

- 3 One of the following:
- **3.1** Lesions are present despite treatment with topical antibiotics, systemic antibiotics, intralesional glucocorticoids, and/or surgical debridement

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve with a MDD of 0.22 (6 every 28 days) for 30 days Second PA: Approve at GPI 8 with Ignore Drug Status of I for 12 mont hs

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis Hidradenitis Suppurativa (HS)		
Approval Length 12/31/2039*		
Guideline Type Prior Authorization – All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand

HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
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1 - Diagnosis of moderate to severe and/or refractory hidradenitis suppurativa (HS) (Hurley II; Hurley III stage)

AND

2 - Prescribed by or in consultation with a dermatologist

- 3 One of the following:
- **3.1** Lesions are present despite treatment with topical antibiotics, systemic antibiotics, intralesional glucocorticoids, and/or surgical debridement

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve with a MDD of 0.22 (6 every 28 days) for 30 days Second PA: Approve at GPI 8 with Ignore Drug Status of I through 12/31/2039.

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis	Psoriatic Arthritis (PsA)	
Approval Length	12 Month(s)*	
Guideline Type	Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand

LILIMIDA	ADALIMIMAD PREFILLED OVERVOE VIT CO		D
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand

PSORIASIS STARTER PACK			
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

- **3** One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

Notes **Place authorization at a GPI 8 with an Ignore Drug Status of I
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Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis	Psoriatic Arthritis (PsA)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand

ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthésitis

OR

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis	Polyarticular Juvenile Idiopathic Arthritis (PJIA), Moderate to Severely Active Rheumatoid Arthritis (RA)	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand

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HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand

HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

- 1 One of the following:
 - Moderate to severely active rheumatoid arthritis (RA)
 - Polyarticular juvenile idiopathic arthritis (PJIA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

OR

Notes	*Place authorization at a GPI 8 with an Ignore Drug Status of I. **Abso
	lute contraindications to methotrexate are pregnancy, nursing, alcoholi
	sm, alcoholic liver disease or other chronic liver disease, immunodefic

iency syndromes, bone marrow hyperplasia, leukopenia, thrombocyto
penia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Humira	, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz
Diagnosis	Polyarticular Juvenile Idiopathic Arthritis (PJIA), Moderate to Severely Active Rheumatoid Arthritis (RA)
Approval Length	12/31/2039
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand

HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand

- 1 Diagnosis of one of the following:
 - Moderate to severely active rheumatoid arthritis (RA)
 - Polyarticular juvenile idiopathic arthritis (PJIA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

OR

		*Place authorization at a GPI 8 with an Ignore Drug Status of I. **Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immunodefic iency syndromes, bone marrow hyperplasia, leukopenia, thrombocyto penia or significant anemia, or hypersensitivity to methotrexate.
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Product Name: Humira	, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz
Diagnosis	Ankylosing Spondylitis (AS)
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand

HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand

1 - Diagnosis of ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

3.2 Continuation of previous therapy with Humira or an Adalimumab biosimilar, confirmed by paid claims or medical records (e.g. chart notes)

Notes	*Diago authorization at a CDI 9 with an Ignore Drug Status of I
Notes	*Place authorization at a GPI 8 with an Ignore Drug Status of I

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz

Diagnosis	Ankylosing Spondylitis (AS)
Approval Length	12/31/2039
Guideline Type	Prior Authorization – All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand

HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

Notes *Place authorization at a GPI 8 with an Ignore Drug Status of	of I
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Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis Non-infectious Uveitis		
Approval Length 12 Month(s)*		
Guideline Type Prior Authorization – IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand

HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40	6627001500F430	Brand
	MG/0.4ML	3027 00 1300F430	
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand

HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of non-infectious uveitis

AND

2 - Prescribed by or in consultation with a rheumatologist and verified by an ophthalmologist or other eye specialist

- 3 One of the following:
- 3.1 Condition classified as intermediate, posterior or panuveitis

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve with a MDD of 0.11 (3 every 28 days) for 30 days
	Second PA: Approve at GPI 8 with Ignore Drug Status of I for 12 mont
	hs

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis Non-infectious Uveitis		
Approval Length	12/31/2039*	
Guideline Type Prior Authorization – All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand

HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand

1 - Diagnosis of non-infectious uveitis

AND

2 - Prescribed by or in consultation with a rheumatologist and verified by an ophthalmologist or other eye specialist

- 3 One of the following:
- 3.1 Condition classified as intermediate, posterior or panuveitis

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same st		
	date:		
	First PA: Approve with a MDD of 0.11 (3 every 28 days) for 30 days		
	Second PA: Approve at GPI 8 with Ignore Drug Status of I through 12/		
	31/2039		

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)	
Approval Length	12 Month(s)*	
Guideline Type	Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand

HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand

PSORIASIS STARTER PACK			
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of moderate to severely active Crohn's disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** Member is considered high-risk based on at least one of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease

• Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)

OR

3.2 Both of the following:

3.2.1 Member is considered low-risk

AND

3.2.2 One of the following:

- Trial and failure, contraindication, or intolerance to one conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with one conventional therapy
- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

OR

3.3 Continuation of previous therapy with Humira or an Adalimumab biosimilar, confirmed by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve with a MDD of 0.22 (6 every 28 days) for 30 days
	Second PA: Approve at GPI 8 with Ignore Drug Status of I for 12 mont
	hs

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis Moderate to Severely Active Crohn's Disease (CD)		
Approval Length 12/31/2039*		
Guideline Type Prior Authorization – All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand

HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40	00070045005400	Brand
TIOWIINA FEN	MG/0.8ML	6627001500F420	DIAIIU
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand

ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of moderate to severely active Crohn's disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- **3** One of the following:
- **3.1** Member is considered high-risk based on at least one of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease
 - Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)

OR

- **3.2** Both of the following:
- 3.2.1 Member is considered low-risk

AND

- **3.2.2** One of the following:
 - Trial and failure, contraindication, or intolerance to one conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
 - Inadequate disease control or inability to achieve remission after an adequate trial of 3
 months with one conventional therapy
 - Demonstrated steroid dependence
 - Conventional therapy clinically inappropriate based on location of disease

OR

3.3 Continuation of previous therapy with Humira or an Adalimumab biosimilar, confirmed by paid claims or medical records (e.g. chart notes)

*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve with a MDD of 0.22 (6 every 28 days) for 30 days
Second PA: Approve at GPI 8 with Ignore Drug Status of I through 12/31/2039

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz		
Diagnosis Moderate to Severely Active Ulcerative Colitis (UC)		
Approval Length 12 Month(s)*		
Guideline Type Prior Authorization – IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand

HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand

HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Member is considered high-risk based on at least one of the following characteristics:
 - Extensive colitis
 - Deep Ulcers
 - Age less than 40 years
 - High CRP and ESR
 - Steroid-requiring disease
 - History of hospitalization
 - C. difficile infection
 - CMV Infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

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3.2 Continuation of previous therapy with Humira or an Adalimumab biosimilar, confirmed by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve with a MDD of 0.22 (6 every 28 days) for 30 days
	Second PA: Approve at GPI 8 with Ignore Drug Status of I for 12 mont
	hs

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz	
Diagnosis	Moderate to Severely Active Ulcerative Colitis (UC)
Approval Length	12/31/2039*
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand

HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand

ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand
	SYRINGE 20 MG/0.2ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML ADALIMUMAB-ADAZ SOLN PREFILLED SYR	SYRINGE 20 MG/0.2ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML ADALIMUMAB-ADAZ SOLN PREFILLED SYR 6627001504E510 6627001504E510 6627001504E510

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Member is considered high-risk based on at least one of the following characteristics:
 - Extensive colitis
 - Deep Ulcers
 - Age less than 40 years
 - High CRP and ESR
 - Steroid-requiring disease
 - History of hospitalization
 - C. difficile infection
 - CMV Infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

OR

3.2 Continuation of previous therapy with Humira or an Adalimumab biosimilar, confirmed by paid claims or medical records (e.g. chart notes)

*For new starts to therapy: Enter 2 PAs as follows with the same start
date:
First PA: Approve with a MDD of 0.22 (6 every 28 days) for 30 days
Second PA: Approve at GPI 8 with Ignore Drug Status of I through 12/
31/2039

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis, JIA, PSA, AS, plaque psoriasis
Approval Length	12/31/2039
Guideline Type	Quantity Exception – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand

HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand

1 - Failure of an adherent 3-month trial of standard maintenance dosing with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)

OR

- **2** There is a confirmed history of an approved quantity limit exception (via prior authorization or historical authorization on file) for any of the following:
 - Humira
 - Adalimumab biosimilar

OR

3 - Continuation of previous therapy with Humira or an Adalimumab biosimilar with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes)

Notes	*Place authorization at a GPI 8 with an Ignore Drug Status of I
1	1

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis, JIA, PSA, AS, plaque psoriasis
Approval Length	12 month(s)
Guideline Type	Quantity Exception – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand

HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand

HYRIMOZ ADALIMUMA 40 MG/0.8MI	AB-ADAZ SOLN PREFILLED SYRINGE	6627001504E520	Brand
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1 - Failure of an adherent 3-month trial of standard maintenance dosing with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)

OR

- **2** There is a confirmed history of an approved quantity limit exception (via prior authorization or historical authorization on file) for any of the following:
 - Humira
 - Adalimumab biosimilar

OR

3 - Continuation of previous therapy with Humira or an Adalimumab biosimilar with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes)

Notes	*Place authorization at a GPI 8 with an Ignore Drug Status of I
140103	Thate authorization at a Of 16 with an ignore brug status of 1

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz			
Diagnosis Crohn's disease, ulcerative colitis			
Approval Length 12 month(s)			
Guideline Type Quantity Exception – IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand

HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
HUMIRA PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F880	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand

ULCERATIVE COLITIS STARTER PACK			
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

- 1 One of the following:
- **1.1** Both of the following:
- **1.1.1** Failure of a two-month trial of monthly therapy after completion of induction dosing regimen

AND

1.1.2 Based on subtherapeutic drug concentrations and absence (or low levels) of drug antibodies

OR

- **1.2** There is a confirmed history of an approved quantity limit exception (via prior authorization or historical authorization on file) for any of the following:
 - Humira

• Adalimumab biosimilar

OR

2 - Continuation of previous therapy with Humira or an Adalimumab biosimilar with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes)

Product Name: Humira, Hadlima, Adalimumab-fkjp, Hyrimoz, Adalimumab-adaz			
Diagnosis Crohn's disease, ulcerative colitis			
Approval Length 12 month(s)			
Guideline Type Quantity Exception – All plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.8ML	6627001500F420	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 40 MG/0.4ML	6627001500F430	Brand
HUMIRA PEN- PEDIATRIC UC STARTER PACK	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- CD/UC/HS STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML	6627001500F440	Brand
HUMIRA PEN- PS/UV STARTER	ADALIMUMAB PEN-INJECTOR KIT 80 MG/0.8ML & 40 MG/0.4ML	6627001500F450	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 10 MG/0.1ML	6627001500F804	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 20 MG/0.2ML	6627001500F809	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001500F820	Brand
HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 40 MG/0.4ML	6627001500F830	Brand

HUMIRA PEDIATRIC CROHNS DISEASE	ADALIMUMAB PREFILLED SYRINGE KIT 80 MG/0.8ML	6627001500F840	Brand
STARTER PACK HUMIRA	ADALIMUMAB PREFILLED SYRINGE KIT 80	6627001500F880	Brand
PEDIATRIC CROHNS DISEASE STARTER PACK	MG/0.8ML & 40 MG/0.4ML		
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001520D510	Brand
HADLIMA PUSHTOUCH	ADALIMUMAB-BWWD SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001520D520	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001520E510	Brand
HADLIMA	ADALIMUMAB-BWWD SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001520E520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP AUTO-INJECTOR KIT 40 MG/0.8ML	6627001535F520	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 20 MG/0.4ML	6627001535F810	Brand
ADALIMUMAB- FKJP	ADALIMUMAB-FKJP PREFILLED SYRINGE KIT 40 MG/0.8ML	6627001535F820	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
ADALIMUMAB- ADAZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.4ML	6627001504D515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 40 MG/0.8ML	6627001504D520	Brand
HYRIMOZ CROHN'S DISEASE AND ULCERATIVE COLITIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML	6627001504D540	Brand
HYRIMOZ PLAQUE PSORIASIS STARTER PACK	ADALIMUMAB-ADAZ SOLN AUTO-INJECTOR 80 MG/0.8ML & 40 MG/0.4ML	6627001504D560	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 10 MG/0.1ML	6627001504E508	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 20 MG/0.2ML	6627001504E513	Brand

HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.4ML	6627001504E515	Brand
HYRIMOZ	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 40 MG/0.8ML	6627001504E520	Brand
HYRIMOZ PEDIATRIC CROHNS DISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYRINGE 80 MG/0.8ML	6627001504E540	Brand
HYRIMOZ PEDIATRIC CROHN'SDISEASE STARTER PACK	ADALIMUMAB-ADAZ SOLN PREFILLED SYR 80 MG/0.8ML & 40 MG/0.4ML	6627001504E560	Brand

- 1 One of the following:
- **1.1** Both of the following:
- **1.1.1** Failure of a two-month trial of monthly therapy after completion of induction dosing regimen

AND

1.1.2 Based on subtherapeutic drug concentrations and absence (or low levels) of drug antibodies

OR

- **1.2** There is a confirmed history of an approved quantity limit exception (via prior authorization or historical authorization on file) for any of the following:
 - Humira
 - Adalimumab biosimilar

OR

2 - Continuation of previous therapy with Humira or an Adalimumab biosimilar with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes)

Notes	*Place authorization at a GPI 8 with an Ignore Drug Status of I
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2. Background

Benefit/Coverage/Program Information

Quantity Limits

Drug Name	Drug Status	Quantity Limits (maintenance/28 days) based on indication	Approval Limits
Adalimumab-bwwd	Preferred	#2	None*
(Hadlima)	Restricted	#4 for HS indication	
Adalimumab-fkjp	Preferred	#2	None*
(unbranded)	Restricted	#4 for HS indication	
Adalimumab-adaz (Hyrimoz)	Preferred	#2	None*
	Restricted	#4 for HS indication	
Adalimumab-adaz-	Preferred	#2	None*
(unbranded Hyrimoz)	Restricted	#4 for HS indication	

^{*}Initial and renewal approvals limited to 12 months for IL and MN plans

3. Revision History

Date	Notes
4/15/2024	Update Program

Adlarity (donepezil)		
The State Annual and the State Annual		

Prior Authorization Guideline

Guideline ID	GL-129155
Guideline Name	Adlarity (donepezil)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Adlarity	
Approval Length	12/31/2039
Guideline Type	Prior Authorization- All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ADLARITY	DONEPEZIL HYDROCHLORIDE TD PATCH WEEKLY 5 MG/DAY	62051025108820	Brand
ADLARITY	DONEPEZIL HYDROCHLORIDE TD PATCH WEEKLY 10 MG/DAY	62051025108830	Brand

Approval Criteria

1 - Diagnosis of dementia associated with Alzheimer's disease

AND

2 - Trial and failure, intolerance, or contraindication to two oral acetylcholinesterase inhibitors (eg donepezil, galantamine), one of which must be donepezil

AND

3 - Trial and failure, intolerance, or contraindication to an acetylcholinesterase inhibitor patch (eg rivastigmine patch)

Product Name: Adlarity	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization-IL and MN plans

Product Name	Generic Name	GPI	Brand/Generic
ADLARITY	DONEPEZIL HYDROCHLORIDE TD PATCH WEEKLY 5 MG/DAY	62051025108820	Brand
ADLARITY	DONEPEZIL HYDROCHLORIDE TD PATCH WEEKLY 10 MG/DAY	62051025108830	Brand

Approval Criteria

1 - Diagnosis of dementia associated with Alzheimer's disease

AND

2 - Trial and failure, intolerance, or contraindication to two oral acetylcholinesterase inhibitors (eg donepezil, galantamine), one of which must be donepezil

AND

3 - Trial and failure, intolerance, or contraindication to an acetylcholinesterase inhibitor patch (eg rivastigmine patch)

Product Name: Adlarity	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization-IL and MN plans

Product Name	Generic Name	GPI	Brand/Generic
ADLARITY	DONEPEZIL HYDROCHLORIDE TD PATCH WEEKLY 5 MG/DAY	62051025108820	Brand
ADLARITY	DONEPEZIL HYDROCHLORIDE TD PATCH WEEKLY 10 MG/DAY	62051025108830	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

2. Revision History

Date	Notes
9/20/2023	New Program

Afrezza (Insulin Regular, Human)				
(2) The beauting word halping for the text and make a count or state that the present according to the state of the state				

Prior Authorization Guideline

Guideline ID	GL-129628
Guideline Name	Afrezza (Insulin Regular, Human)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Afrezza		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 4 UNIT/CARTRIDGE	27104010002940	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 8 UNIT/CARTRIDGE	27104010002950	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 12 UNIT/CARTRIDGE	27104010002955	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHAL POWD 90 X 4 UNIT & 90 X 8 UNIT	27104010002978	Brand

AFREZZA	INSULIN REGULAR (HUMAN) INH POWD 90 X 8 UNIT & 90 X 12 UNIT	27104010002988	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INH POWD 60X4 & 60X8 & 60X12 UT/CART	27104010002990	Brand

1 - Diagnosis of diabetes mellitus

AND

2 - Prescription is initiated by, or in consultation with, an Endocrinologist

AND

3 - Documented disability that does not physically allow the administration of insulin from conventional vials or pens

AND

4 - Does not have a documented chronic lung disease (asthma, COPD, etc.)

AND

5 - Is a nonsmoker

Product Name: Afrezza					
Approval Length		12 month(s)			
Therapy Stage		Reauthorization			
Guideline Type		Prior Authorization - IL and MN Plans			
Product Name	Generic Name		GPI	Brand/Generic	
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 4 UNIT/CARTRIDGE		27104010002940	Brand	

AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 8 UNIT/CARTRIDGE	27104010002950	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 12 UNIT/CARTRIDGE	27104010002955	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHAL POWD 90 X 4 UNIT & 90 X 8 UNIT	27104010002978	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INH POWD 90 X 8 UNIT & 90 X 12 UNIT	27104010002988	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INH POWD 60X4 & 60X8 & 60X12 UT/CART	27104010002990	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Afrezza		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 4 UNIT/CARTRIDGE	27104010002940	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 8 UNIT/CARTRIDGE	27104010002950	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHALATION POWDER 12 UNIT/CARTRIDGE	27104010002955	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INHAL POWD 90 X 4 UNIT & 90 X 8 UNIT	27104010002978	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INH POWD 90 X 8 UNIT & 90 X 12 UNIT	27104010002988	Brand
AFREZZA	INSULIN REGULAR (HUMAN) INH POWD 60X4 & 60X8 & 60X12 UT/CART	27104010002990	Brand

Approval Criteria

1 - Diagnosis of diabetes mellitus



2 - Prescription is initiated by, or in consultation with, an Endocrinologist

AND

3 - Documented disability that does not physically allow the administration of insulin from conventional vials or pens

AND

4 - Does not have a documented chronic lung disease (asthma, COPD, etc.)

AND

5 - Is a nonsmoker

2. Revision History

Date	Notes
10/6/2023	New Program

Alosetron				
(g) "Perhandrang usura terdagian. Terbang karatan sasan, ususul, o salam indi baba bid patan bid parata bid pa				

Prior Authorization Guideline

Guideline ID	GL-144840
Guideline Name	Alosetron
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	1/20/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Generic Alosetron				
Approval Length		12 month(s)		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Gene	eric Name	GPI	Brand/Generic
ALOSETRON HYDROCHLORIDE	ALOS	ETRON HCL TAB 0.5 MG (BASE EQUIV)	52554015100310	Generic
ALOSETRON HYDROCHLORIDE	ALOS	ETRON HCL TAB 1 MG (BASE EQUIV)	52554015100320	Generic
Approval Criteria				

- 1 One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of diarrhea- predominant irritable bowel syndrome (IBS)

AND

1.1.2 Member is female

AND

1.1.3 Trial and failure of one-month trial of conventional therapy (such as loperamide or diphenoxylate/atropine)

OR

1.2 Continuation of prior therapy with alosetron, verified by paid claims, medical records (e.g. chart notes), or provider attestation.

Product Name: Generic Alosetron	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
ALOSETRON HYDROCHLORIDE	ALOSETRON HCL TAB 0.5 MG (BASE EQUIV)	52554015100310	Generic
ALOSETRON HYDROCHLORIDE	ALOSETRON HCL TAB 1 MG (BASE EQUIV)	52554015100320	Generic

Approval Criteria

- 1 One of the following:
 - **1.1** All of the following:
 - **1.1.1** Diagnosis of diarrhea- predominant irritable bowel syndrome (IBS)

AND

1.1.2 Member is female

AND

1.1.3 Trial and failure of one-month trial of conventional therapy (such as loperamide or diphenoxylate/atropine)

OR

1.2 Member is new to the plan (within the past 90 days) and submission of medical notes (e.g., chart notes) from the past 12 months that the person is continuing therapy with the requested medication

2. Revision History

Date	Notes
3/25/2024	New program

Ampyra (Dalfampridine)			
The based region community of the first term to contact a state and factor to provide an executive and the contract and the c			

Prior Authorization Guideline

Guideline ID	GL-129138
Guideline Name	Ampyra (Dalfampridine)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Dalfampridine	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
DALFAMPRIDINE ER	DALFAMPRIDINE TAB ER 12HR 10 MG	62406030007420	Generic

Approval Criteria

1 - Diagnosis of multiple sclerosis

AND

2 - Person is ambulatory with or without assistance

AND

3 - Baseline assessment (ex: timed 25-foot walk) or supporting documentation indicating difficulty ambulating (ex: gait contributing to falls, etc.)

Product Name: Generic Dalfampridine	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
DALFAMPRIDINE ER	DALFAMPRIDINE TAB ER 12HR 10 MG	62406030007420	Generic

Approval Criteria

1 - Prescriber provides clinical documentation from the previous 12 months that the person has a diagnosis of multiple sclerosis and remains ambulatory (with or without assistance).

2. Revision History

Date	Notes
9/20/2023	New Program

Antifibrotic Agents
The State of the S

Prior Authorization Guideline

Guideline ID	GL-129091
Guideline Name	Antifibrotic Agents
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic pirfenidone		
12/31/2039		
Prior Authorization - All plans except IL and MN Plans		
Seneric		

1 - Diagnosis of idiopathic pulmonary fibrosis	(IPF) confirmed by high-resolution computed
tomography	

2 - Member is 18 years of age or older

AND

3 - Prescribed by or in consultation with a pulmonologist

Product Name: Generic pirfenidone	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
PIRFENIDONE	PIRFENIDONE CAP 267 MG	45550060000120	Generic
PIRFENIDONE	PIRFENIDONE TAB 267 MG	45550060000325	Generic
PIRFENIDONE	PIRFENIDONE TAB 801 MG	45550060000345	Generic

Approval Criteria

1 - Diagnosis of idiopathic pulmonary fibrosis (IPF) confirmed by high-resolution computed tomography

AND

2 - Member is 18 years of age or older

AND

3 - Prescribed by or in consultation with a pulmonologist

Product Name: Ofev	
Approval Length	12/31/2039
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
OFEV	NINTEDANIB ESYLATE CAP 100 MG (BASE EQUIVALENT)	45554050200120	Brand
OFEV	NINTEDANIB ESYLATE CAP 150 MG (BASE EQUIVALENT)	45554050200130	Brand

Approval Criteria

- **1** ONE of the following:
- **1.1** Diagnosis of idiopathic pulmonary fibrosis (IPF) confirmed by high-resolution computed tomography

OR

1.2 Chronic Fibrosing Interstitial Lung Diseases with a Progressive Phenotype

OR

- **1.3** Both of the following:
 - Diagnosis of systemic sclerosis associated lung disease (SSc-ILD)
 - Trial and failure, contraindication or intolerance to cyclophosphamide

AND

2 - Member is 18 years of age or older

3 - Prescribed by or in consultation with a pulmonologist

Product Name: Ofev	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
OFEV	NINTEDANIB ESYLATE CAP 100 MG (BASE EQUIVALENT)	45554050200120	Brand
OFEV	NINTEDANIB ESYLATE CAP 150 MG (BASE EQUIVALENT)	45554050200130	Brand

Approval Criteria

- **1** ONE of the following:
- **1.1** Diagnosis of idiopathic pulmonary fibrosis (IPF) confirmed by high-resolution computed tomography

OR

1.2 Chronic Fibrosing Interstitial Lung Diseases with a Progressive Phenotype

OR

- **1.3** Both of the following:
 - Diagnosis of systemic sclerosis associated lung disease (SSc-ILD)
 - Trial and failure, contraindication or intolerance to cyclophosphamide

AND

2 - Member is 18 years of age or older

AND

3 - Prescribed by or in consultation with a pulmonologist

Product Name: Generic pirfenidone, Ofev	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
PIRFENIDONE	PIRFENIDONE CAP 267 MG	45550060000120	Generic
PIRFENIDONE	PIRFENIDONE TAB 267 MG	45550060000325	Generic
PIRFENIDONE	PIRFENIDONE TAB 801 MG	45550060000345	Generic
OFEV	NINTEDANIB ESYLATE CAP 100 MG (BASE EQUIVALENT)	45554050200120	Brand
OFEV	NINTEDANIB ESYLATE CAP 150 MG (BASE EQUIVALENT)	45554050200130	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
9/19/2023	2024 New Implementation

Arikayce (amikacin inhaled)		
So the followage was to deplay to Tree the state of teach of and any first this passes the country for the state of the st		

Guideline ID	GL-128153
Guideline Name	Arikayce (amikacin inhaled)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Arikayce*	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
ARIKAYCE	AMIKACIN SULFATE LIPOSOME INHAL SUSP 590 MG/8.4ML (BASE EQ)	07000010121830	Brand

Approval Criteria

1 - Diagnosis of Mycobacterium avium complex (MAC) lung disease

2 - Prescribed by, or in consultation with, an Infectious Disease expert

AND

3 - Submission of medical records (e.g., chart notes) of positive sputum cultures despite at least 6 months of multidrug background therapy

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
	new to plan, reautionzation official applies

Product Name: Arikayce*	
Approval Length	6 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - All Plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
ARIKAYCE	AMIKACIN SULFATE LIPOSOME INHAL SUSP 590 MG/8.4ML (BASE EQ)	07000010121830	Brand

Approval Criteria

1 - Diagnosis of Mycobacterium avium complex (MAC) lung disease

AND

2 - Prescribed by, or in consultation with, an Infectious Disease expert

AND

	cal records (e.g., chart notes) of positive sputum cultures despite at drug background therapy
Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

Product Name: Arikayce*	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
ARIKAYCE	AMIKACIN SULFATE LIPOSOME INHAL SUSP 590 MG/8.4ML (BASE EQ)	07000010121830	Brand

1 - Diagnosis of Mycobacterium avium complex (MAC) lung disease

AND

2 - Prescribed by, or in consultation with, an Infectious Disease expert

AND

3 - Person achieves and/or maintains negative sputum culture status by 6 months

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers
	will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

Date	Notes
11/3/2023	New Program

Atacand (candesartan)
The bit sell integression to indeplay on. The fire ray have been record, marked, or defect leady that for the politics disc convenils and invadion.

Guideline ID	GL-144534	
Guideline Name	Atacand (candesartan)	
Formulary	Quartz	

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Generic Candesartan, Candesartan/hctz				
Approval Length 12/31/203		9		
Guideline Type	Prior Auth	orization - All plans excep	t IL and MN	
Product Name		Generic Name	GPI	Brand/Generic
CANDESARTAN CILEXETIL		CANDESARTAN CILEXETIL TAB 4 MG	36150020100310	Generic
CANDESARTAN CILEXETIL		CANDESARTAN CILEXETIL TAB 8 MG	36150020100320	Generic
CANDESARTAN CILEXETIL		CANDESARTAN CILEXETIL TAB 16 MG	36150020100330	Generic
CANDESARTAN CILEXETIL		CANDESARTAN CILEXETIL TAB 32 MG	36150020100340	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE		CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 16-12.5 MG	36994002200320	Generic

CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 32-12.5 MG	36994002200340	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 32-25 MG	36994002200350	Generic

1 - Diagnosis of heart failure

Product Name: Generic Candesartan, Candesartan/hctz	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 4 MG	36150020100310	Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 8 MG	36150020100320	Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 16 MG	36150020100330	Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 32 MG	36150020100340	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 16-12.5 MG	36994002200320	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 32-12.5 MG	36994002200340	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 32-25 MG	36994002200350	Generic

Approval Criteria

1 - Diagnosis of heart failure

Product Name: Generic Candesartan, Candesartan/hctz

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 4 MG	36150020100310	Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 8 MG	36150020100320	Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 16 MG	36150020100330	Generic
CANDESARTAN CILEXETIL	CANDESARTAN CILEXETIL TAB 32 MG	36150020100340	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 16-12.5 MG	36994002200320	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 32-12.5 MG	36994002200340	Generic
CANDESARTAN CILEXETIL/HYDROCHLOROTHIAZIDE	CANDESARTAN CILEXETIL- HYDROCHLOROTHIAZIDE TAB 32-25 MG	36994002200350	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Date	Notes
3/18/2024	Updated product name

Auryxia (Ferric Citrate)		
The billioning search shipping to Trult has been been round, created, a dead ship but his to constitut and has been been been been been been been bee		

Guideline ID	GL-129081	
Guideline Name	Auryxia (Ferric Citrate)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Auryxia				
Approval Length		12/31/2039		
Guideline Type		Prior Authorization - All Plans Except IL and MN Plans		
Product Generic Name Name		GPI	Brand/Generic	
AURYXIA	FERRIC CITRATE TAB 1 GM (210 MG FERRIC IRON)		52800030100320	Brand

Approval Criteria

- 1 One of the following:
 - **1.1** All of the following:
 - 1.1.1 Diagnosis of chronic kidney disease (CKD)

AND
1.1.2 Member has hyperphosphatemia requiring dialysis
AND
1.1.3 Trial and failure or intolerance (e.g. side effects) from Both of the following:
Sevelamer product (i.e., Renagel, Renvela)Fosrenol (lanthanum)
OR
1.2 All of the following:
1.2.1 Diagnosis of iron deficiency anemia
AND
1.2.2 Member has chronic kidney disease (CKD)
AND
1.2.3 Memebr is not on dialysis
AND
1.2.4 Trial and failure, or intolerance (e.g. side effects) to at least 2 forms of oral iron products (i.e., ferrous sulfate, polysaccharide complex, ferrous fumarate, ferrous gluconate)

Product Name: Auryxia		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	

Guideline Type		Prior Authorization - IL and MN Plans*			
Product Name			GPI	Brand/Generic	
AURYXIA	FERRIC CITRATE TAB 1 GM (210 MG FERRIC IRON)		52800030100320	Brand	
Approval Criteria					
1 - One of the following:					

1.1 All of the following:

1.1.1 Diagnosis of chronic kidney disease (CKD)

1.2.2 Member has chronic kidney disease (CKD)

1.1.2 Member has hyperphosphatemia requiring dialysis
AND
1.1.3 Trial and failure or intolerance (e.g. side effects) from Both of the following:

Sevelamer product (i.e., Renagel, Renvela)
Fosrenol (lanthanum)

OR
1.2 All of the following:
1.2.1 Diagnosis of iron deficiency anemia

AND

AND

AND

1.2.3 Member is not on dialysis

AND

1.2.4 Trial and failure, or intolerance (e.g. side effects) to at least 2 forms of oral iron products (i.e., ferrous sulfate, polysaccharide complex, ferrous fumarate, ferrous gluconate)

Notes	*Members new to the plan (as evidenced by coverage effective date o
	f less than or equal to 90 days) must meet the initial criteria coverage

Product Name: Auryxia		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans*	

Product Name	Generic Name	GPI	Brand/Generic
AURYXIA	FERRIC CITRATE TAB 1 GM (210 MG FERRIC IRON)	52800030100320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Notes	*Members new to the plan (as evidenced by coverage effective date o
	f less than or equal to 90 days) must meet the initial criteria coverage

Date	Notes
8/23/2023	2024 New Implementation

Au	stedo	(deute	etrabe	nazine	9)	
Deckstring	uuroot ke diqiqqut. Tre lle ray haar baar	menod, unamed, or dilded. Verily that he led yo	ich in der somerifik auf koalen.			

Guideline ID	GL-129069
Guideline Name	Austedo (deutetrabenazine)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Na	me: Austed	o, Austedo XR		
Approval L	ength	12/31/2039		
Guideline 7	уре	Prior Authorization - All Plans Excep	ot IL and MN Plans	5
Product Name	Generic Na	me	GPI	Brand/Generic
AUSTEDO PATIENT TITRATION KIT	DEUTETRAB 9 MG & 12 M	ENAZINE TAB TITRATION PACK 6 MG & G	6238003000B720	Brand
AUSTEDO XR PATIENT TITRATION KIT	DEUTETRAB MG & 12 MG	ENAZINE TAB ER TITRATION PACK 6 & 24 MG	6238003000C120	Brand
AUSTEDO	DEUTETRAB	ENAZINE TAB 6 MG	62380030000310	Brand

AUSTEDO	DEUTETRABENAZINE TAB 9 MG	62380030000320	Brand
AUSTEDO	DEUTETRABENAZINE TAB 12 MG	62380030000330	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 6 MG	62380030007510	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 12 MG	62380030007520	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 24 MG	62380030007530	Brand

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		OI.	นาษ	101	lowing:
	_				

- **1.1** All of the following:
- **1.1.1** Diagnosis of chorea associated with Huntington's disease

AND

1.1.2 Prescribed by or in consultation with a neurologist or other expert in the treatment of Huntington's disease

OR

- **1.2** All of the following:
- 1.2.1 Diagnosis of tardive dyskinesia

AND

- **1.2.2** One of the following:
 - Symptoms persist despite discontinuation of the offending dopamine receptor blocking drug
 - Submission of medical records (e.g. chart notes) why discontinuation of the drug is not a treatment option for the person based on their diagnosis and previous treatment history

1.2.3 Trial and inadequate response to clonazepam (e.g., did not control symptoms, caused significant side effects)

AND

1.2.4 Trial and failure, contraindication or intolerance to an adequate trial of trihexyphenidyl with documentation of tardive dystonia

AND

1.2.5 Prescribed by or in consultation with a neurologist or other expert in the treatment of tardive dyskinesia/movement disorders

Product Name: Austed	o, Austedo XR
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
AUSTEDO PATIENT TITRATION KIT	DEUTETRABENAZINE TAB TITRATION PACK 6 MG & 9 MG & 12 MG	6238003000B720	Brand
AUSTEDO XR PATIENT TITRATION KIT	DEUTETRABENAZINE TAB ER TITRATION PACK 6 MG & 12 MG & 24 MG	6238003000C120	Brand
AUSTEDO	DEUTETRABENAZINE TAB 6 MG	62380030000310	Brand
AUSTEDO	DEUTETRABENAZINE TAB 9 MG	62380030000320	Brand
AUSTEDO	DEUTETRABENAZINE TAB 12 MG	62380030000330	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 6 MG	62380030007510	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 12 MG	62380030007520	Brand

AUSTEDO	DEUTETRABENAZINE TAB ER 24HR 24 MG	62380030007530	Brand
XR			

- **1** One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of chorea associated with Huntington's disease

AND

1.1.2 Prescribed by or in consultation with a neurologist or other expert in the treatment of Huntington's disease

OR

- **1.2** All of the following:
- **1.2.1** Diagnosis of tardive dyskinesia

AND

- **1.2.2** One of the following:
 - Symptoms persist despite discontinuation of the offending dopamine receptor blocking drug
 - Submission of medical records (e.g. chart notes) why discontinuation of the drug is not a treatment option for the person based on their diagnosis and previous treatment history

AND

1.2.3 Trial and inadequate response to clonazepam (e.g., did not control symptoms, caused significant side effects)

1.2.4 Trial and failure, contraindication or intolerance to an adequate trial of trihexyphenidyl with documentation of tardive dystonia

AND

1.2.5 Prescribed by or in consultation with a neurologist or other expert in the treatment of tardive dyskinesia/movement disorders

Notes	*Members new to the plan (as evidenced by coverage effective date o
	f less than or equal to 90 days) must meet initial criteria for coverage

Product Name: Austed	lo, Austedo XR
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
AUSTEDO PATIENT TITRATION KIT	DEUTETRABENAZINE TAB TITRATION PACK 6 MG & 9 MG & 12 MG	6238003000B720	Brand
AUSTEDO XR PATIENT TITRATION KIT	DEUTETRABENAZINE TAB ER TITRATION PACK 6 MG & 12 MG & 24 MG	6238003000C120	Brand
AUSTEDO	DEUTETRABENAZINE TAB 6 MG	62380030000310	Brand
AUSTEDO	DEUTETRABENAZINE TAB 9 MG	62380030000320	Brand
AUSTEDO	DEUTETRABENAZINE TAB 12 MG	62380030000330	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 6 MG	62380030007510	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 12 MG	62380030007520	Brand
AUSTEDO XR	DEUTETRABENAZINE TAB ER 24HR 24 MG	62380030007530	Brand

Approval Criteria	
	cal records (e.g., chart notes) documenting that within the past 12 continuing therapy with the requested drug
Notes	*Members new to the plan (as evidenced by coverage effective date o f less than or equal to 90 days) must meet initial criteria for coverage

Date	Notes
9/20/2023	2024 New Implementation

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Guideline ID	GL-128137
Guideline Name	Auvelity (dextromethorphan-bupropion)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Auvelity	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type Step Therapy - IL and MN Plans	

L	Product Name	Generic Name	GPI	Brand/Generic
	AUVELITY	DEXTROMETHORPHAN HBR-BUPROPION HCL TAB ER 45-105 MG	58999902300420	Brand

Approval Criteria

1 - Trial and failure, contraindication, or intolerance of at least two preferred antidepressants

within the Selective Serotonin Reuptake inhibitor (SSRI) or Serotonin Norepinephrine Reuptake inhibitor (SNRI) drug classes

- citalopram
- escitalopram
- sertraline
- paroxetine
- fluoxetine
- venlafaxine
- duloxetine

Product Name: Auvelity	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
AUVELITY	DEXTROMETHORPHAN HBR-BUPROPION HCL TAB ER 45-105 MG	58999902300420	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Auvelity	
Approval Length 12/31/2039	
Guideline Type	Step Therapy - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
AUVELITY	DEXTROMETHORPHAN HBR-BUPROPION HCL TAB ER 45-105 MG	58999902300420	Brand

Approval Criteria

1 - Trial and failure, contraindication, or intolerance of at least two preferred antidepressants

within the Selective Serotonin Reuptake inhibitor (SSRI) or Serotonin Norepinephrine Reuptake inhibitor (SNRI) drug classes

- citalopram
- escitalopram
- sertraline
- paroxetine
- fluoxetine
- venlafaxine
- duloxetine

Date	Notes
8/21/2023	New Program

Azelex, Finacea (Azelaic Acid)		
Sambar dangunan kadapun beda ng kasalam sarang samag a sama kenji berbata yanan ka mendikan kadalan		

Guideline ID	GL-127879
Guideline Name	Azelex, Finacea (Azelaic Acid)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Brand Finacea, Generic Azelaic Acid, Brand Azelex	
Approval Length	12 month(s)
Therapy Stage Initial Authorization	
Guideline Type Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
AZELEX	AZELAIC ACID CREAM 20%	90050005103720	Brand
FINACEA	AZELAIC ACID FOAM 15%	90060010003920	Brand
AZELAIC ACID	AZELAIC ACID GEL 15%	90060010004020	Generic

Approval Criteria

- 1 Both of the following:
- **1.1** Trial and failure of one topical tretinoin
 - tretinoin 0.01% gel
 - tretinoin 0.025% gel/cream
 - tretinoin 0.05% gel/cream
 - tretinoin 0.1% cream

1.2 Trial of topical adapalene (0.1% gel/cream, 0.3% gel)

OR

2 - For Generic Azelaic acid 15% or Brand Finacea 15% Foam: Diagnosis of rosacea

Product Name: Brand Finacea, Generic Azelaic Acid, Brand Azelex		
Approval Length	12 month(s)	
Therapy Stage	ge Reauthorization	
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
AZELEX	AZELAIC ACID CREAM 20%	90050005103720	Brand
FINACEA	AZELAIC ACID FOAM 15%	90060010003920	Brand
AZELAIC ACID	AZELAIC ACID GEL 15%	90060010004020	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Brand Finacea, Generic Azelaic Acid, Brand Azelex	
Approval Length	12/31/2039

Guideline Type		Step Therapy - All Plans except IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
AZELEX	AZELAIC ACID CREAM 20%		90050005103720	Brand
FINACEA	AZELAIC ACID FOAM 15%		90060010003920	Brand
AZELAIC ACID			90060010004020	Generic

- 1 Both of the following:
- 1.1 Trial and failure of one topical tretinoin
 - tretinoin 0.01% gel
 - tretinoin 0.025% gel/cream
 - tretinoin 0.05% gel/cream
 - tretinoin 0.1% cream

AND

1.2 Trial of topical adapalene (0.1% gel/cream, 0.3% gel)

OR

2 - For Generic Azelaic acid 15% or Brand Finacea 15% Foam: Diagnosis of rosacea

Date	Notes
8/25/2023	New Programs

Baxdela (Delafloxacin)
(g) "Perhandrag sour to dipper." Tellow, has been source, county or distinct the protect for annual dark basis.

Guideline ID	GL-136393
Guideline Name	Baxdela (Delafloxacin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Baxdel			
Approval Length See Note*			
Guideline Type Prior Authorization - All Plans except IL and MN Plans		3	
Product Generic Name GPI Brand/Gen		Brand/Generic	

Product Name	Generic Name	GPI	Brand/Generic
BAXDELA	DELAFLOXACIN MEGLUMINE TAB 450 MG (BASE EQUIV)	05000025100320	Brand
BAXDELA	DELAFLOXACIN MEGLUMINE FOR IV SOLN 300 MG (BASE EQUIV)	05000025102120	Brand

Approval Criteria

1 - One of the following:

1.1 Person has been receiving drug during hospitalization and needs to complete the course of therapy as an outpatient.

OR

- **1.2** Both of the following:
- **1.2.1** Outpatient treatment of bacterial resistant strains as ordered by, or in documented consultation with an Infectious Disease Specialist

AND

1.2.2 Report of cultures and susceptibilities documenting resistance to preferred alternatives needs to be provided for approval

Notes * Approve for duration of treatment, usually 6-14 days for	or 1 fill
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Product Name: Baxdela	
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
BAXDELA	DELAFLOXACIN MEGLUMINE TAB 450 MG (BASE EQUIV)	05000025100320	Brand
BAXDELA	DELAFLOXACIN MEGLUMINE FOR IV SOLN 300 MG (BASE EQUIV)	05000025102120	Brand

Approval Criteria

1 - Person has been receiving drug during hospitalization and needs to complete the course of therapy as an outpatient.

OR

2 - Both of the following

2.1 Outpatient treatment of bacterial resistant strains as ordered by, or in documented consultation with an Infectious Disease Specialist.

AND

2.2 Report of cultures and susceptibilities documenting resistance to preferred alternatives needs to be provided for approval

OR

3 - For Illinois Plans Only - the requested drug is being used for the long-term treatment of tick-borne disease

Date	Notes
11/17/2023	Update guideline

Belsomra (suvorexant)			
The interferoment in the last to the control of the			

Guideline ID	GL-136538
Guideline Name	Belsomra (suvorexant)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Belsomra	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
BELSOMRA	SUVOREXANT TAB 5 MG	60500070000305	Brand
BELSOMRA	SUVOREXANT TAB 10 MG	60500070000310	Brand
BELSOMRA	SUVOREXANT TAB 15 MG	60500070000315	Brand
BELSOMRA	SUVOREXANT TAB 20 MG	60500070000320	Brand

1 - Person needs the medication for sleep

AND

2 - Trial and failure, contraindication, or intolerance to two preferred alternatives (e.g., zolpidem, eszopiclone, zaleplon, trazodone, mirtazapine, quetiapine, temazepam, etc.)

Product Name: Belsomra	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
BELSOMRA	SUVOREXANT TAB 5 MG	60500070000305	Brand
BELSOMRA	SUVOREXANT TAB 10 MG	60500070000310	Brand
BELSOMRA	SUVOREXANT TAB 15 MG	60500070000315	Brand
BELSOMRA	SUVOREXANT TAB 20 MG	60500070000320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Belsomra	
Approval Length 12/31/2039	
Guideline Type	Prior Authorization-All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
BELSOMRA	SUVOREXANT TAB 5 MG	60500070000305	Brand
BELSOMRA	SUVOREXANT TAB 10 MG	60500070000310	Brand
BELSOMRA	SUVOREXANT TAB 15 MG	60500070000315	Brand
BELSOMRA	SUVOREXANT TAB 20 MG	60500070000320	Brand

1 - Person needs the medication for sleep

AND

2 - Trial and failure, contraindication, or intolerance to two preferred alternatives (e.g., zolpidem, eszopiclone, zaleplon, trazodone, mirtazapine, quetiapine, temazepam, etc.)

Date	Notes
12/8/2023	Examples included in the criteria.

Be	exarotene		
* The best of	d image current is a displayed. The file may have learn record, renamed, or disblack Verily that it is in	k points in the consectific and leaders.	

Guideline ID	GL-128907
Guideline Name	Bexarotene
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Bexarotene Gel				
Approval Length 12 month(s)				
Guideline Type Prior authorization - IL and MN Plans				
Product Generic Name		GPI	Brand/Generic	
BEXAROTENE	BEXARO1	ENE GEL 1%	90376220004020	Generic

Approval Criteria

- 1 All of the following:
 - **1.1** One of the following:

1.1.1 The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the person presents with*				
OR				
1.1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the person*				
AND				
1.2 Prescribed and monitored by an Oncologist, Hematologist, or other specialist in the treatment of malignancy				
OR				
2 - One of the following:				
2.1 Both of the following:				
2.1.1 If the request is for Minnesota Plans only, the requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the person based on one of the following:				
 United States Pharmacopeia Drug Information The American Hospital Formulary Service Drug Information One article in a major peer- reviewed medical journal recognizes the safety and efficacy of the requested drug in the person's specific condition 				
AND				
2.1.2 Prescribed and monitored by an Oncologist, Hematologist, or other specialist in the treatment of malignancy				
OR				
2.2 Both of the following:				

- **2.2.1** If the request is for IL plans only, the requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the person based on one of the following:
 - Thomson Micromedex Drugdex
 - The American Hospital Formulary Service Drug Information
 - Elsevier Gold Standard's Clinical Pharmacology
 - Two articles in a peer- reviewed medical journal from the United States or Great Britain recognize the safety and efficacy of the requested drug in the person's specific condition

2.2.2 Prescribed and monitored by an Oncologist, Hematologist, or other specialist in the treatment of malignancy

Notes	*includes any relevant genetic testing, mutations, etc.
110103	includes any relevant genetic testing, matations, etc.

Product Name: Generic Bexarotene Gel		
Approval Length 12/31/2039		
Guideline Type Prior authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
BEXAROTENE	BEXAROTENE GEL 1%	90376220004020	Generic

Approval Criteria

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the person presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the person*

	AND	
2 - Prescribed and monitored by an Oncologist, Hematologist, or other specialist in the treatment of malignancy		
Notes	*includes any relevant genetic testing, mutations, etc.	

Date	Notes
11/3/2023	New Program

Briviact (Brivaracetam)					
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Guideline ID	GL-127878
Guideline Name	Briviact (Brivaracetam)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Briviact		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
BRIVIACT	BRIVARACETAM TAB 10 MG	72600015000310	Brand
BRIVIACT	BRIVARACETAM TAB 25 MG	72600015000320	Brand
BRIVIACT	BRIVARACETAM TAB 50 MG	72600015000330	Brand
BRIVIACT	BRIVARACETAM TAB 75 MG	72600015000340	Brand
BRIVIACT	BRIVARACETAM TAB 100 MG	72600015000350	Brand
BRIVIACT	BRIVARACETAM ORAL SOLN 10 MG/ML	72600015002020	Brand

- **1** Trial and failure, contraindication, or intolerance to at least TWO preferred anticonvulsants:
 - lamotrigine
 - levetiracetam
 - carbamazepine
 - valproate
 - oxcarbazepine
 - gabapentin
 - pregabalin
 - topiramate
 - phenytoinzonisamide
 - primidone

Product Name: Briviact		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
BRIVIACT	BRIVARACETAM TAB 10 MG	72600015000310	Brand
BRIVIACT	BRIVARACETAM TAB 25 MG	72600015000320	Brand
BRIVIACT	BRIVARACETAM TAB 50 MG	72600015000330	Brand
BRIVIACT	BRIVARACETAM TAB 75 MG	72600015000340	Brand
BRIVIACT	BRIVARACETAM TAB 100 MG	72600015000350	Brand
BRIVIACT	BRIVARACETAM ORAL SOLN 10 MG/ML	72600015002020	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Briviact

Approval Length	12/31/2039
Guideline Type	Step Therapy - All Plans Except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
BRIVIACT	BRIVARACETAM TAB 10 MG	72600015000310	Brand
BRIVIACT	BRIVARACETAM TAB 25 MG	72600015000320	Brand
BRIVIACT	BRIVARACETAM TAB 50 MG	72600015000330	Brand
BRIVIACT	BRIVARACETAM TAB 75 MG	72600015000340	Brand
BRIVIACT	BRIVARACETAM TAB 100 MG	72600015000350	Brand
BRIVIACT	BRIVARACETAM ORAL SOLN 10 MG/ML	72600015002020	Brand

- **1** Trial and failure, contraindication, or intolerance to at least TWO preferred anticonvulsants:
 - lamotrigine
 - levetiracetam
 - carbamazepine
 - valproate
 - oxcarbazepine
 - gabapentin
 - pregabalin
 - topiramate
 - phenytoin
 - zonisamide
 - primidone

Date	Notes
8/25/2023	New Program

Broa	Broad Spectrum Antifungal			
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Guideline ID	GL-129108
Guideline Name	Broad Spectrum Antifungal
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Cresemba, Generic posaconazole tablet, Generic voriconazole					
Approval Length		12 month(s)			
Guideline Type		Prior Authorization			
Product Name	Gene	ric Name	GPI	Brand/Generic	
CRESEMBA		CONAZONIUM SULFATE CAP 186 MG JCONAZOLE 100 MG)	11407030100120	Brand	
CRESEMBA	_	CONAZONIUM SULF FOR IV SOL 372 MG JCONAZOLE 200MG)	11407030102130	Brand	
POSACONAZOLE	POSAG MG	CONAZOLE TAB DELAYED RELEASE 100	11407060000620	Generic	
POSACONAZOLE DR	POSACONAZOLE TAB DELAYED RELEASE 100 MG		11407060000620	Generic	
VORICONAZOLE	VORIC	ONAZOLE TAB 50 MG	11407080000320	Generic	
VORICONAZOLE	VORIC	ONAZOLE TAB 200 MG	11407080000340	Generic	

VORICONAZOLE	VORICONAZOLE FOR SUSP 40 MG/ML	11407080001920	Generic
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1 - Diagnosis of confirmed (or suspected) serious fungal infection with probable resistance to other preferred antifungals, or other antifungals have been not tolerated, or significant drugdrug interactions exist with other antifungals

OR

2 - Prescribed by or in consultation with an Infectious Disease specialist

OR

3 - For generic posaconazole tablet only, used for prophylaxis of serious fungal infections in patients with hematologic malignancies with severely compromised immunity

OR

4 - For continuation of therapy initiated as an inpatient

OR

5 - For Illinois plans only - the requested FDA approved drug is being used for the long-term treatment of tick-borne disease.

OR

6 - For Minnesota plans only - member has stage four metastatic cancer and the requested drug is being used to prevent or treat cancer-related fungal infections

Product Name: Generic posaconazole suspension, Noxafil suspension packet		
Approval Length	12 month(s)	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
POSACONAZOLE	POSACONAZOLE SUSP 40 MG/ML	11407060001820	Generic
NOXAFIL	POSACONAZOLE FOR DELAYED RELEASE SUSP PACKET 300 MG	11407060003020	Brand

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1.1 Diagnosis of confirmed (or suspected) serious fungal infection with probable resistance to other preferred antifungals, or other antifungals have been not tolerated, or significant drugdrug interactions exist with other antifungals

OR

1.2 Prescribed by or in consultation with an Infectious Disease specialist

OR

1.3 Used for prophylaxis of serious fungal infections in patients with hematologic malignancies with severely compromised immunity

OR

1.4 For continuation of therapy initiated as an inpatient

OR

1.5 For Minnesota plans only - member has stage four metastatic cancer and the requested drug is being used to prevent or treat cancer-related fungal infections

OR

1.6 For Illinois plans only - the requested FDA approved drug is being used for the long-term treatment of tick-borne disease.

2 - Member is unable to tolerate solid dosage form

Date	Notes
9/7/2023	2024 New Implementation

Bylvay (odevixibat)
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Guideline ID	GL-135532
Guideline Name	Bylvay (odevixibat)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Bylvay	
Diagnosis	Progressive Familial Intrahepatic Cholestasis (PFIC))
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
BYLVAY	ODEVIXIBAT CAP 400 MCG	52350060000120	Brand
BYLVAY	ODEVIXIBAT CAP 1200 MCG	52350060000140	Brand
BYLVAY (PELLETS)	ODEVIXIBAT PELLETS CAP SPRINKLE 200 MCG	52350060006810	Brand
BYLVAY (PELLETS)	ODEVIXIBAT PELLETS CAP SPRINKLE 600 MCG	52350060006830	Brand

Approval Criteria
1 - Diagnosis of progressive familial intrahepatic cholestasis (PFIC)
AND
2 - Disease is confirmed by one of the following:
 Genetic testing Diagnostic test (e.g., liver function test, liver ultrasound and biopsy, bile analysis)
AND
3 - Genetic testing does not indicate PFIC type 2 with ABCB11 variant encoding for nonfunctioning or absence of bile salt export pump protein (BSEP-3)
AND
4 - Member is experiencing moderate to severe cholestatic pruritus
AND
5 - Member has serum bile acid greater than 3x the upper limit of normal (ULN)
AND
6 - Member has not had a liver transplant, biliary diversion surgery within the past 6 months, or decompensated liver disease
AND
7 - Trial and failure, contraindication or intolerance to TWO of the following medications for

pruritis:

- Ursodeoxycholic acid (e.g., Ursodiol) Antihistamines (e.g., diphenhydramine, hydroxyzine)
- Rifampin
- Bile acid sequestrants (e.g., Questran, Colestid, Welchol)

- **8** Prescribed by or in consultation with one of the following:
 - hepatologist
 - gastroenterologist

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies. **For members new to plan (as evidenced by coverage effective date of less than or equal to 90 days) prescriber provides submission of me dical records (e.g., chart notes) documenting that from the previous 1 2 months, member demonstrates an improvement or stabilization in pr uritus and the member is tolerating therapy.

Product Name: Bylvay	
Diagnosis	Alagille syndrome)
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
BYLVAY	ODEVIXIBAT CAP 400 MCG	52350060000120	Brand
BYLVAY	ODEVIXIBAT CAP 1200 MCG	52350060000140	Brand
BYLVAY (PELLETS)	ODEVIXIBAT PELLETS CAP SPRINKLE 200 MCG	52350060006810	Brand
BYLVAY (PELLETS)	ODEVIXIBAT PELLETS CAP SPRINKLE 600 MCG	52350060006830	Brand

Approval Criteria

1 - Diagnosis of Alagille syndrome (ALGS)		
AND		
2 - Molecular genetic testing confirms mutations in the JAG1 or NOTCH2 gene		
AND		
3 - One of the following:		
 Total serum bile acid greater than 3x the upper limit of normal (ULN) Conjugated bilirubin greater than 1 mg/dL Fat soluble vitamin deficiency otherwise unexplainable Gamma-glutamyl transpeptidase (GGT) greater than 3x ULN 		
AND		
4 - Member is experiencing moderate to severe cholestatic pruritus		
AND		
5 - Member has not had a liver transplant or decompensated liver disease		
AND		
6 - Trial and failure, contraindication or intolerance to TWO of the following medications for pruritis:		
 Ursodeoxycholic acid (e.g., Ursodiol) Antihistamines (e.g., diphenhydramine, hydroxyzine) Rifampin 		
Bile acid sequestrants (e.g., Questran, Colestid, Welchol)		
AND		
7 - Trial and failure, contraindication or intolerance to maralixibat		

- **8** Prescribed by or in consultation with one of the following:
 - hepatologist
 - Expert in the treatment of cholestasis

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.
	**For members new to plan (as evidenced by coverage effective date of less than or equal to 90 days) prescriber provides submission of me dical records (e.g., chart notes) documenting that from the previous 1 2 months, member demonstrates an improvement or stabilization in pruritus and the member is tolerating therapy.

Product Name: Bylvay	
Diagnosis	All Indications)
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
BYLVAY	ODEVIXIBAT CAP 400 MCG	52350060000120	Brand
BYLVAY	ODEVIXIBAT CAP 1200 MCG	52350060000140	Brand
BYLVAY (PELLETS)	ODEVIXIBAT PELLETS CAP SPRINKLE 200 MCG	52350060006810	Brand
BYLVAY (PELLETS)	ODEVIXIBAT PELLETS CAP SPRINKLE 600 MCG	52350060006830	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months of therapy, member is experiencing improvement or stabilization in pruritus compared to baseline (e.g., change in member reported pruritus, change in sleeping habits due to itch)

and the member is tolerating therapy (e.g., does not have chronic diarrhea requiring ongoing intravenous fluids, bile acid reduction)		
Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies. **For members new to plan (as evidenced by coverage effective date of less than or equal to 90 days) prescriber provides submission of me dical records (e.g., chart notes) documenting that from the previous 1 2 months, member demonstrates an improvement or stabilization in pruritus and the member is tolerating therapy.	

Date	Notes
11/2/2023	2024 New Implementation

Cablivi (caplacizumab-yhdp)			
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Guideline ID	GL-128994	
Guideline Name	Cablivi (caplacizumab-yhdp)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Na	ame: Cablivi			
Approval Length		1 month (30days)		
Guideline Type		Prior Authorization - All Plans Except IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
CABLIVI CAPLACIZUMAB-YHDP FOR INJ KIT 11 MG		85151020806420	Brand	

Approval Criteria

1 - Diagnosis of severe acquired thrombotic thrombocytopenic purpura (aTTP) with at least one ADAMST13 level below 20 percent

2 - Member is 18 years of age or older

AND

- 3 Both of the following:
- **3.1** Member had been receiving plasma exchange (PEX) in combination with Cablivi (either as an inpatient or in an outpatient clinic setting)

AND

3.2 PEX has been discontinued and Cablivi therapy will continue

AND

4 - Used in combination with immunosuppressive therapy (e.g., systemic corticosteroids or rituximab)

AND

5 - Member has not had greater than 2 recurrences of aTTP while on Cablivi therapy

Product Name: Cablivi	
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CABLIVI	CAPLACIZUMAB-YHDP FOR INJ KIT 11 MG	85151020806420	Brand

Approval Criteria

1 - Diagnosis of severe acquired thrombotic thrombocytopenic purpura (aTTP) with at least one ADAMST13 level below 20 percent
AND
2 - Member is 18 years of age or older
AND
3 - Both of the following:
3.1 Member had been receiving plasma exchange (PEX) in combination with Cablivi (either as an inpatient or in an outpatient clinic setting)
AND
3.2 PEX has been discontinued and Cablivi therapy will continue
AND
4 - Used in combination with immunosuppressive therapy (e.g., systemic corticosteroids or rituximab)
AND
5 - Member has not had greater than 2 recurrences of aTTP while on Cablivi therapy
AND
6 - Cablivi (caplacizumab) will be self-administered

Date	Notes
9/8/2023	2024 New Implementation

Camzyos (mavacamten)			
(g) The interface of the large of the large content of the large of th			

Guideline ID	GL-130131
Guideline Name	Camzyos (mavacamten)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Camzyos	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
CAMZYOS	MAVACAMTEN CAP 2.5 MG	40190050000110	Brand
CAMZYOS	MAVACAMTEN CAP 5 MG	40190050000120	Brand
CAMZYOS	MAVACAMTEN CAP 10 MG	40190050000130	Brand
CAMZYOS	MAVACAMTEN CAP 15 MG	40190050000140	Brand

1 - Diagnosis of obstructive hypertrophic cardiomyopathy with New York Heart Association (NYHA) class II-III symptoms

AND

2 - Left ventricular ejection fraction (LVEF) greater than or equal to 55%

AND

3 - Member is 18 years of age or older

AND

4 - Prescribed by, or in consultation with, a Cardiologist or other expert in the treatment of hypertrophic cardiomyopathy

AND

- **5** Submission of medical records (e.g., chart notes) documenting trial and failure, contraindication, or intolerance to BOTH of the following:
 - Beta-blockers (i.e., carvedilol, labetalol, metoprolol, propranolol)
 - Calcium channel blockers (i.e., diltiazem, verapamil)

Notes	Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan regulatorization criteria applies.		
	new to plan, reauthorization criteria applies		

Product Name: Camzyos	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
CAMZYOS	MAVACAMTEN CAP 2.5 MG	40190050000110	Brand
CAMZYOS	MAVACAMTEN CAP 5 MG	40190050000120	Brand
CAMZYOS	MAVACAMTEN CAP 10 MG	40190050000130	Brand
CAMZYOS	MAVACAMTEN CAP 15 MG	40190050000140	Brand

1 - Diagnosis of obstructive hypertrophic cardiomyopathy with New York Heart Association (NYHA) class II-III symptoms

AND

2 - Member is 18 years of age or older

AND

3 - Person has been evaluated by a cardiologist, or other expert in the treatment of hypertrophic cardiomyopathy, within the previous 12 months

AND

4 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member demonstrates positive clinical response to therapy

Notes	Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu
	rer-sponsored free drug program, provider samples, and/or vouchers
	will go through initial criteria, otherwise for continuation of therapy for
	new to plan, reauthorization criteria applies

Date	Notes

10/25/2023	2024 New Implementation

Cardura XL (doxazosin ER)				
Commission and the second seco				

Guideline ID	GL-129156
Guideline Name	Cardura XL (doxazosin ER)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Cardura XL	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CARDURA XL	DOXAZOSIN MESYLATE TAB ER 24 HR 4 MG (BASE EQUIV)	56852025207520	Brand
CARDURA XL	DOXAZOSIN MESYLATE TAB ER 24 HR 8 MG (BASE EQUIV)	56852025207530	Brand

Approval Criteria

1 - The prescriber provides an evidence-based clinical rationale for why the side effects are not likely to occur with the extended-release formulation

AND

2 - Use of at least one additional preferred alpha blocker (such as terazosin or prazosin) did not control symptoms or caused side effects

AND

3 - Trial and failure, contraindication or intolerance of immediate-release doxazosin (unable to achieve symptom control due to therapy-limiting side effects)

Product Name: Cardura XL	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CARDURA XL	DOXAZOSIN MESYLATE TAB ER 24 HR 4 MG (BASE EQUIV)	56852025207520	Brand
CARDURA XL	DOXAZOSIN MESYLATE TAB ER 24 HR 8 MG (BASE EQUIV)	56852025207530	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Na	Product Name: Cardura XL			
Approval Le	ength 12/31/2039			
Guideline T	уре	Prior Authorization - All Plans excep	t IL and MN Plans	,
Product Generic Name Name		me	GPI	Brand/Generic

CARDURA XL	DOXAZOSIN MESYLATE TAB ER 24 HR 4 MG (BASE EQUIV)	56852025207520	Brand
CARDURA XL	DOXAZOSIN MESYLATE TAB ER 24 HR 8 MG (BASE EQUIV)	56852025207530	Brand

1 - The prescriber provides an evidence-based clinical rationale for why the side effects are not likely to occur with the extended-release formulation

AND

2 - Use of at least one additional preferred alpha blocker (such as terazosin or prazosin) did not control symptoms or caused side effects

AND

3 - Trial and failure, contraindication or intolerance of immediate-release doxazosin (unable to achieve symptom control due to therapy-limiting side effects)

Date	Notes
9/11/2023	New Program

Cayston (Aztreonam Inhalation Solution		

Guideline ID	GL-129106
Guideline Name	Cayston (Aztreonam Inhalation Solution)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Cayston				
Approval L	ength	12/31/2039		
Guideline Type Prior Authorization - All plans except IL and MN Plans				
Product Name	Generic Name		GPI	Brand/Generic
CAYSTON	AZTREONAM LYSINE FOR INHAL SOLN 75 MG (BASE EQUIVALENT)		16140010402120	Brand

Approval Criteria

1 - Diagnosis of cystic fibrosis

2 - Member has a history of recurrent Pseudomonas aeruginosa lung infections

AND

3 - Medication will be used for inhalation only

AND

- 4 One of the following:
 - Recurrence despite prior use of tobramycin inhalation solution
 - Submission of medical records (e.g., chart notes) documenting tobramycin resistance

Product Name: Cayston	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CAYSTON	AZTREONAM LYSINE FOR INHAL SOLN 75 MG (BASE EQUIVALENT)	16140010402120	Brand

Approval Criteria

1 - Diagnosis of cystic fibrosis

AND

2 - Member has a history of recurrent Pseudomonas aeruginosa lung infections

3 - Medication will be used for inhalation only

AND

- 4 One of the following:
 - Recurrence despite prior use of tobramycin inhalation solution
 - Submission of medical records (e.g., chart notes) documenting tobramycin resistance

Product Name: Cayston	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CAYSTON	AZTREONAM LYSINE FOR INHAL SOLN 75 MG (BASE EQUIVALENT)	16140010402120	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
7/31/2023	2024 New Implementation

Chronic Constipation Medications	
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Guideline ID	GL-144842
Guideline Name	Chronic Constipation Medications
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/18/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Linzess, Trulance, Motegrity				
Diagnosis Chronic Constipation				
Approval Le	ngth	12/31/2039		
Guideline Ty	/ре	Prior Authorization - All plans excep	t IL and MN Plans	
Product Name	Generic N	ame GPI Brand/Generic		
LINZESS	LINACLOTI	DE CAP 72 MCG	52557050000110	Brand
LINZESS	ESS LINACLOTIDE CAP 145 MCG		52557050000120	Brand
LINZESS	LINACLOTIDE CAP 290 MCG		52557050000140	Brand
TRULANCE	RULANCE PLECANATIDE TAB 3 MG 52543060000320 Brand			Brand

MOTEGRITY	PRUCALOPRIDE SUCCINATE TAB 1 MG (BASE EQUIVALENT)	52560060200320	Brand
MOTEGRITY	PRUCALOPRIDE SUCCINATE TAB 2 MG (BASE EQUIVALENT)	52560060200330	Brand

1	_	One	of	the	follo	wing:
	-	OHE	Οı	เมเต	IUIIU	willig.

- **1.1** All of the following:
- **1.1.1** Diagnosis of Chronic Constipation

AND

1.1.2 Member is 18 years of age or older

AND

1.1.3 Trial and failure, contraindication or intolerance to two first line therapies (e.g., Miralax, stimulants, fiber supplements, stool softeners)

AND

1.1.4 For Linzess Only Trial and failure, contraindication or intolerance to lubiprostone

AND

1.1.5 For Trulance and Motegrity Only - Trial and failure, contraindication or intolerance to lubiprostone and linaclotide

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

Product Name: Linzess, Trulance, Motegrity			
Diagnosis Chronic Constipation			
Approval Length 12 month(s)			
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
LINZESS	LINACLOTIDE CAP 72 MCG	52557050000110	Brand
LINZESS	LINACLOTIDE CAP 145 MCG	52557050000120	Brand
LINZESS	LINACLOTIDE CAP 290 MCG	52557050000140	Brand
TRULANCE	PLECANATIDE TAB 3 MG	52543060000320	Brand
MOTEGRITY	PRUCALOPRIDE SUCCINATE TAB 1 MG (BASE EQUIVALENT)	52560060200320	Brand
MOTEGRITY	PRUCALOPRIDE SUCCINATE TAB 2 MG (BASE EQUIVALENT)	52560060200330	Brand

- 1 One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of Chronic Constipation

AND

1.1.2 Member is 18 years of age or older

AND

1.1.3 Trial and failure, contraindication or intolerance to two first line therapies (e.g., Miralax, stimulants, fiber supplements, stool softeners)

AND

1.1.4 For Linzess Only - Trial and failure, contraindication or intolerance to lubiprostone

1.1.5 For Trulance and Motegrity Only - Trial and failure, contraindication or intolerance to lubiprostone and linaclotide

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

Product Name: Linzess, Trulance			
Diagnosis Irritable Bowel Syndrome - Constipation (IBS-C)			
Approval Length 12/31/2039			
Guideline Type Prior Authorization - All plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
LINZESS	LINACLOTIDE CAP 72 MCG	52557050000110	Brand
LINZESS	LINACLOTIDE CAP 145 MCG	52557050000120	Brand
LINZESS	LINACLOTIDE CAP 290 MCG	52557050000140	Brand
TRULANCE	PLECANATIDE TAB 3 MG	52543060000320	Brand

Approval Criteria

- 1 One of the following:
 - **1.1** All of the following:
 - 1.1.1 Diagnosis of Irritable Bowel Syndrome Constipation (IBS-C)

AND

1.1.2 Member is 18 years of age or older

1.1.3 Trial and failure, contraindication or intolerance of at least two alternative therapies (e.g., Miralax, fiber, stimulants, dicyclomine, hyoscyamine, tricyclic or SSRI antidepressants)

AND

1.1.4 For Linzess Only - Trial and failure, contraindication or intolerance to lubiprostone

AND

1.1.5 For Trulance Only - Trial and failure, contraindication or intolerance to lubiprostone and linaclotide

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

Product Name: Linzess, Trulance				
Diagnosis Irritable Bowel Syndrome - Constipation (IBS-C)				
Approval Length 12 month(s)				
Guideline Type Prior Authorization - IL and MN Plans				

Product Name	Generic Name	GPI	Brand/Generic
LINZESS	LINACLOTIDE CAP 72 MCG	52557050000110	Brand
LINZESS	LINACLOTIDE CAP 145 MCG	52557050000120	Brand
LINZESS	LINACLOTIDE CAP 290 MCG	52557050000140	Brand
TRULANCE	PLECANATIDE TAB 3 MG	52543060000320	Brand

Approval Criteria

- 1 One of the following:
- **1.1** All of the following:
- 1.1.1 Diagnosis of Irritable Bowel Syndrome Constipation (IBS-C)

1.1.2 Member is 18 years of age or older

AND

1.1.3 Trial and failure, contraindication or intolerance of at least two alternative therapies (e.g., Miralax, fiber, stimulants, dicyclomine, hyoscyamine, tricyclic or SSRI antidepressants)

AND

1.1.4 For Linzess Only - Trial and failure, contraindication or intolerance to lubiprostone

AND

1.1.5 For Trulance Only - Trial and failure, contraindication or intolerance to lubiprostone and linaclotide

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

Product Name: Symproic, Movantik				
Diagnosis Opioid-Induced Constipation				
Approval Length	12/31/2039			
Guideline Type	Prior Authorization - All plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
MOVANTIK	MOVANTIK NALOXEGOL OXALATE TAB 12.5 MG (BASE EQUIVALENT)		Brand
MOVANTIK	NALOXEGOL OXALATE TAB 25 MG (BASE EQUIVALENT)	52580060300330	Brand
SYMPROIC	NALDEMEDINE TOSYLATE TAB 0.2 MG (BASE EQUIVALENT)	52580057200320	Brand

1	-	One	of	the	fol	lowing:
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- **1.1** All of the following:
- 1.1.1 Diagnosis of Opioid-Induced Constipation

AND

1.1.2 Member is on chronic opioid therapy

AND

1.1.3 Member is 18 years of age or older

AND

1.1.4 Trial and failure, contraindication or intolerance to a concurrent combination of a stimulant (e.g., Senna) and an osmotic laxative (e.g., Miralax)

AND

1.1.5 For Movantik Only - Trial and failure, contraindication or intolerance to lubiprostone

AND

1.1.6 For Symproic Only - Trial and failure, contraindication or intolerance to lubiprostone and naloxegol

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

Product Name: Symproic, Movantik				
Diagnosis Opioid-Induced Constipation				
Approval Length	12 month(s)			
Guideline Type Prior Authorization - IL and MN Plans				

Product Name	Generic Name	GPI	Brand/Generic
MOVANTIK	MOVANTIK NALOXEGOL OXALATE TAB 12.5 MG (BASE EQUIVALENT)		Brand
MOVANTIK	NALOXEGOL OXALATE TAB 25 MG (BASE EQUIVALENT)	52580060300330	Brand
SYMPROIC	NALDEMEDINE TOSYLATE TAB 0.2 MG (BASE EQUIVALENT)	52580057200320	Brand

Approval Criteria

- 1 One of the following:
- **1.1** All of the following:
- 1.1.1 Diagnosis of Opioid-Induced Constipation

AND

1.1.2 Member is on chronic opioid therapy

AND

1.1.3 Member is 18 years of age or older

AND

1.1.4 Trial and failure, contraindication or intolerance to a concurrent combination of a stimulant (e.g., Senna) and an osmotic laxative (e.g., Miralax)

AND

1.1.5 For Movantik Only - Trial and failure, contraindication or intolerance to lubiprostone

AND

1.1.6 For Symproic Only - Trial and failure, contraindication or intolerance to lubiprostone and naloxegol

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

Product Name: Linzess, Trulance, Motegrity, Symproic, Movantik	
Diagnosis	Metastatic Cancer
Approval Length	12 month(s)
Guideline Type Prior Authorization - MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
LINZESS	LINACLOTIDE CAP 72 MCG	52557050000110	Brand
LINZESS	LINACLOTIDE CAP 145 MCG	52557050000120	Brand
LINZESS	LINACLOTIDE CAP 290 MCG	52557050000140	Brand
MOVANTIK	NALOXEGOL OXALATE TAB 12.5 MG (BASE EQUIVALENT)	52580060300320	Brand
MOVANTIK	TIK NALOXEGOL OXALATE TAB 25 MG (BASE EQUIVALENT) 52580060300330 Brand		Brand
SYMPROIC	NALDEMEDINE TOSYLATE TAB 0.2 MG (BASE EQUIVALENT)	52580057200320	Brand

TRULANCE	PLECANATIDE TAB 3 MG	52543060000320	Brand
MOTEGRITY	PRUCALOPRIDE SUCCINATE TAB 1 MG (BASE EQUIVALENT)	52560060200320	Brand
MOTEGRITY	PRUCALOPRIDE SUCCINATE TAB 2 MG (BASE EQUIVALENT)	52560060200330	Brand

1 - Diagnosis of stage four metastatic cancer

AND

2 - Member is on opioid therapy to treat cancer-related pain with opioid-induced constipation

Product Name: Linzess	
Diagnosis Functional Constipation	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
LINZESS	LINACLOTIDE CAP 72 MCG	52557050000110	Brand
LINZESS	LINACLOTIDE CAP 145 MCG	52557050000120	Brand
LINZESS	LINACLOTIDE CAP 290 MCG	52557050000140	Brand

Approval Criteria

- 1 One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of Functional Constipation

AND

1.1.2 Member is between the age of 6 and 17 years of age

AND

1.1.3 Trial and failure, contraindication or intolerance to an adequate trial of over-the-counter osmotic laxatives (e.g., polyethylene glycol, magnesium hydroxide)

AND

1.1.4 Trial and failure, contraindication or intolerance to an adequate trial of over-the-counter stimulant laxative (e.g. bisacodyl, senna)

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

Product Name: Linzess	
Diagnosis	Functional Constipation
Approval Length	12 month(s)
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
LINZESS	LINACLOTIDE CAP 72 MCG	52557050000110	Brand
LINZESS	LINACLOTIDE CAP 145 MCG	52557050000120	Brand
LINZESS	LINACLOTIDE CAP 290 MCG	52557050000140	Brand

Approval Criteria

- 1 One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of Functional Constipation

AND

1.1.2 Member is between the age of 6 and 17 years of age

AND

1.1.3 Trial and failure, contraindication or intolerance to an adequate trial of over-the-counter osmotic laxatives (e.g., polyethylene glycol, magnesium hydroxide)

AND

1.1.4 Trial and failure, contraindication or intolerance to an adequate trial of over-the-counter stimulant laxative (e.g. bisacodyl, senna)

OR

1.2 Continuation of prior therapy with the requested drug, verified by paid claims, or medical records (e.g. chart notes), or provider attestation.

2. Revision History

Date	Notes
4/15/2024	New Program

Cimzia (certolizumab)
(ii) become more simple. While the local county and the local transition county are being the local transition county and the local transition county are being the local transition.

Prior Authorization Guideline

Guideline ID	GL-145329
Guideline Name	Cimzia (certolizumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Cimzia				
Diagnosis		Plaque Psoriasis		
Approval Length		12/31/2039*		
Guideline Type		Prior Authorization – All Plans except IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
CIMZIA	CERTOLIZUM X 200 MG/ML	MAB PEGOL PREFILLED SYRINGE KIT 2	5250502010F840	Brand
CIMZIA STARTER KIT			5250502010F860	Brand
CIMZIA CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG		52505020106420	Brand	

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g. nails, scalp, genitals, or intertriginous areas

AND

3.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days
	Second PA: Approve GPI 10 through 12/31/2039

Product Name: Cimzia	
Diagnosis Plaque Psoriasis	
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g. nails, scalp, genitals, or intertriginous areas

AND

3.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

	OR
3.2 Continuation of pr (e.g. chart notes)	ior therapy with certolizumab verified by paid claims or medical records
Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days Second PA: Approve at GPI 10 for 12 months

Product Name: Cimzia			
Diagnosis	Psoriatic Arthritis (PsA)		
Approval Length	12/31/2039*		
Guideline Type Prior Authorization – All Plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

3 - One of the following:

- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

3.2 Continuation of prior therapy with certolizumab verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cimzia		
Diagnosis	Psoriatic Arthritis (PsA)	
Approval Length	12 Month(s)*	
Guideline Type	Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days Second PA: Approve at GPI 10 for 12 months

Product Name: Cimzia		
Diagnosis	Ankylosing Spondylitis (AS)	
Approval Length	12/31/2039*	
Guideline Type	Prior Authorization – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Diagnosis of ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cimzia	
Diagnosis	Ankylosing Spondylitis (AS)
Approval Length	12 Month(s)*
Guideline Type	Prior Authorization – IL and MN Plans

ı	Product Name	Generic Name	GPI	Brand/Generic
	CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand

CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Diagnosis of ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days
	Second PA: Approve at GPI 10 for 12 months

Product Name: Cimzia	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis (RA)
Approval Length	12/31/2039*
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Diagnosis of moderate to severely active Rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - Methotrexate (MTX)**
 - Leflunomide
 - Hydroxychloroquine
 - Sulfasalazine

OR

Notes	**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant anemia, or hypersensitivity to methotrexate.

*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days
Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cimzia	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis (RA)
Approval Length	12 Month(s)*
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Diagnosis of moderate to severely active Rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - Methotrexate (MTX)**
 - Leflunomide
 - Hydroxychloroquine
 - Sulfasalazine

UK

3.2 Continuation of prior therapy with certolizumab verified by paid claims or medical records (e.g. chart notes)

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Notes	**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant anemia, or hypersensitivity to methotrexate.
	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days Second PA: Approve at GPI 10for 12 months

Product Name: Cimzia	
Diagnosis	Non-radiographic axial spondyloarthritis (nr-axSpA)
Approval Length	12/31/2039*
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

Approval Criteria

1 - Diagnosis of active non-radiographic axial spondyloarthritis (nr-axSpA) with elevated labs (e.g. C-reactive protein) and/or sacroiliitis on imaging

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

3.2 Continuation of prior therapy with certolizumab verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cimzia	
Diagnosis	Non-radiographic axial spondyloarthritis (nr-axSpA)
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

Approval Criteria

1 - Diagnosis of active non-radiographic axial spondyloarthritis (nr-axSpA) with elevated labs (e.g. C-reactive protein) and/or sacroiliitis on imaging

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days
Second PA: Approve at GPI 10 for 12 months

Product Name: Cimzia	
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)
Approval Length	12/31/2039*
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

Approval Criteria
1 - Diagnosis of moderate to severely active Crohn's disease (CD)
AND
2 - Prescribed by or in consultation with a gastroenterologist
AND
3 - One of the following:
3.1 One of the following:
3.1.1 Member is considered high-risk based on ONE of the following characteristics:
 Age less than 30 years at diagnosis Extensive anatomic involvement Perianal and/or severe rectal disease Deep ulcers Prior surgical resection Stricturing and/or penetrating behavior Fistulizing disease Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)
OR
3.1.2 Both of the following:
3.1.2.1 Member is considered low-risk
AND

3.1.2.2 One of the following:

- Trial and failure, contraindication, or intolerance to one conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with one conventional therapy

- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

OR

3.2 Continuation of prior therapy with certolizumab verified by paid claims or medical records (e.g. chart notes)

*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for 60 days Second PA: Approve at GPI 10 through 12/31/2039
Second PA. Approve at GPT to through 12/31/2039

Product Name: Cimzia	
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)
Approval Length	12 Month(s)*
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active Crohn's disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** One of the following:
- **3.1.1** Member is considered high-risk based on ONE of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease
 - Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)

OR

- **3.1.2** Both of the following:
- **3.1.2.1** Member is considered low-risk

AND

- **3.1.2.2** One of the following:
 - Trial and failure, contraindication, or intolerance to one conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
 - Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with one conventional therapy
- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve at GPI 14 with a MDD of 0.06 (3 every 56 days) for

60 days Second PA: Approve at GPI 10 for 12 months

Product Name: Cimzia			
Diagnosis Moderate to Severely Active Rheumatoid Arthritis, PSA, AS, nr-axSpA, plaque psoriasis			
Approval Length	12 month(s)		
Guideline Type	Quantity Exception - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Failure of an adherent 3-month trial of standard maintenance dosing with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)

OR

2 - Continuation of previous therapy with certolizumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

Product Name: Cimzia		
Diagnosis Moderate to Severely Active Rheumatoid Arthritis, PSA, AS, nr-axSpA, plaque psoriasis		
Approval Length	12/31/2099	
Guideline Type Quantity Exception – All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand

	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

1 - Failure of an adherent 3-month trial of standard maintenance dosing with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)

OR

2 - Continuation of previous therapy with certolizumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

Product Name: Cimzia		
Diagnosis	Crohn's disease	
Approval Length	12 month(s)	
Guideline Type	Quantity Exception - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

Approval Criteria

- 1 Both of the following:
- **1.1** Failure of a two-month trial of monthly therapy after completion of induction dosing regimen

AND

1.2 Based on subtherapeutic drug concentrations and absence (or low levels) of drug antibodies

OR

2 - Continuation of previous therapy with certolizumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes)

Product Name: Cimzia		
Diagnosis Crohn's disease		
Approval Length 12/31/2099		
Guideline Type Quantity Exception – All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
CIMZIA	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 2 X 200 MG/ML	5250502010F840	Brand
CIMZIA STARTER KIT	CERTOLIZUMAB PEGOL PREFILLED SYRINGE KIT 6 X 200 MG/ML	5250502010F860	Brand
CIMZIA	CERTOLIZUMAB PEGOL FOR INJ KIT 2 X 200 MG	52505020106420	Brand

Approval Criteria

- 1 Both of the following:
- **1.1** Failure of a two-month trial of monthly therapy after completion of induction dosing regimen

AND

1.2 Based on subtherapeutic drug concentrations and absence (or low levels) of drug antibodies

OR

2 - Continuation of previous therapy with certolizumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

2. Revision History

Date	Notes
4/17/2024	Update COT language

Clomipramine (anafranil)		
(g) The Mandrings control budgeton. Totals may be about a result, or about any body to be a purely be of source the art budget.		

Prior Authorization Guideline

Guideline ID	GL-144844
Guideline Name	Clomipramine (anafranil)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	6/15/2016
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Generic Clomipramine				
Diagnosis		Obsessive compulsive disorder:		
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name Gen		eric Name	GPI	Brand/Generic
CLOMIPRAMINE CLON		MIPRAMINE HCL CAP 25 MG	58200025100120	Generic
CLOMIPRAMINE CLOMIPR HYDROCHLORIDE		MIPRAMINE HCL CAP 25 MG	58200025100120	Generic

CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic

- 1 One of the following:
- **1.1** Trial and failure, contraindication or intolerance to, two preferred antidepressants within the Serotonin Selective Reuptake Inhibitor (SSRI) category category (e.g. citalopram, escitalopram, sertraline, fluoxetine, paroxetine)

OR

1.2 Continuation of prior therapy with clomipramine, verified by paid claims, medical records (e.g. chart notes), or provider attestation.

Product Name: Generic Clomipramine		
Diagnosis	Other mood or anxiety disorders	
Approval Length	12 month(s)	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 25 MG	58200025100120	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 25 MG	58200025100120	Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic

- 1 One of the following:
- **1.1** Trial and failure, contraindication or intolerance to, two preferred antidepressants within the Serotonin Selective Reuptake Inhibitor (SSRI) or Selective Norepinephrine Reuptake Inhibitor (SNRI) categories (e.g. citalopram, escitalopram, sertraline, fluoxetine, paroxetine, duloxetine, venlafaxine)

OR

1.2 Continuation of prior therapy with clomipramine, verified by paid claims, medical records (e.g. chart notes), or provider attestation

Product Name: Generic Clomipramine		
Diagnosis	Obsessive compulsive disorder:	
Approval Length	12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 25 MG	58200025100120	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 25 MG	58200025100120	Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic

Approval Criteria

- 1 One of the following:
- **1.1** Trial and failure, contraindication or intolerance to, two preferred antidepressants within

the Serotonin Selective Reuptake Inhibitor (SSRI) category (e.g. citalopram, escitalopram, sertraline, fluoxetine, paroxetine

OR

1.2 Continuation of prior therapy with clomipramine, verified by paid claims, medical records (e.g. chart notes), or provider attestation

Product Name: Generic Clomipramine		
Diagnosis	Other mood or anxiety disorders	
Approval Length	12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 25 MG	58200025100120	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 25 MG	58200025100120	Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 50 MG	58200025100130	Generic
CLOMIPRAMINE HCL	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic
CLOMIPRAMINE HYDROCHLORIDE	CLOMIPRAMINE HCL CAP 75 MG	58200025100140	Generic

Approval Criteria

- 1 One of the following:
- **1.1** Trial and failure, contraindication or intolerance to, two preferred antidepressants within the Serotonin Selective Reuptake Inhibitor (SSRI) or Selective Norepinephrine Reuptake Inhibitor (SNRI) categories (e.g. citalopram, escitalopram, sertraline, fluoxetine, paroxetine, duloxetine, venlafaxine)

OR

1.2 Continuation of prior therapy with clomipramine, verified by paid claims, medical records (e.g. chart notes), or provider attestation

2. Revision History

Date	Notes
3/25/2024	New Program

Codeine and Tramadol-Containing	Troducts
(g) binding were higher both as basined and a deal of the binding words around a format.	

Prior Authorization Guideline

Guideline ID	GL-144535	
Guideline Name	Codeine and Tramadol-Containing Products	
Formulary	Quartz	

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: generic Acetaminophen/Codeine, Brand Rydex,generic BUTALBITAL/ACETAMINOPHEN/CAFFEINE/CODEINE, Brand Capcof, generic Virtussin A/C, generic codeine/guaifenesin, generic Maxi-Tuss AC, generic guaiatussin AC, generic G Tussin AC, generic guaifenesin AC, Brand M-Clear WC, Brand Ninjacof-XG, generic codeine, generic Ascomp/codeine, generic BUTALBITAL/ASPIRIN/CAFFEINE/CODEINE, generic PROMETHAZINE/CODEINE, generic PROMETHAZINE VC/CODEINE brand TUXARIN ER, Generic tramadol 12 month(s) Approval Length Therapy Stage **Initial Authorization** Prior Authorization - IL and MN Plans Guideline Type **Product Name** Generic Name GPI Brand/Gener ic ACETAMINOPHEN ACETAMINOPHEN/CODEINE PHOSPHATE 6599100205031 Generic W/ CODEINE TAB 300-15 MG

A OFTANINO DUENIO O DEINE	A OFTA MINIOR USE	050040000500	
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-15 MG	6599100205031 0	Generic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE #3	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-60 MG	6599100205032 0	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-60 MG	6599100205032 0	Generic
RYDEX	PSEUDOEPHEDRIN E-BROMPHEN- CODEINE LIQ 10- 1.33-6.33 MG/5ML	4399530319092 2	Brand
BUTALBITAL/ACETAMINOPHEN/CAFFEINE/COD EINE	BUTALBITAL- ACETAMINOPHEN- CAFF W/ COD CAP 50-300-40-30 MG	6599100410011 3	Generic
BUTALBITAL/ACETAMINOPHEN/CAFFEINE/COD EINE	BUTALBITAL- ACETAMINOPHEN- CAFF W/ COD CAP 50-325-40-30 MG	6599100410011 5	Generic
CAPCOF	PHENYLEPHRINE- CHLORPHEN W/ CODEINE SYRUP 5- 2-10 MG/5ML	4399530314122 0	Brand
VIRTUSSIN A/C	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
CODEINE/GUAIFENESIN	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
MAXI-TUSS AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN/CODEINE	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIATUSSIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic

G TUSSIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN/CODEINE PHOSPHATE	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
M-CLEAR WC	GUAIFENESIN- CODEINE SOLN 100-6.33 MG/5ML	4399700228201 8	Brand
NINJACOF-XG	GUAIFENESIN- CODEINE LIQUID 200-8 MG/5ML	4399700228094 2	Brand
CODEINE SULFATE	CODEINE SULFATE TAB 15 MG	6510002020030 5	Generic
CODEINE SULFATE	CODEINE SULFATE TAB 30 MG	6510002020031 0	Generic
CODEINE SULFATE	CODEINE SULFATE TAB 60 MG	6510002020031 5	Generic
ASCOMP/CODEINE	BUTALBITAL- ASPIRIN-CAFF W/ CODEINE CAP 50- 325-40-30 MG	6599100430011 5	Generic
BUTALBITAL/ASPIRIN/CAFFEINE/CODEINE	BUTALBITAL- ASPIRIN-CAFF W/ CODEINE CAP 50- 325-40-30 MG	6599100430011 5	Generic
PROMETHAZINE/CODEINE	PROMETHAZINE W/ CODEINE SYRUP 6.25-10 MG/5ML	4399520234121 0	Generic
PROMETHAZINE VC/CODEINE	PROMETHAZINE- PHENYLEPHRINE- CODEINE SYRUP 6.25-5-10 MG/5ML	4399530310121 0	Generic
TUXARIN ER	CODEINE PHOS- CHLORPHENIRAMI NE MALEATE TAB ER 12HR 54.3-8 MG	4399520232743 0	Brand
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL TAB 50 MG	6510009510032 0	Generic
TRAMADOL HCL	TRAMADOL HCL TAB 50 MG	6510009510032 0	Generic
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL TAB 100 MG	6510009510034 0	Generic

TRAMADOL HYDROCHLORIDE	TRAMADOL HCL ORAL SOLN 5 MG/ML	6510009510200 5	Generic	
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1 - Age greater than 11 years

Product Name: generic Acetaminophen/Codeine, Brand Rydex,generic BUTALBITAL/ACETAMINOPHEN/CAFFEINE/CODEINE, Brand Capcof, generic Virtussin A/C, generic codeine/guaifenesin, generic Maxi-Tuss AC, generic guaiatussin AC, generic G Tussin AC, generic guaifenesin AC, Brand M-Clear WC, Brand Ninjacof-XG, generic codeine, generic Ascomp/codeine, generic BUTALBITAL/ASPIRIN/CAFFEINE/CODEINE, generic PROMETHAZINE/CODEINE, generic PROMETHAZINE VC/CODEINE brand TUXARIN ER, Generic tramadol

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Gener ic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-15 MG	6599100205031 0	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-15 MG	6599100205031 0	Generic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE #3	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-60 MG	6599100205032 0	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-60 MG	6599100205032 0	Generic
RYDEX	PSEUDOEPHEDRIN E-BROMPHEN-	4399530319092 2	Brand

	CODEINE LIQ 10- 1.33-6.33 MG/5ML		
BUTALBITAL/ACETAMINOPHEN/CAFFEINE/COD EINE	BUTALBITAL- ACETAMINOPHEN- CAFF W/ COD CAP 50-300-40-30 MG	6599100410011 3	Generic
BUTALBITAL/ACETAMINOPHEN/CAFFEINE/COD EINE	BUTALBITAL- ACETAMINOPHEN- CAFF W/ COD CAP 50-325-40-30 MG	6599100410011 5	Generic
CAPCOF	PHENYLEPHRINE- CHLORPHEN W/ CODEINE SYRUP 5- 2-10 MG/5ML	4399530314122 0	Brand
VIRTUSSIN A/C	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
CODEINE/GUAIFENESIN	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
MAXI-TUSS AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN/CODEINE	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIATUSSIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
G TUSSIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN/CODEINE PHOSPHATE	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
M-CLEAR WC	GUAIFENESIN- CODEINE SOLN 100-6.33 MG/5ML	4399700228201 8	Brand
NINJACOF-XG	GUAIFENESIN- CODEINE LIQUID 200-8 MG/5ML	4399700228094 2	Brand
CODEINE SULFATE	CODEINE SULFATE TAB 15 MG	6510002020030 5	Generic
CODEINE SULFATE	CODEINE SULFATE TAB 30 MG	6510002020031 0	Generic

CODEINE SULFATE	CODEINE SULFATE TAB 60 MG	6510002020031 5	Generic
ASCOMP/CODEINE	BUTALBITAL- ASPIRIN-CAFF W/ CODEINE CAP 50- 325-40-30 MG	6599100430011 5	Generic
BUTALBITAL/ASPIRIN/CAFFEINE/CODEINE	BUTALBITAL- ASPIRIN-CAFF W/ CODEINE CAP 50- 325-40-30 MG	6599100430011 5	Generic
PROMETHAZINE/CODEINE	PROMETHAZINE W/ CODEINE SYRUP 6.25-10 MG/5ML	4399520234121 0	Generic
PROMETHAZINE VC/CODEINE	PROMETHAZINE- PHENYLEPHRINE- CODEINE SYRUP 6.25-5-10 MG/5ML	4399530310121 0	Generic
TUXARIN ER	CODEINE PHOS- CHLORPHENIRAMI NE MALEATE TAB ER 12HR 54.3-8 MG	4399520232743 0	Brand
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL TAB 50 MG	6510009510032 0	Generic
TRAMADOL HCL	TRAMADOL HCL TAB 50 MG	6510009510032 0	Generic
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL TAB 100 MG	6510009510034 0	Generic
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL ORAL SOLN 5 MG/ML	6510009510200 5	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: generic Acetaminophen/Codeine, Brand Rydex,generic
BUTALBITAL/ACETAMINOPHEN/CAFFEINE/CODEINE, Brand Capcof, generic Virtussin A/C,
generic codeine/guaifenesin, generic Maxi-Tuss AC, generic guaiatussin AC, generic G Tussin
AC, generic guaifenesin AC, Brand M-Clear WC, Brand Ninjacof-XG, generic codeine, generic
Ascomp/codeine, generic BUTALBITAL/ASPIRIN/CAFFEINE/CODEINE, generic
PROMETHAZINE/CODEINE, generic PROMETHAZINE VC/CODEINE brand TUXARIN ER,
Generic tramadol

Approval Length

12/31/2039

Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Gener ic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-15 MG	6599100205031 0	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-15 MG	6599100205031 0	Generic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE #3	ACETAMINOPHEN W/ CODEINE TAB 300-30 MG	6599100205031 5	Generic
ACETAMINOPHEN/CODEINE PHOSPHATE	ACETAMINOPHEN W/ CODEINE TAB 300-60 MG	6599100205032 0	Generic
ACETAMINOPHEN/CODEINE	ACETAMINOPHEN W/ CODEINE TAB 300-60 MG	6599100205032 0	Generic
RYDEX	PSEUDOEPHEDRIN E-BROMPHEN- CODEINE LIQ 10- 1.33-6.33 MG/5ML	4399530319092 2	Brand
BUTALBITAL/ACETAMINOPHEN/CAFFEINE/COD EINE	BUTALBITAL- ACETAMINOPHEN- CAFF W/ COD CAP 50-300-40-30 MG	6599100410011 3	Generic
BUTALBITAL/ACETAMINOPHEN/CAFFEINE/COD EINE	BUTALBITAL- ACETAMINOPHEN- CAFF W/ COD CAP 50-325-40-30 MG	6599100410011 5	Generic
CAPCOF	PHENYLEPHRINE- CHLORPHEN W/ CODEINE SYRUP 5- 2-10 MG/5ML	4399530314122 0	Brand
VIRTUSSIN A/C	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
CODEINE/GUAIFENESIN	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
MAXI-TUSS AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic

GUAIFENESIN/CODEINE	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIATUSSIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
G TUSSIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN AC	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
GUAIFENESIN/CODEINE PHOSPHATE	GUAIFENESIN- CODEINE SOLN 100-10 MG/5ML	4399700228202 0	Generic
M-CLEAR WC	GUAIFENESIN- CODEINE SOLN 100-6.33 MG/5ML	4399700228201 8	Brand
NINJACOF-XG	GUAIFENESIN- CODEINE LIQUID 200-8 MG/5ML	4399700228094 2	Brand
CODEINE SULFATE	CODEINE SULFATE TAB 15 MG	6510002020030 5	Generic
CODEINE SULFATE	CODEINE SULFATE TAB 30 MG	6510002020031 0	Generic
CODEINE SULFATE	CODEINE SULFATE TAB 60 MG	6510002020031 5	Generic
ASCOMP/CODEINE	BUTALBITAL- ASPIRIN-CAFF W/ CODEINE CAP 50- 325-40-30 MG	6599100430011 5	Generic
BUTALBITAL/ASPIRIN/CAFFEINE/CODEINE	BUTALBITAL- ASPIRIN-CAFF W/ CODEINE CAP 50- 325-40-30 MG	6599100430011 5	Generic
PROMETHAZINE/CODEINE	PROMETHAZINE W/ CODEINE SYRUP 6.25-10 MG/5ML	4399520234121 0	Generic
PROMETHAZINE VC/CODEINE	PROMETHAZINE- PHENYLEPHRINE- CODEINE SYRUP 6.25-5-10 MG/5ML	4399530310121 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL CAP ER 24HR BIPHASIC RELEASE 100 MG	6510009510707 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL CAP ER 24HR BIPHASIC RELEASE 200 MG	6510009510708 0	Generic

TRAMADOL HCL ER	TRAMADOL HCL CAP ER 24HR BIPHASIC RELEASE 300 MG	6510009510709 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL TAB ER 24HR 100 MG	6510009510752 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL TAB ER 24HR 200 MG	6510009510753 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL TAB ER 24HR 300 MG	6510009510754 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL TAB ER 24HR BIPHASIC RELEASE 100 MG	6510009510756 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL TAB ER 24HR BIPHASIC RELEASE 200 MG	6510009510757 0	Generic
TRAMADOL HCL ER	TRAMADOL HCL TAB ER 24HR BIPHASIC RELEASE 300 MG	6510009510758 0	Generic
TUXARIN ER	CODEINE PHOS- CHLORPHENIRAMI NE MALEATE TAB ER 12HR 54.3-8 MG	4399520232743 0	Brand
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL TAB 50 MG	6510009510032 0	Generic
TRAMADOL HCL	TRAMADOL HCL TAB 50 MG	6510009510032 0	Generic
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL TAB 100 MG	6510009510034 0	Generic
TRAMADOL HYDROCHLORIDE	TRAMADOL HCL ORAL SOLN 5 MG/ML	6510009510200 5	Generic

1 - Age greater than11 years

Date	Notes
3/18/2024	Updated product name

Compounded Hormones						
The State Programmer for England. The Birth Rep State Seen Record, resident, or delete	ne valg der let is pain a fir en enrich er indien.					

Guideline ID	GL-145620
Guideline Name	Compounded Hormones
Formulary	Quartz

Guideline Note:

Effective Date:	4/11/2024
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1. Criteria

Product Name: Compounded progesterone to maintain pregnancy in the first trimester				
Approval Length		4 month(s)		
Guideline Type		Prior Authorization - All plans except IL and MN		
Product Name	Gene	Generic Name GPI Brand/Generic		Brand/Generic
PROGESTERONE	PROG	ESTERONE CAP 100 MG	26000040000120	Generic
PROGESTERONE	PROG	ESTERONE CAP 200 MG	26000040000140	Generic
PROGESTERONE	PROG	ESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic
PROGESTERONE WETTABLE	PROG	ESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE MILLED	PROG	ESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (YAM)	PROG	ESTERONE (BULK) POWDER	96727643212900	Brand

PROGESTERONE PROGESTERONE (BULK) POWDER WETTABLE (SOY)	96727643212900	Brand
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1 - Medication will be used to maintain pregnancy

AND

2 - Submission of medical records (e.g., chart notes) documenting the woman is currently pregnant in her 1st trimester

Product Name: Compounded progesterone to maintain pregnancy beyond the first trimester		
Approval Length 6 month(s)		
Guideline Type Prior Authorization – All plans except MN and IL		

Product Name	Generic Name	GPI	Brand/Generic
PROGESTERONE	PROGESTERONE CAP 100 MG	26000040000120	Generic
PROGESTERONE	PROGESTERONE CAP 200 MG	26000040000140	Generic
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic
PROGESTERONE WETTABLE	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE MILLED	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (YAM)	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (SOY)	PROGESTERONE (BULK) POWDER	96727643212900	Brand

Approval Criteria

1 - Medication will be used to maintain pregnancy

AND

2 - Submission of medical records (e.g., chart notes) documenting the woman has a singleton pregnancy

AND

3 - Woman is beyond the 1st trimester

AND

4 - Submission of medical records (e.g., chart notes) documenting a history of preterm birth

Product Name: Compounded progesterone to maintain pregnancy		
Approval Length 12 month(s)		
Guideline Type Prior Authorization - IL and MN plans		

Product Name	Generic Name	GPI	Brand/Generic
PROGESTERONE	PROGESTERONE CAP 100 MG	26000040000120	Generic
PROGESTERONE	PROGESTERONE CAP 200 MG	26000040000140	Generic
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic
PROGESTERONE WETTABLE	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE MILLED	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (YAM)	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (SOY)	PROGESTERONE (BULK) POWDER	96727643212900	Brand

Approval Criteria

- 1 BOTH of the following:
 - Medication will be used to maintain pregnancy
 - Submission of medical records (e.g., chart notes) documenting the woman is currently pregnant in her 1st trimester

OR

2 - All of the following:

- Medication will be used to maintain pregnancy
- Submission of medical records (e.g., chart notes) woman has a singleton pregnancy
- Woman is beyond the 1st trimester
- Submission of medical records (e.g., chart notes) documenting a history of preterm birth

Product Name: Compounded progesterone to treat infertility				
Approval Length	12 month(s)			
Guideline Type	Prior Authorization - IL Plans			
Product Name Gen	eric Name	GPI	Brand/Generic	

Product Name	Generic Name	GPI	Brand/Generic
PROGESTERONE	PROGESTERONE CAP 100 MG	26000040000120	Generic
PROGESTERONE	PROGESTERONE CAP 200 MG	26000040000140	Generic
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic
PROGESTERONE WETTABLE	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE MILLED	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (YAM)	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (SOY)	PROGESTERONE (BULK) POWDER	96727643212900	Brand

Approval Criteria

1 - Provider attests patient has infertility coverage as outlined in Illinois Insurance Code 215 ILCS 5/356m

Product Name: Estrogen, testosterone, and progesterone use outside of pregnancy				
Approval Length 12 month(s)				
Therapy Stage	Initial Authorization			
Guideline Type	Guideline Type Prior Authorization			

Product Name	Generic Name	GPI	Brand/Generic	
PROGESTERONE	PROGESTERONE CAP 100 MG	26000040000120	Generic	
PROMETRIUM	PROGESTERONE CAP 100 MG	26000040000120	Brand	
PROGESTERONE	PROGESTERONE CAP 200 MG	26000040000140	Generic	
PROMETRIUM	PROGESTERONE CAP 200 MG	26000040000140	Brand	
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic	
PROGESTERONE 10% KIT	*PROGESTERONE MICRONIZED TD CREAM 10% (CMPD KIT)*	26000040103730	Brand	
EC-RX PROGESTERONE 10%	*PROGESTERONE MICRONIZED TD CREAM 10% (CMPD KIT)*	26000040103730	Brand	
PROGESTERONE WETTABLE	PROGESTERONE (BULK) POWDER	96727643212900	Brand	
PROGESTERONE MILLED	PROGESTERONE (BULK) POWDER	96727643212900	Brand	
PROGESTERONE WETTABLE (YAM)	PROGESTERONE (BULK) POWDER	96727643212900	Brand	
PROGESTERONE WETTABLE (SOY)	PROGESTERONE (BULK) POWDER	96727643212900	Brand	
PROGESTERONE	PROGESTERONE (BULK) POWDER	96727643212900	Brand	
PROGESTERONE ULTRA MICRONIZED	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand	
PROGESTERONE MICRONIZED	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand	
PROGESTERONE MICRONIZED PREMIUM	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand	
PROGESTERONE MICRONIZED (SOY)	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand	
PROGESTERONE MICRONIZED (YAM)	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand	
TESTOSTERONE	TESTOSTERONE TD SOLN 30 MG/ACT	23100030002020	Generic	
TESTOSTERONE TOPICAL SOLUTION	TESTOSTERONE TD SOLN 30 MG/ACT	23100030002020	Generic	
TESTOSTERONE	TESTOSTERONE TD GEL 25 MG/2.5GM (1%)	23100030004025	Generic	
TESTIM	TESTOSTERONE TD GEL 50 MG/5GM (1%)	23100030004030	Brand	
TESTOSTERONE	TESTOSTERONE TD GEL 50 MG/5GM (1%)	23100030004030	Generic	

VOOELVO	TEOTOOTEDONE TO OEL 50 MO/50M (40)		Durand
VOGELXO	TESTOSTERONE TD GEL 50 MG/5GM (1%)	23100030004030	Brand
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 12.5 MG/ACT (1%)	23100030004040	Generic
VOGELXO PUMP	TESTOSTERONE TD GEL 12.5 MG/ACT (1%)	23100030004040	Brand
TESTOSTERONE	TESTOSTERONE TD GEL 20.25 MG/1.25GM (1.62%)	23100030004044	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 40.5 MG/2.5GM (1.62%)	23100030004047	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Generic
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Generic
ANDROGEL PUMP	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Brand
FORTESTA	TESTOSTERONE TD GEL 10MG/ACT (2%)	23100030004070	Brand
TESTOSTERONE	TESTOSTERONE TD GEL 10MG/ACT (2%)	23100030004070	Generic
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 25 MG	23100030008910	Brand
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 50 MG	23100030008915	Brand
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 100 MG	23100030008930	Brand
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 200 MG	23100030008940	Brand
DEPO- TESTOSTERONE	TESTOSTERONE CYPIONATE IM INJ IN OIL 100 MG/ML	23100030102010	Brand
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 100 MG/ML	23100030102010	Generic
DEPO- TESTOSTERONE	TESTOSTERONE CYPIONATE IM INJ IN OIL 200 MG/ML	23100030102015	Brand
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 200 MG/ML	23100030102015	Generic
TESTOSTERONE CYPIONATE	TESTOSTERONE CYP IM OR SUBCUTANEOUS INJ IN OIL 200 MG/ML	23100030102070	Brand
TESTOSTERONE ENANTHATE	TESTOSTERONE ENANTHATE IM INJ IN OIL 200 MG/ML	23100030202010	Generic
TESTOSTERONE NON- MICRONIZED SOY	TESTOSTERONE (BULK) POWDER	96805050502900	Brand
TESTOSTERONE MICRONIZED SOY	TESTOSTERONE (BULK) POWDER	96805050502900	Brand
TESTOSTERONE NON- MICRONIZED	TESTOSTERONE (BULK) POWDER	96805050502900	Brand
TESTOSTERONE	TESTOSTERONE (BULK) POWDER	96805050502900	Brand

TESTOSTERONE MICRONIZED	TESTOSTERONE MICRONIZED (BULK) POWDER	96805050522900	Brand
TESTOSTERONE MICRONIZED (YAM)	TESTOSTERONE MICRONIZED (BULK) 96805050522900 POWDER		Brand
TESTOSTERONE MICRONIZED (SOY)	TESTOSTERONE MICRONIZED (BULK) POWDER	96805050522900	Brand
TESTOSTERONE MICRONIZED YAM	TESTOSTERONE MICRONIZED (BULK) CRYSTALS	96805050523800	Brand
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE (BULK) POWDER	96805050552900	Brand
TESTOSTERONE ENANTHATE	TESTOSTERONE ENANTHATE (BULK) POWDER	96805050602900	Brand
ESTRADIOL BENZOATE	ESTRADIOL BENZOATE (BULK) POWDER	96507860062900	Brand
ESTRADIOL CYPIONATE	ESTRADIOL CYPIONATE (BULK) POWDER	96507860092900	Brand
ELESTRIN	ESTRADIOL GEL 0.06% (0.52 MG/0.87 GM METERED-DOSE PUMP)	24000035004008	Brand
ESTROGEL	ESTRADIOL GEL 0.06% (0.75 MG/1.25 GM METERED-DOSE PUMP)	24000035004010	Brand
ESTRADIOL	ESTRADIOL MICRONIZED (BULK) POWDER	96507860172900	Brand
ESTRADIOL MICRONIZED	ESTRADIOL MICRONIZED (BULK) POWDER	96507860172900	Brand
ESTRADIOL	ESTRADIOL TAB 0.5 MG	24000035000303	Generic
ESTRADIOL	ESTRADIOL TAB 1 MG	24000035000305	Generic
ESTRADIOL	ESTRADIOL TAB 2 MG	24000035000310	Generic
ESTRADIOL	ESTRADIOL TD GEL 0.25 MG/0.25GM (0.1%)	24000035004035	Generic
ESTRADIOL	ESTRADIOL TD GEL 0.5 MG/0.5GM (0.1%)	24000035004040	Generic
ESTRADIOL	ESTRADIOL TD GEL 0.75 MG/0.75GM (0.1%)	24000035004042	Generic
ESTRADIOL	ESTRADIOL TD GEL 1 MG/GM (0.1%)	24000035004045	Generic
ESTRADIOL	ESTRADIOL TD GEL 1.25 MG/1.25GM (0.1%)	24000035004050	Generic
ESTRADIOL	ESTRADIOL VAGINAL CREAM 0.1 MG/GM	55350020003705	Generic
ESTRADIOL	ESTRADIOL VAGINAL TAB 10 MCG	55350020000310	Generic
YUVAFEM	ESTRADIOL VAGINAL TAB 10 MCG	55350020000310	Generic
ESTRADIOL VALERATE	ESTRADIOL VALERATE (BULK) CRYSTALS	96507860243800	Brand
ESTRADIOL VALERATE	ESTRADIOL VALERATE (BULK) POWDER	96507860242900	Brand

ESTRIOL	ESTRIOL MICRONIZED (BULK) POWDER	96507861572900	Brand
ESTRIOL MICRONIZED	ESTRIOL MICRONIZED (BULK) POWDER	96507861572900	Brand
ESTRONE	ESTRONE (BULK) CRYSTALS	96507862423800	Brand
ESTRONE	ESTRONE (BULK) POWDER	96507862422900	Brand
CRINONE	PROGESTERONE VAGINAL GEL 4%	55370060004010	Brand
CRINONE	PROGESTERONE VAGINAL GEL 8%	55370060004020	Brand
FIRST- PROGESTERONE VGS 100 COMPOUNDING KIT	PROGESTERONE VAGINAL SUPPOSITORY 100 MG	55370060005210	Brand
FIRST- PROGESTERONE VGS 200 COMPOUNDING KIT	PROGESTERONE VAGINAL SUPPOSITORY 200 MG	55370060005220	Brand

1 - Trial and	failure to a	all preferred	alternatives	available	on the	formulary	of the r	equested
hormone								

AND

2 - Meets off-label criteria

AND

- **3** For testosterone only, both of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting a diagnosis of primary or secondary hypogonadism or mixed hypogonadism that clinically appropriate laboratory data demonstrate androgen deficiency*

AND

3.2 Member is symptomatic with symptoms other than sexual dysfunction

workers) below the lower limit of normal as defined by the laborate eference range. A single low testosterone is not diagnostic for and	Notes	* Androgen deficiency is defined as a fasting, morning testosterone le vel (drawn between 7 and 10 AM or within 3 hours of waking for shift workers) below the lower limit of normal as defined by the laboratory r eference range. A single low testosterone is not diagnostic for androg en deficiency and must be confirmed with a second fasting, morning testosterone level.
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Product Name: Estrogen, testosterone, and progesterone use outside of pregnancy				
Approval Length	12 month(s)			
Therapy Stage	Reauthorization			
Guideline Type	Prior Authorization			

Product Name	Generic Name	GPI	Brand/Generic
PROGESTERONE	PROGESTERONE CAP 100 MG	26000040000120	Generic
PROMETRIUM	PROGESTERONE CAP 100 MG	26000040000120	Brand
PROGESTERONE	PROGESTERONE CAP 200 MG	26000040000140	Generic
PROMETRIUM	PROGESTERONE CAP 200 MG	26000040000140	Brand
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic
PROGESTERONE 10% KIT	*PROGESTERONE MICRONIZED TD CREAM 10% (CMPD KIT)*	26000040103730	Brand
EC-RX PROGESTERONE 10%	*PROGESTERONE MICRONIZED TD CREAM 10% (CMPD KIT)*	26000040103730	Brand
PROGESTERONE WETTABLE	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE MILLED	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (YAM)	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE WETTABLE (SOY)	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE	PROGESTERONE (BULK) POWDER	96727643212900	Brand
PROGESTERONE ULTRA MICRONIZED	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand
PROGESTERONE MICRONIZED	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand
PROGESTERONE MICRONIZED PREMIUM	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand

PROGESTERONE MICRONIZED (SOY)	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand
PROGESTERONE MICRONIZED (YAM)	PROGESTERONE MICRONIZED (BULK) POWDER	96727643252900	Brand
TESTOSTERONE	TESTOSTERONE TD SOLN 30 MG/ACT	23100030002020	Generic
TESTOSTERONE TOPICAL SOLUTION	TESTOSTERONE TD SOLN 30 MG/ACT	23100030002020	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 25 MG/2.5GM (1%)	23100030004025	Generic
TESTIM	TESTOSTERONE TD GEL 50 MG/5GM (1%)	23100030004030	Brand
TESTOSTERONE	TESTOSTERONE TD GEL 50 MG/5GM (1%)	23100030004030	Generic
VOGELXO	TESTOSTERONE TD GEL 50 MG/5GM (1%)	23100030004030	Brand
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 12.5 MG/ACT (1%)	23100030004040	Generic
VOGELXO PUMP	TESTOSTERONE TD GEL 12.5 MG/ACT (1%)	23100030004040	Brand
TESTOSTERONE	TESTOSTERONE TD GEL 20.25 MG/1.25GM (1.62%)	23100030004044	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 40.5 MG/2.5GM (1.62%)	23100030004047	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Generic
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Generic
ANDROGEL PUMP	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Brand
FORTESTA	TESTOSTERONE TD GEL 10MG/ACT (2%)	23100030004070	Brand
TESTOSTERONE	TESTOSTERONE TD GEL 10MG/ACT (2%)	23100030004070	Generic
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 25 MG	23100030008910	Brand
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 50 MG	23100030008915	Brand
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 100 MG	23100030008930	Brand
TESTOSTERONE	TESTOSTERONE IMPLANT PELLETS 200 MG	23100030008940	Brand
DEPO- TESTOSTERONE	TESTOSTERONE CYPIONATE IM INJ IN OIL 100 MG/ML	23100030102010	Brand
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 100 MG/ML	23100030102010	Generic
DEPO- TESTOSTERONE	TESTOSTERONE CYPIONATE IM INJ IN OIL 200 MG/ML	23100030102015	Brand
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 200 MG/ML	23100030102015	Generic

TESTOSTERONE CYPIONATE	TESTOSTERONE CYP IM OR SUBCUTANEOUS INJ IN OIL 200 MG/ML	23100030102070	Brand
TESTOSTERONE ENANTHATE	TESTOSTERONE ENANTHATE IM INJ IN OIL 23100030202010 Get 200 MG/ML		Generic
TESTOSTERONE NON- MICRONIZED SOY	TESTOSTERONE (BULK) POWDER	DSTERONE (BULK) POWDER 96805050502900 Brand	
TESTOSTERONE MICRONIZED SOY	TESTOSTERONE (BULK) POWDER	96805050502900	Brand
TESTOSTERONE NON- MICRONIZED	TESTOSTERONE (BULK) POWDER	96805050502900	Brand
TESTOSTERONE	TESTOSTERONE (BULK) POWDER	96805050502900	Brand
TESTOSTERONE MICRONIZED	TESTOSTERONE MICRONIZED (BULK) POWDER	96805050522900	Brand
TESTOSTERONE MICRONIZED (YAM)	TESTOSTERONE MICRONIZED (BULK) POWDER	96805050522900	Brand
TESTOSTERONE MICRONIZED (SOY)	TESTOSTERONE MICRONIZED (BULK) POWDER	96805050522900	Brand
TESTOSTERONE MICRONIZED YAM	TESTOSTERONE MICRONIZED (BULK) CRYSTALS	96805050523800	Brand
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE (BULK) POWDER	96805050552900	Brand
TESTOSTERONE ENANTHATE	TESTOSTERONE ENANTHATE (BULK) POWDER	96805050602900	Brand
ESTRADIOL BENZOATE	ESTRADIOL BENZOATE (BULK) POWDER	96507860062900	Brand
ESTRADIOL CYPIONATE	ESTRADIOL CYPIONATE (BULK) POWDER	96507860092900	Brand
ELESTRIN	ESTRADIOL GEL 0.06% (0.52 MG/0.87 GM METERED-DOSE PUMP)	24000035004008	Brand
ESTROGEL	ESTRADIOL GEL 0.06% (0.75 MG/1.25 GM METERED-DOSE PUMP)	24000035004010	Brand
ESTRADIOL	ESTRADIOL MICRONIZED (BULK) POWDER	96507860172900	Brand
ESTRADIOL MICRONIZED	ESTRADIOL MICRONIZED (BULK) POWDER	96507860172900	Brand
ESTRADIOL	ESTRADIOL TAB 0.5 MG	24000035000303	Generic
ESTRADIOL	ESTRADIOL TAB 1 MG	24000035000305	Generic
ESTRADIOL	ESTRADIOL TAB 2 MG	24000035000310	Generic
ESTRADIOL	ESTRADIOL TD GEL 0.25 MG/0.25GM (0.1%)	24000035004035	Generic

ESTRADIOL	ESTRADIOL TD GEL 0.5 MG/0.5GM (0.1%)	24000035004040	Generic
ESTRADIOL	ESTRADIOL TD GEL 0.75 MG/0.75GM (0.1%)	24000035004042	Generic
ESTRADIOL	ESTRADIOL TD GEL 1 MG/GM (0.1%)	24000035004045	Generic
ESTRADIOL	ESTRADIOL TD GEL 1.25 MG/1.25GM (0.1%)	24000035004050	Generic
ESTRADIOL	ESTRADIOL VAGINAL CREAM 0.1 MG/GM	55350020003705	Generic
ESTRADIOL	ESTRADIOL VAGINAL TAB 10 MCG	55350020000310	Generic
YUVAFEM	ESTRADIOL VAGINAL TAB 10 MCG	55350020000310	Generic
ESTRADIOL VALERATE	ESTRADIOL VALERATE (BULK) CRYSTALS	96507860243800	Brand
ESTRADIOL VALERATE	ESTRADIOL VALERATE (BULK) POWDER	96507860242900	Brand
ESTRIOL	ESTRIOL MICRONIZED (BULK) POWDER	96507861572900	Brand
ESTRIOL MICRONIZED	ESTRIOL MICRONIZED (BULK) POWDER	96507861572900	Brand
ESTRONE	ESTRONE (BULK) CRYSTALS	96507862423800	Brand
ESTRONE	ESTRONE (BULK) POWDER	96507862422900	Brand
CRINONE	PROGESTERONE VAGINAL GEL 4%	55370060004010	Brand
CRINONE	PROGESTERONE VAGINAL GEL 8%	55370060004020	Brand
FIRST- PROGESTERONE VGS 100 COMPOUNDING KIT	PROGESTERONE VAGINAL SUPPOSITORY 100 MG	55370060005210	Brand
FIRST- PROGESTERONE VGS 200 COMPOUNDING KIT	PROGESTERONE VAGINAL SUPPOSITORY 200 MG	55370060005220	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member demonstrates positive clinical response to therapy

Notes	* Androgen deficiency is defined as a fasting, morning testosterone le vel (drawn between 7 and 10 AM or within 3 hours of waking for shift workers) below the lower limit of normal as defined by the laboratory r eference range. A single low testosterone is not diagnostic for androg en deficiency and must be confirmed with a second fasting, morning
	testosterone level.

Date	Notes
4/11/2024	Update Compounded Progesterone for infertility section

Compounded Prescriptions	
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Guideline ID	GL-129124	
Guideline Name	ompounded Prescriptions	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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Note:

These criteria will be applied only if a compound claim requires prior authorization (e.g., most expensive ingredient requires prior authorization)

1. Criteria

Product Name: 0	Compo	unded Prescription		
Approval Length		12 month(s)		
Guideline Type Prior Authorization – MN plans only				
Product Name	ne Generic Name		GPI	Brand/Generic
A	•_			
Approval Criter	та			
1 - For Minnesota plans only - One of the following:				

1.1 Both of the following:

1.1.1 The compound is prescribed for a member with emotional disturbance or mental illness

AND

1.1.2 One of the following:

- Submission of medical records (e.g., chart notes) documenting that all equivalent drugs in the formulary for each active ingredient were considered and it has been determined that the compound prescribed will best treat the person's condition
- For continuation of care (formulary changes or new member) the member has been treated for 90 days prior to the change, the medication is working, and the prescriber attests that the compound prescribed will best treat the member's condition.

OR

1.2 ALL of the following:

- Stage four metastatic cancer and prescribed drug is used for cancer related treatment including but not limited to: pain, constipation, nausea, fatigue related to chemotherapy or bacterial, fungal or viral infection
- Medication is not commercially available in a formulation that is suitable for the person
- Medication is on the formulary or the medication is considered medically necessary by Quartz
- None of the products in the compound are otherwise excluded from coverage as defined by the person's benefit

OR

1.3 All of the following:

- Each active ingredient in the compounded drug is FDA-approved or national compendia* supported for the condition being treated
- The therapeutic amounts are supported by national compendia* or two peer-reviewed literature for the condition being treated in the requested route of delivery
- Compound is not commercially available in a formulation that is suitable for the person
- Each active ingredient in the compounded drug is on the formulary or meets the nonformulary criteria
- None of the active ingredient(s) in the compound are otherwise excluded from coverage as defined by the person's benefit

 None of the active ingredient(s) in the compound are experimental or limited by the FDA to investigational use only

Product Name: Compo	unded Prescription		
Approval Length	12 month(s)		
Guideline Type	Prior Authorization – IL plans only		
Product Name Gene	ric Name	GPI	Brand/Generic

Approval Criteria

1 - For Illinois plans only - ONE of the following:

1.1 ALL of the following:

- Diagnosis of long-term treatment of tick-borne disease
- Medication is not commercially available in a formulation that is suitable for the person
- Medication is on the formulary or the medication is considered medically necessary by Quartz
- None of the products in the compound are otherwise excluded from coverage as defined by the person's benefit

OR

1.2 ALL of the following:

- Request is for a medication for a mental health condition under the mental and behavioral disorder chapter of the International Classification of Disease or is listed in the most recent version of the Diagnostic and Statistical Manual of Mental Disorders
- Medication is not commercially available in a formulation that is suitable for the person
- Medication is on the formulary or the medication is considered medically necessary by Quartz
- None of the products in the compound are otherwise excluded from coverage as defined by the person's benefit
- Determination should not be more restrictive than for non-behavioral health or substance use disorder diagnosis

OR

1.3 BOTH of the following:

- Request is for a medication for treating a substance use disorder
- Determination should be based on criteria established by American Society of Addiction Medicine and should not be more restrictive than non-behavioral health or substance use disorder diagnosis

OR

1.4 All of the following:

- Each active ingredient in the compounded drug is FDA-approved or national compendia* supported for the condition being treated
- The therapeutic amounts are supported by national compendia* or two peer-reviewed literature for the condition being treated in the requested route of delivery
- Compound is not commercially available in a formulation that is suitable for the person
- Each active ingredient in the compounded drug is on the formulary or meets the nonformulary criteria
- None of the active ingredient(s) in the compound are otherwise excluded from coverage as defined by the person's benefit
- None of the active ingredient(s) in the compound are experimental or limited by the FDA to investigational use only

Product Name: Compo	unded Prescription		
Approval Length	12 month(s)		
Guideline Type	Prior Authorization – All plans except IL and MN		
Product Name Gener	ric Name	GPI	Brand/Generic

Approval Criteria

1 - Each active ingredient in the compounded drug is FDA-approved or national compendia* supported for the condition being treated

AND

2 - The therapeutic amounts are supported by national compendia* or two peer-reviewed literature for the condition being treated in the requested route of delivery

AND

3 - Compound is not commercially available in a formulation that is suitable for the person

AND

4 - Each active ingredient in the compounded drug is on the formulary or meets the nonformulary criteria

AND

5 - None of the active ingredient(s) in the compound are otherwise excluded from coverage as defined by the person's benefit

AND

6 - None of the active ingredient(s) in the compound are experimental or limited by the FDA to investigational use only

2. Background

*Compendia Requirements For all non-antineoplastic medications • American Hospital Formulary Service Drug Information (AHFSDI); OR • FDA Uses/Non-FDA Uses section in DRUGDEX Evaluation with a Strength of Recommendation rating of Ilb or better (see DRUGDEX Strength of Recommendation table below); OR

	One major peer reviewed medical journal submitted by the prescriber that presents data supporting the proposed off-label use or uses as generally safe and effective unless there is clear and convincing contradictory evidence presented in a major peer-reviewed medical journal
For an antineoplastic medication	American Hospital Formulary Service Drug Information (AHFSDI); OR
	National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B (see NCCN Categories of Evidence and Consensus table below); OR
	FDA Uses/Non-FDA Uses section in DRUGDEX Evaluation with a Strength of Recommendation rating of IIb or better (see DRUGDEX Strength of Recommendation table below); OR
	Clinical Pharmacology (Gold Standard); OR
	One peer-reviewed published medical literature submitted by the prescriber:
	American Journal of Medicine
	Annals of Internal Medicine
	o Annals of Oncology
	o Annals of Surgical Oncology
	 Biology of Blood and Marrow Transplantation
	o Blood

 Bone Marrow Transplantation
 British Journal of Cancer
 British Journal of Hematology
 British Medical Journal
o Cancer
 Clinical Cancer Research
o Drugs
 European Journal of Cancer (formerly the European Journal of Cancer and Clinical Oncology)
 Gynecologic Oncology
 International Journal of Radiation, Oncology, Biology, and Physics
 The Journal of the American Medical Association
 Journal of Clinical Oncology
 Journal of the National Cancer Institute
 Journal of the National Comprehensive Cancer Network (NCCN)
 o Journal of Urology
∘ Lancet
 Lancet Oncology
o Leukemia
 The New England Journal of Medicine
o Radiation Oncology
Wolters Kluwer Lexi-Drugs rated as "Evidence Level A" with a "Strong" recommendation (see Lexi-Drugs Strength of Recommendation table below)

DRUGDEX Strength of Recommendation:

Class	Recommendation	Description
Class I	Recommended	The given test or treatment has been proven useful, and should be performed or administered.
Class IIa	Recommended, In Most Cases	The given test or treatment is generally considered to be useful, and is indicated in most cases.
Class IIb	Recommended, in Some Cases	The given test or treatment may be useful, and is indicated in some, but not most, cases.
Class III	Not Recommended	The given test or treatment is not useful, and should be avoided
Class Indeterminate	Evidence Inconclusive	

NCCN Categories of Evidence and Consensus:

Category	Level of Consensus
1	Based upon high-level evidence, there is uniform NCCN consensus that the intervention is appropriate.
2A	Based upon lower-level evidence, there is uniform NCCN consensus that the intervention is appropriate.
2B	Based upon lower-level evidence, there is NCCN consensus that the intervention is appropriate.
3	Based upon any level of evidence, there is major NCCN disagreement that the intervention is appropriate.

Lexi-Drugs: Strength of Recommendation for Inclusion in Lexi-Drugs for Oncology Off-Label Use and Level of Evidence Scale for Oncology Off-Label Use

Strength of Recommendation for Inclusion

Strong (for proposed off-label use)	The evidence persuasively supports the off-label use (ie, Level of Evidence A).
Equivocal (for proposed off-label use)	The evidence to support the off-label use is of uncertain clinical significance (ie, Level of Evidence B, C). Additional studies may be necessary to further define the role of this medication for the off-label use.
Against proposed off-label use	The evidence either advocates against the off-label use or suggests a lack of support for the off-label use (independent of Level of Evidence). Additional studies are necessary to define the role of this medication for the off-label use.

Level of Evidence Scale for Oncology Off-Label Use

A	Consistent evidence from well-performed randomized, controlled trials or overwhelming evidence of some other form (eg, results of the introduction of penicillin treatment) to support off-label use. Further research is unlikely to change confidence in the estimate of benefit.
В	Evidence from randomized, controlled trials with important limitations (eg, inconsistent results, methodologic flaws, indirect, imprecise); or very strong evidence of some other research design. Further research (if performed) is likely to have an impact on confidence in the estimate of benefit and risk and may change the estimate.
С	Evidence from observational studies (eg, retrospective case series/reports providing significant impact on patient care); unsystematic clinical experience; or potentially flawed randomized, controlled trials (eg, when limited options exist for condition). Any estimate of effect is uncertain.

G	Use has been substantiated by inclusion in at least one evidence-based or consensus-based clinical practice guideline.

Date	Notes
12/8/2023	2024 New Implementation

Corla	Corlanor (ivabradine)				
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Guideline ID	GL-129113	
Guideline Name	Corlanor (ivabradine)	
Formulary	• Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Corlanor		
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - ALL Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CORLANOR	IVABRADINE HCL TAB 5 MG (BASE EQUIV)	40700035100320	Brand
CORLANOR	IVABRADINE HCL TAB 7.5 MG (BASE EQUIV)	40700035100330	Brand
CORLANOR	IVABRADINE HCL ORAL SOLN 5 MG/5ML (BASE EQUIV)	40700035102020	Brand

Approval Criteria

1 - ONE of the following:

- **1.1** ALL of the following:
- 1.1.1 Diagnosis of stable, symptomatic heart failure in sinus rhythm

AND

- **1.1.2** Both of the following:
 - Left ventricular ejection fraction less than or equal to 35%
 - Resting heart rate greater than or equal to 70 beats per minute

AND

1.1.3 Prescribed by or in consultation with a cardiologist

OR

- **1.2** BOTH of the following:
- 1.2.1 Diagnosis of Inappropriate Sinus Tachycardia

AND

- **1.2.2** One of the following:
- **1.2.2.1** Member has symptoms despite use of maximally tolerated beta blocker therapy

OR

1.2.2.2 Member has contraindication to beta blocker use

Product Name: Corlanor	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
CORLANOR	IVABRADINE HCL TAB 5 MG (BASE EQUIV)	40700035100320	Brand
CORLANOR	IVABRADINE HCL TAB 7.5 MG (BASE EQUIV)	40700035100330	Brand
CORLANOR	IVABRADINE HCL ORAL SOLN 5 MG/5ML (BASE EQUIV)	40700035102020	Brand

- **1** ONE of the following:
- **1.1** ALL of the following:
- **1.1.1** Diagnosis of stable, symptomatic heart failure in sinus rhythm

AND

- **1.1.2** Both of the following:
 - Left ventricular ejection fraction less than or equal to 35%
 - Resting heart rate greater than or equal to 70 beats per minute

AND

1.1.3 Prescribed by or in consultation with a cardiologist

OR

- **1.2** BOTH of the following:
- 1.2.1 Diagnosis of Inappropriate Sinus Tachycardia

AND

- **1.2.2** One of the following:
- 1.2.2.1 Member has symptoms despite use of maximally tolerated beta blocker therapy

1.2.2.2 Member has	OR contraindication to beta blocker use
Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) must meet initial criteria for coverage

Product Name: Corlanor		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans*		

Product Name	Generic Name	GPI	Brand/Generic
CORLANOR	IVABRADINE HCL TAB 5 MG (BASE EQUIV)	40700035100320	Brand
CORLANOR	IVABRADINE HCL TAB 7.5 MG (BASE EQUIV)	40700035100330	Brand
CORLANOR	IVABRADINE HCL ORAL SOLN 5 MG/5ML (BASE EQUIV)	40700035102020	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Notes	*Member new to the plan (as evidenced by coverage effective date of
	less than or equal to 90 days) must meet initial criteria for coverage

Date	Notes
9/8/2023	2024 implementation

Cortic	Corticotropin Gel				
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Guideline ID	GL-144536
Guideline Name	Corticotropin Gel
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Brand Acthar, Brand Corticotropin		
Approval Length 12 month(s)		
Therapy Stage	Therapy Stage Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
ACTHAR	CORTICOTROPIN INJ GEL 80 UNIT/ML	30300010004010	Brand
CORTROPHIN	CORTICOTROPIN INJ GEL 80 UNIT/ML	30300010004010	Brand

Approval Criteria

1 - One of the following:

- **1.1** All of the following:
- **1.1.1** Diagnosis of infantile spasm with electroencephalogram pattern consistent with hypsarrhythmia

AND

1.1.2 Prescribed by, or in consultation with a Neurologist

AND

1.1.3 Member is less than 2 years of age

OR

- **1.2** Both of the following:
- 1.2.1 FDA approved diagnosis with evidence-based supporting literature/guideline

AND

1.2.2 Trial and failure, contraindication, or intolerance to an adequate trial of preferred formulary medications appropriate for the condition

Product Name: Brand Acthar, Brand Corticotropin		
Approval Length 3 Month(s) with partial fill (max 15 days/prescription)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization – All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
ACTHAR	CORTICOTROPIN INJ GEL 80 UNIT/ML	30300010004010	Brand
CORTROPHIN	CORTICOTROPIN INJ GEL 80 UNIT/ML	30300010004010	Brand

- 1 One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of infantile spasm with electroencephalogram pattern consistent with hypsarrhythmia

AND

1.1.2 Prescribed by, or in consultation with a Neurologist

AND

1.1.3 Member is less than 2 years of age

OR

- **1.2** Both of the following:
- 1.2.1 FDA approved diagnosis with evidence-based supporting literature/guideline

AND

1.2.2 Trial and failure, contraindication, or intolerance to an adequate trial of preferred formulary medications appropriate for the condition

Product Name: Brand Acthar, Brand Corticotropin			
Approval Length	12 month(s)		
Therapy Stage	Reauthorization		
Guideline Type	Prior Authorization - All Plans		
Product Generic Name GPI Brand/6		Brand/Generic	

Product Name	Generic Name	GPI	Brand/Generic
ACTHAR	CORTICOTROPIN INJ GEL 80 UNIT/ML	30300010004010	Brand
CORTROPHIN	CORTICOTROPIN INJ GEL 80 UNIT/ML	30300010004010	Brand

Approval Criteria
1 - One of the following:
1.1 All of the following:
1.1.1 Diagnosis of infantile spasm with electroencephalogram pattern consistent with hypsarrhythmia
AND
1.1.2 Prescribed by, or in consultation with a Neurologist
AND
1.1.3 Member is less than 2 years of age
OR
1.2 Both of the following:
1.2.1 FDA approved diagnosis with evidence-based supporting literature/guideline
AND
1.2.2 Trial and failure, contraindication, or intolerance to an adequate trial of preferred formulary medications appropriate for the condition
AND
2 - Submission of medical records (e.g., chart notes) with documentation of evidence-based rationale for continued use and evidence of member response to therapy from the previous period.

Date	Notes
3/18/2024	Updated product name

Cosentyx (secukinumab)			
(2) The interest and the large in the first term and count of states and states to proceed and states of the interest and the			

Guideline ID	GL-144845
Guideline Name	Cosentyx (secukinumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/20/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Cosentyx					
Diagnosis		Plaque Psoriasis			
Approval Length		12/31/2039*			
Guideline Type		Prior Authorization - All plans except IL and MN Plans			
Product Generic Name		Name	GPI	Brand/Generic	
COSENTYX SENSOREADY PEN		UMAB SUBCUTANEOUS SOLN AUTO- R 150 MG/ML	9025057500D520	Brand	
COSENTYX SENSOREADY PEN		UMAB SUBCUTANEOUS AUTO-INJ 150 00 MG DOSE)	9025057500D530	Brand	

COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

- **1** Diagnosis of moderate to severe plaque psoriasis with ONE of the following:
 - Significant functional disability
 - BSA involvement greater than 3%
 - Debilitating palmar/plantar psoriasis or other vulnerable areas that are difficult to treat such as nails, hairy/scalp areas, genitals, or intertriginous areas

Α	N	D

2 - Member is greater than 6 years old

AND

3 - Prescribed by or in consultation with a dermatologist

AND

- 4 One of the following:
- **4.1** Trial and failure, contraindication or intolerance to BOTH of the following:
- **4.1.1** Topical treatment (e.g., topical corticosteroids, calcipotriene, retinoids, calcineurin inhibitors)

AND

4.1.2 ONE of the following:

- Certolizumab
- Etanercept
- Adalimumab (biosimilars or Humira)
- Risankizumab
- Ustekinumab
- Guselkumab

OR

4.2 Continuation of prior therapy with secukinumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.29 (8ml every 28 days) f or 30 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cosentyx		
Diagnosis Plaque Psoriasis		
Approval Length	12 month(s)	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

Approval Criteria

1 - Diagnosis of moderate to severe plaque psoriasis with ONE of the following:
Significant functional disability
BSA involvement greater than 3%
 Debilitating palmar/plantar psoriasis or other vulnerable areas that are difficult to treat such as nails, hairy/scalp areas, genitals, or intertriginous areas
AND
2 - Member is greater than 6 years old
AND
3 - Prescribed by or in consultation with a dermatologist
AND
4 - One of the following:
4.1 Trial and failure, contraindication or intolerance to BOTH of the following:
4.1.1 Topical treatment (e.g., topical corticosteroids, calcipotriene, retinoids)
AND
4.1.2 ONE of the following:
Certolizumab
Etanercept
Adalimumab (biosimilars or Humira)Risankizumab
Ustekinumab
Guselkumab
OR
4.2 Continuation of prior therapy with secukinumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.29 (8ml every 28 days) f
	Second PA: Approve at GPI 10 for 12 months

Product Name: Cosentyx		
Diagnosis	Psoriatic Arthritis (PsA)	
Approval Length	12/31/2039*	
Guideline Type	Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis

AND

2 - Prescribed by or in consultation with a Dermatologist or Rheumatologist

AND

3 - One of the following:

- **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - · Actively inflamed joints
 - Axial disease
 - · Active skin, nail, or scalp psoriasis involvement
 - Dactylitis
 - Enthesitis

AND

- **3.1.2** Trial and failure or intolerance to a minimum 3 month trial or contraindication to ONE of the following:
 - adalimumab
 - certolizumab
 - etanercept
 - upadacitinib
 - risankizumab
 - guselkumab
 - golimumab
 - tofacitinib/tofacitinib XR
 - ustekinumab

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days) f or 30 days Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cosentyx		
Diagnosis	Psoriatic Arthritis (PsA)	
Approval Length	12 months*	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis

AND

2 - Prescribed by or in consultation with a Dermatologist or Rheumatologist

AND

- 3 One of the following:
 - **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - Actively inflamed joints
 - Axial disease
 - Active skin, nail, or scalp psoriasis involvement
 - Dactylitis
 - Enthesitis

AND

- 3.1.2 Trial and failure or intolerance to a minimum 3 month trial or contraindication to ONE of the following:
 - adalimumab
 - certolizumab
 - etanercept
 - upadacitinib
 - risankizumab
 - guselkumab

 - golimumab tofacitinib/tofacitinib XR
 - ustekinumab

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days) f
	or 30 days
	Second PA: Approve at GPI 10 for 12 months

Product Name: Cosentyx	
Diagnosis Ankylosing spondylitis (AS)	
Approval Length	12/31/2039*
Guideline Type	Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand

COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of Ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Trial and failure of a 1-month trial of scheduled prescription doses of two different NSAIDs (e.g., naproxen, nabumetone, diclofenac, etc.)

AND

- **3.1.2** Trial and failure or intolerance to a minimum 3 month trial or contraindication to ONE of the following:
 - adalimumab
 - certolizumab
 - etanercept
 - upadacitinib
 - golimumab
 - tofacitinib/tofacitinib XR

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days) f
	or 30 days Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cosentyx	
Diagnosis	Ankylosing spondylitis (AS)
Approval Length	12 months*
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of Ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

3 - One of the following:

- **3.1** All of the following:
- **3.1.1** Trial and failure of a 1-month trial of scheduled prescription doses of two different NSAIDs (e.g., naproxen, nabumetone, diclofenac, etc.)

AND

- **3.1.2** Trial and failure or intolerance to a minimum 3 month trial or contraindication to ONE of the following:
 - adalimumab
 - certolizumab
 - etanercept
 - upadacitinib
 - golimumab
 - tofacitinib/tofacitinib XR

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days)fo
	r 30 days
	Second PA: Approve at GPI 10 for 12 months

Product Name: Cosentyx	
Diagnosis	Non-radiographic axial spondyloarthritis (nr-axSpA)
Approval Length	12/31/2039*
Guideline Type	Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand

COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of active Non-radiographic axial spondyloarthritis (nr-axSpA)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Trial and failure of a 1-month trial of scheduled prescription doses of two different NSAIDs (e.g., naproxen, nabumetone, diclofenac, etc.)

AND

- **3.1.2** Trial and failure or intolerance to a minimum 3 month trial or contraindication to ONE of the following:
 - certolizumab
 - upadacitinib

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days) f
	or 30 days Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cosentyx		
Diagnosis	Non-radiographic axial spondyloarthritis (nr-axSpA)	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of active Non-radiographic axial spondyloarthritis (nr-axSpA)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

3 - One of the following:

- **3.1** All of the following:
- **3.1.1** Trial and failure of a 1-month trial of scheduled prescription doses of two different NSAIDs (e.g., naproxen, nabumetone, diclofenac, etc.)

AND

- **3.1.2** Trial and failure or intolerance to a minimum 3 month trial or contraindication to ONE of the following:
 - certolizumab
 - upadacitinib

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days) f or 30 days
	Second PA: Approve at GPI 10 for 12 months

Product Name: Cosentyx	
Diagnosis Enthesitis-related arthritis (ERA)	
Approval Length 12/31/2039*	
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand

COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of Enthesitis-related arthritis (ERA)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

3 - Member is greater than 4 years old

AND

- 4 One of the following:
 - **4.1** All of the following:
- **4.1.1** Trial and failure, contraindication or intolerance to at least one NSAID (e.g., naproxen, nabumetone, diclofenac, etc.)

AND

4.1.2 Trial and failure, contraindication or intolerance to one DMARD (e.g., sulfasalazine, methotrexate)

OR

*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days) f
or 30 days Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cosentyx		
Diagnosis	Enthesitis-related arthritis (ERA)	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - Diagnosis of Enthesitis-related arthritis (ERA)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

3 - Member is greater than 4 years old

AND

- 4 One of the following:
- **4.1** All of the following:
- **4.1.1** Trial and failure, contraindication or intolerance to at least one NSAID (e.g., naproxen, nabumetone, diclofenac, etc.)

AND

4.1.2 Trial and failure, contraindication or intolerance to one DMARD (e.g., sulfasalazine, methotrexate)

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve at GPI 14 with a MDD of 0.15 (4ml every 28 days) f
	or 30 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Cosentyx		
Diagnosis Plaque psoriasis, AS, PSA, ERA		
Approval Length	12 month(s)	
Guideline Type	Quantity Exception - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand

COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - FDA labeled regimen (based on weight or lack of response to lower doses)

OR

2 - Trial and failure of a 3-month course of standard dosing and the prescriber provides published literature supporting the requested regimen for the person's diagnosis

OR

3 - Continuation of previous therapy with secukinumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

Product Name: Cosentyx		
Diagnosis Plaque psoriasis, AS, PSA, ERA		
Approval Length	12/31/2039	
Guideline Type	Quantity Exception – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	9025057500D520	Brand
COSENTYX SENSOREADY PEN	SECUKINUMAB SUBCUTANEOUS AUTO-INJ 150 MG/ML (300 MG DOSE)	9025057500D530	Brand
COSENTYX UNOREADY	SECUKINUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	9025057500D550	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	9025057500E510	Brand

COSENTYX	SECUKINUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	9025057500E520	Brand
COSENTYX	SECUKINUMAB SUBCUTANEOUS PREF SYR 150 MG/ML (300 MG DOSE)	9025057500E530	Brand

1 - FDA labeled regimen (based on weight or lack of response to lower doses)

OR

2 - Trial and failure of a 3-month course of standard dosing and the prescriber provides published literature supporting the requested regimen for the person's diagnosis

OR

3 - Continuation of previous therapy with seculinumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

2. Revision History

Date	Notes
3/25/2024	New Program

Cystic Fibrosis Transmembrane Receptor (CFTR) Modifiers		
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Prior Authorization Guideline

Guideline ID	GL-137861	
Guideline Name	Cystic Fibrosis Transmembrane Receptor (CFTR) Modifiers	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
P&T Approval Date:	
P&T Revision Date:	

1. Criteria

Product Name: Kalydeco, Orkambi, Symdeko, Trikafta		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 80-40-60 MG& IVACAF 59.5MG THPK GRAN	4530990340B120	Brand
TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 100-50-75 MG& IVACAF 75MG THPK GRAN	4530990340B140	Brand
TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 50-25-37.5 MG & IVACAFTOR 75 MG TBPK	4530990340B720	Brand

TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 100-50-75 MG &IVACAFTOR 150 MG TBPK	4530990340B740	Brand
KALYDECO	IVACAFTOR TAB 150 MG	45302030000320	Brand
KALYDECO	IVACAFTOR PACKET 13.4 MG	45302030003005	Brand
KALYDECO	IVACAFTOR PACKET 25 MG	45302030003010	Brand
KALYDECO	IVACAFTOR PACKET 50 MG	45302030003020	Brand
KALYDECO	IVACAFTOR PACKET 75 MG	45302030003030	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR TAB 100-125 MG	45309902300310	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR TAB 200-125 MG	45309902300320	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR GRANULES PACKET 75- 94 MG	45309902303005	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR GRANULES PACKET 100- 125 MG	45309902303010	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR GRANULES PACKET 150- 188 MG	45309902303020	Brand
SYMDEKO	TEZACAFTOR-IVACAFTOR 50-75 MG & IVACAFTOR 75 MG TAB TBPK	4530990280B710	Brand
SYMDEKO	TEZACAFTOR-IVACAFTOR 100-150 MG & IVACAFTOR 150 MG TAB TBPK	4530990280B720	Brand

- **1** Submission of medical records (e.g., chart notes) documenting ALL of the following:
- **1.1** Diagnosis of cystic fibrosis (CF)

AND

- **1.2** Patient has at least one of the following mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene:
 - Homozygous F508del CFTR mutation
 - Heterozygous F508del CFTR mutation
 - Heterozygous CFTR mutation that does not include F508del (Mutation is responsive to the drug as noted in the product labeling)

AND

1.3 Patient has chronic sinopulmonary, gastrointestinal or nutritional abnormalities related to cystic fibrosis (CF) requiring medical treatment
AND
2 - Prescribed by or in consultation with one of the following:
 Pulmonologist Specialist in the care of cystic fibrosis (CF)
AND
3 - ONE of the following:
3.1 For members with homozygous F508del CFTR mutation, one of the following:
3.1.1 For Trikafta requests ONLY: Member is 2 years of age or older
OR
3.1.2 For Orkambi requests ONLY, one of the following::
3.1.2.1 Member is between 1 and 2 years of age
OR
3.1.2.2 Both of the following:
 Member is 2 years of age or older Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta
OR
3.1.3 For Symdeko requests ONLY, all of the following:
Member is 6 years of age or older

- Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta
- Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Orkambi

OR

- **3.2** For members with heterozygous F508del CFTR mutation, one of the following:
- 3.2.1 For Trikafta requests ONLY: Member is 2 years of age or older

OR

3.2.2 For Kalydeco requests ONLY, member is 1 months of age or older

OR

- **3.2.3** For Symdeko requests ONLY, both of the following:
 - Member is 6 years of age or older
 - Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta

OR

- **3.3** For members with heterozygous CFTR mutation that does not include F508del (Mutation is responsive to the drug as noted in the product labeling), one of the following:
 - 3.3.1 For Kalydeco requests ONLY, member is 1 months of age or older

OR

3.3.2 For Trikafta requests ONLY, member is 2 years of age or older

OR

3.3.3 For Symdeko requests ONLY, both of the following:

- Member is 6 years of age or older Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta

Product Name: Kalydeco, Orkambi, Symdeko, Trikafta			
Approval Length 12 month(s)			
Therapy Stage	herapy Stage Reauthorization		
Guideline Type Prior Authorization			

Product Name	Generic Name	GPI	Brand/Generic
TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 80-40-60 MG& IVACAF 59.5MG THPK GRAN	4530990340B120	Brand
TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 100-50-75 MG& IVACAF 75MG THPK GRAN	4530990340B140	Brand
TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 50-25-37.5 MG & IVACAFTOR 75 MG TBPK	4530990340B720	Brand
TRIKAFTA	ELEXACAF-TEZACAF-IVACAF 100-50-75 MG &IVACAFTOR 150 MG TBPK	4530990340B740	Brand
KALYDECO	IVACAFTOR TAB 150 MG	45302030000320	Brand
KALYDECO	IVACAFTOR PACKET 13.4 MG	45302030003005	Brand
KALYDECO	IVACAFTOR PACKET 25 MG	45302030003010	Brand
KALYDECO	IVACAFTOR PACKET 50 MG	45302030003020	Brand
KALYDECO	IVACAFTOR PACKET 75 MG	45302030003030	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR TAB 100-125 MG	45309902300310	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR TAB 200-125 MG	45309902300320	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR GRANULES PACKET 75- 94 MG	45309902303005	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR GRANULES PACKET 100- 125 MG	45309902303010	Brand
ORKAMBI	LUMACAFTOR-IVACAFTOR GRANULES PACKET 150- 188 MG	45309902303020	Brand
SYMDEKO	TEZACAFTOR-IVACAFTOR 50-75 MG & IVACAFTOR 75 MG TAB TBPK	4530990280B710	Brand
SYMDEKO	TEZACAFTOR-IVACAFTOR 100-150 MG & IVACAFTOR 150 MG TAB TBPK	4530990280B720	Brand

- **1** Submission of medical records (e.g., chart notes) from the previous 12 months demonstrating positive clinical response to therapy by one of the following:
 - FEV1 stabilization or improvement from baseline
 - Reduction in the number of pulmonary exacerbations that require antibiotics in the past year
 - Improvement in BMI from baseline
 - Member-specific description of benefit

AND

- **2** Submission of medical records (e.g., chart notes) documenting patient has at least one of the following mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene:
 - Homozygous F508del CFTR mutation
 - Heterozygous F508del CFTR mutation
 - Heterozygous CFTR mutation that does not include F508del (Mutation is responsive to the drug as noted in the product labeling)

AND

- **3** ONE of the following:
- **3.1** For members with homozygous F508del CFTR mutation, one of the following:
- 3.1.1 For Trikafta requests ONLY: Member is 2 years of age or older

OR

- **3.1.2** For Orkambi requests ONLY, one of the following:
- 3.1.2.1 Member is between 1 and 2 years of age

OR

- **3.1.2.2** Both of the following:
- Member is 2 years of age or older

 Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta

OR

- **3.1.3** For Symdeko requests ONLY, all of the following:
 - Member is 6 years of age or older
 - Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta
 - Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Orkambi

OR

- **3.2** For members with heterozygous F508del CFTR mutation, one of the following:
- 3.2.1 For Trikafta requests ONLY: Member is 2 years of age or older

OR

3.2.2 For Kalydeco requests ONLY, member is 1 months of age or older

OR

- 3.2.3 For Symdeko requests ONLY, both of the following:
 - Member is 6 years of age or older
 - Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta

OR

- **3.3** For members with heterozygous CFTR mutation that does not include F508del (Mutation is responsive to the drug as noted in the product labeling), one of the following:
 - 3.3.1 For Kalydeco requests ONLY, member is 1 months of age or older

OR

3.3.2 For Trikafta requests ONLY, member is 2 years of age or older

OR

- **3.3.3** For Symdeko requests ONLY, both of the following:
 - Member is 6 years of age or older
 - Submission of medical records (e.g., chart notes) documenting trial and failure to a minimum 6-month trial, contraindication, or intolerance to Trikafta

2. Revision History

Date	Notes
12/15/2023	New Program

Diacomit (Stiripentol)				
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Prior Authorization Guideline

Guideline ID	GL-136422	
Guideline Name	Diacomit (Stiripentol)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Diacomit		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
DIACOMIT	STIRIPENTOL CAP 250 MG	72600070000120	Brand
DIACOMIT	STIRIPENTOL CAP 500 MG	72600070000130	Brand
DIACOMIT	STIRIPENTOL PACKET 250 MG	72600070003020	Brand
DIACOMIT	STIRIPENTOL PACKET 500 MG	72600070003030	Brand

1 - Diagnosis of Dravet Syndrome

AND

2 - Prescribed by, or in consultation with, a neurologist

AND

3 - Age greater than or equal to 2 years

AND

4 - Used in combination with clobazam and valproate

Product Name: Diacomit		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
DIACOMIT	STIRIPENTOL CAP 250 MG	72600070000120	Brand
DIACOMIT	STIRIPENTOL CAP 500 MG	72600070000130	Brand
DIACOMIT	STIRIPENTOL PACKET 250 MG	72600070003020	Brand
DIACOMIT	STIRIPENTOL PACKET 500 MG	72600070003030	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Diacomit

Approval Length	12/31/2039	
Guideline Type	Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
DIACOMIT	STIRIPENTOL CAP 250 MG	72600070000120	Brand
DIACOMIT	STIRIPENTOL CAP 500 MG	72600070000130	Brand
DIACOMIT	STIRIPENTOL PACKET 250 MG	72600070003020	Brand
DIACOMIT	STIRIPENTOL PACKET 500 MG	72600070003030	Brand

1 - Diagnosis of Dravet Syndrome

AND

2 - Prescribed by, or in consultation with, a neurologist

AND

3 - Age greater than or equal to 2 years

AND

4 - Used in combination with clobazam and valproate

2. Revision History

Date	Notes
12/8/2023	New program

Dificid (Fidaxomicin)		
The State Progress of Subspace The Board States Considerated, in States		

Prior Authorization Guideline

Guideline ID	GL-129944
Guideline Name	Dificid (Fidaxomicin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Draduet	Canaria Na		CDI	Drand/Canaria
Guideline 7	Гуре	Prior Authorization		
Approval L	ength	12 month(s) with a fill count = 1		
Product Na	Product Name: Dificid			

Product Name	Generic Name	GPI	Brand/Generic
DIFICID	FIDAXOMICIN TAB 200 MG	03530025000320	Brand
DIFICID	FIDAXOMICIN FOR SUSP 40 MG/ML	03530025001920	Brand

Approval Criteria

- 1 All of the following:
- **1.1** Outpatient initiation of treatment

AND
1.2 Relapse or recurrence after a greater than or equal to 10 days treatment course with vancomycin
AND
1.3 One of the following:
1.3.1 Submission of medical records (i.e., PCR positive, toxin assay, or colonoscopy) of recurrent C difficile infection
OR
1.3.2 Submission of medical records (e.g., chart notes) documenting low levels of neutralizing antibodies to C. difficile
OR
2 - Both of the following:
2.1 Continuation of hospital therapy
AND
2.2 Member has been receiving as an inpatient during hospitalization and needs to complete the course of therapy as an outpatient
OR
3 - (Illinois plans only) – the requested drug is being used for the long-term treatment of tickborne disease
OR

- **4** (Minnesota plans only) Both of the following:
 - Member has stage four metastatic cancer
 - Requested drug is being used to treat a cancer-related C. difficile infection

2. Revision History

Date	Notes
10/25/2023	2024 New Implementation

Dojolvi (Triheptanoin)			
(3) hashed may sear to displace the base to see a seed, consider and further to prove the contributions			

Prior Authorization Guideline

Guideline ID	GL-131134
Guideline Name	Dojolvi (Triheptanoin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Dojolvi	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - ALL Plans

Product Name	Generic Name	GPI	Brand/Generic
DOJOLVI	TRIHEPTANOIN ORAL LIQUID 100%	80200080000920	Brand

Approval Criteria

1 - Diagnosis of long-chain fatty acid oxidation disorder

AND

- 2 Disease confirmed by one of the following:
 - elevation of acylcarnitine
 - enzyme activity assay below lower limit of normal
 - genetic testing showing mutation associated with long-chain fatty acid oxidation disorders

AND

3 - Trial and failure or contraindication to over-the-counter medium-chain fatty acid products for management of long-chain fatty acid oxidation disorder, including submission of medical records (e.g., chart notes) documenting specific adherence intervention deployed by a health care provider

AND

4 - Prescribed by or in consultation with a metabolic disease provider who specializes in management of fatty acid oxidation disorders

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it >\ se

Product Name: Dojolvi		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - ALL Plans	

Product Name	Generic Name	GPI	Brand/Generic
DOJOLVI	TRIHEPTANOIN ORAL LIQUID 100%	80200080000920	Brand

Approval Criteria

1 - Diagnosis of long-chain fatty acid oxidation disorder

AND

- 2 Disease confirmed by one of the following:
 - elevation of acylcarnitine
 - enzyme activity assay below lower limit of normal
 - genetic testing showing mutation associated with long-chain fatty acid oxidation disorders

AND

3 - Trial and failure or contraindication to over-the-counter medium-chain fatty acid products for management of long-chain fatty acid oxidation disorder, including submission of medical records (e.g., chart notes) documenting specific adherence intervention deployed by a health care provider

AND

4 - Prescribed by or in consultation with a metabolic disease provider who specializes in management of fatty acid oxidation disorders

AND

5 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member has shown improvement with requested drug (e.g., improved cardiac symptoms/function, decreased hospitalizations or urgent care visits, decreased hypoglycemic episodes, etc.)

Notes	*Member new to the plan (as evidenced by coverage effective date of
	less than or equal to 90 days) who initiated therapy using a manufactu
	rer-sponsored free drug program, provider samples, and/or vouchers
	will go through initial criteria, otherwise for continuation of therapy for
	new to plan, reauthorization criteria applies

2. Revision History

Date	Notes
8/20/2023	2024 New Implementation

Dry Eye Disease					
	The birth the process in trigger. The first term moved, wound, is about the first being prime to a country or trades.				

Prior Authorization Guideline

Guideline ID	GL-127812
Guideline Name	Dry Eye Disease
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Cequa, Tyrvaya, Xiidra	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CEQUA	CYCLOSPORINE (OPHTH) SOLN 0.09% (PF)	86720020002040	Brand
XIIDRA	LIFITEGRAST OPHTH SOLN 5%	86734050002020	Brand
TYRVAYA	VARENICLINE TARTRATE NASAL SOLN 0.03 MG/ACT	86280080202020	Brand

Approval Criteria

1 - Trial and failure of cyclosporine 0.05% eye drops

Product Name: Cequa, Tyrvaya, Xiidra	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CEQUA	CYCLOSPORINE (OPHTH) SOLN 0.09% (PF)	86720020002040	Brand
XIIDRA	LIFITEGRAST OPHTH SOLN 5%	86734050002020	Brand
TYRVAYA	VARENICLINE TARTRATE NASAL SOLN 0.03 MG/ACT	86280080202020	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Cequa, Tyrvaya, Xiidra	
Approval Length 12/31/2039	
Guideline Type Step Therapy - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CEQUA	CYCLOSPORINE (OPHTH) SOLN 0.09% (PF)	86720020002040	Brand
XIIDRA	LIFITEGRAST OPHTH SOLN 5%	86734050002020	Brand
TYRVAYA	VARENICLINE TARTRATE NASAL SOLN 0.03 MG/ACT	86280080202020	Brand

Approval Criteria

1 - Trial and failure of cyclosporine 0.05% eye drops

2. Revision History

Date	Notes
8/21/2023	New Program

Dupixent (dupilumab)					
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Prior Authorization Guideline

Guideline ID	GL-145322
Guideline Name	Dupixent (dupilumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	4/1/2017
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Dupixent				
Diagnosis Atopic Dermatitis				
Approval L	pproval Length 12 Month(s)*			
Guideline ⁻	Гуре	Prior Authorization - IL and MN Plans		
Product Name	Generic Na	eneric Name GPI Brand/Gen		Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML		9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML		9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED 9027302000E SYRINGE 100 MG/0.67ML		9027302000E510	Brand

DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

1 - Diagnosis of moderate to severe atopic dermatitis based on body surface area (greater than 10%), extent of disability, extent of pruritus, or impact on sleep, quality of life, current use of systemic immunomodulators)

AND

2 - Prescribed by, or in consultation with, a specialist experienced with the treatment of moderate to severe atopic dermatitis (e.g., Dermatologist, Pediatric Dermatologist, Allergist, Immunologist)

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance with at least TWO of the following:
 - Topical corticosteroid (e.g., clobetasol, betamethasone, triamcinolone)
 - Topical calcineurin inhibitor (e.g., pimecrolimus, tacrolimus)
 - Topical phosphodiesterase 4 (PDE-4) inhibitor (e.g., Crisaborole)
 - Topical janus kinase (JAK) inhibitor (e.g., ruxolitinib)
 - Phototherapy

OR

3.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For New Starts to Therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.22 (6 per 28 days) for 30
	Second PA: Approve at GPI 10 for 12 months

Product Name: Dupixent	
Diagnosis	Atopic Dermatitis
Approval Length	12/39/2039
Guideline Type	Prior Authorization - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

1 - Diagnosis of moderate to severe atopic dermatitis based on body surface area (greater than 10%), extent of disability, extent of pruritus, or impact on sleep, quality of life, current use of systemic immunomodulators)

AND

2 - Prescribed by, or in consultation with, a specialist experienced with the treatment of moderate to severe atopic dermatitis (e.g., Dermatologist, Pediatric Dermatologist, Allergist, Immunologist)

AND

- **3** One of the following:
- **3.1** Trial and failure, contraindication, or intolerance with at least TWO of the following:
 - Topical corticosteroid (e.g., clobetasol, betamethasone, triamcinolone)
 - Topical calcineurin inhibitor (e.g., pimecrolimus, tacrolimus)
 - Topical phosphodiesterase 4 (PDE-4) inhibitor (e.g., Crisaborole)
 - Topical janus kinase (JAK) inhibitor (e.g., ruxolitinib)

Phototherapy

OR

3.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve at GPI 14 with a MDD of 0.22 (6 per 28 days) for 30
	days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Dupixent	
Diagnosis	Severe Asthma
Approval Length	12 Month(s)*
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

Approval Criteria

1 - Diagnosis of eosinophilic asthma

AND

2 - Prescribed by, or in consultation with, an asthma specialist (e.g., Allergist, Immunologist, Pulmonologist)

- **3** One of the following:
- **3.1** Both of the following
- **3.1.1** Submission of medical records (e.g., chart notes) of one of the following:
 - Blood eosinophil count of greater than or equal to 150 cells/mm3 (other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease must be ruled out)
 - Oral corticosteroid dependent asthma

AND

- **3.1.2** One of the following:
- **3.1.2.1** Symptoms are not well controlled or poorly controlled despite an adherent (adherent treatment is defined as a medication possession ratio (MPR) greater than or equal to 70% based on the previous 120 days of prescription claims) greater than or equal to 3-month trial of medium to high- dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier

OR

3.1.2.2 Patient has intolerance to medium to high dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier (see background for exceptions)

OR

3.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with MDD of 0.22 (6 per 28 days) for 30 d
	ays
	Second PA: Approve at GPI 10 for 12 months

Product Name: Dupixent	
Diagnosis	Severe Asthma
Approval Length	12/31/2039*
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

1 - Diagnosis of eosinophilic asthma

AND

2 - Prescribed by, or in consultation with, an asthma specialist (e.g., Allergist, Immunologist, Pulmonologist)

AND

- 3 One of the following:
- 3.1 Both of the following
- **3.1.1** Submission of medical records (e.g., chart notes) of one of the following:
 - Blood eosinophil count of greater than or equal to 150 cells/mm3 (other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease must be ruled out)
 - Oral corticosteroid dependent asthma

- **3.1.2** One of the following:
- **3.1.2.1** Symptoms are not well controlled or poorly controlled despite an adherent (adherent treatment is defined as a medication possession ratio (MPR) greater than or equal to 70% based on the previous 120 days of prescription claims) greater than or equal to 3-month trial of medium to high- dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier

OR

3.1.2.2 Patient has intolerance to medium to high dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier (see background for exceptions)

OR

3.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with MDD of 0.22 (6 per 28 days) for 30 d
	ays Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Dupixent			
Diagnosis Nasal Polyps			
Approval Length	roval Length 12 month(s)		
Guideline Type Prior Authorization - IL or MN Plans Only			

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand

DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

- 1 Diagnosis of chronic rhinosinusitis with nasal polyposis with ALL of the following:
 - At least eight weeks of moderate to severe nasal congestion/blockage/obstruction or diminished sense of smell or rhinorrhea
 - Documented nasal polyps by direct exam, endoscopy, or sinus CT scan (ex: nasal polyp score five out of eight)
 - No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation

AND

2 - Prescribed by, or in consultation with, a specialist experienced in the treatment of nasal polyps (ex: Otolaryngologist, Allergist)

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Trial and failure, contraindication, or intolerance to one of the following:
 - Oral corticosteroids for nasal polyps
 - Prior surgery for nasal polyps greater than six months ago
 - IM corticosteroid injections for polyps with one previous steroid nasal spray
 - To greater than 2 nasal steroid sprays (i.e., failed two nasal sprays)

AND

3.1.2 Requested drug will be used in combination with a nasal corticosteroid medication

OR

3.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Dupixent	
Diagnosis Nasal Polyps	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

Approval Criteria

- 1 Diagnosis of chronic rhinosinusitis with nasal polyposis with ALL of the following:
 - At least eight weeks of moderate to severe nasal congestion/blockage/obstruction or diminished sense of smell or rhinorrhea
 - Documented nasal polyps by direct exam, endoscopy, or sinus CT scan (ex: nasal polyp score five out of eight)
 - No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation

AND

2 - Prescribed by, or in consultation with, a specialist experienced in the treatment of nasal polyps (ex: Otolaryngologist, Allergist)

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Trial and failure, contraindication, or intolerance to one of the following:
 - Oral corticosteroids for nasal polyps
 - Prior surgery for nasal polyps greater than six months ago
 - IM corticosteroid injections for polyps with one previous steroid nasal spray
 - To greater than 2 nasal steroid sprays (i.e., failed two nasal sprays)

3.1.2 Requested drug will be used in combination with a nasal corticosteroid medication

OR

3.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Dupixent		
Diagnosis Eosinophilic Esophagitis		
Approval Length	12 month(s)	
Guideline Type	Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

Approval Criteria

1 - Diagnosis of eosinophilic esophagitis confirmed by biopsy

AND

2 - Prescribed by, or in consultation with, a specialist experienced in the treatment of eosinophilic esophagitis (ex: Allergist, Gastroenterologist, Immunologist)

AND

3 - Member is 12 years of age or older

AND

- 4 One of the following:
- **4.1** Both of the following:
- **4.1.1** Trial and failure, intolerance, or contraindication to dietary therapy (e.g. elemental or amino acid-based formulary diet or empiric elimination diet that removes common triggers such as milk, wheat, soy, eggs, nuts, and seafood)

AND

4.1.2 Trial and failure, intolerance, or contraindication to proton pump inhibitors (e.g., omeprazole, pantoprazole, lansoprazole)

OR

4.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Dupixent		
Diagnosis Eosinophilic Esophagitis		
Approval Length	12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

1 - Diagnosis of eosinophilic esophagitis confirmed by biopsy

AND

2 - Prescribed by, or in consultation with, a specialist experienced in the treatment of eosinophilic esophagitis (ex: Allergist, Gastroenterologist, Immunologist)

AND

3 - Member is 12 years of age or older

AND

- 4 One of the following:
- **4.1** Both of the following:
- **4.1.1** Trial and failure, intolerance, or contraindication to dietary therapy (e.g. elemental or amino acid-based formulary diet or empiric elimination diet that removes common triggers such as milk, wheat, soy, eggs, nuts, and seafood)

AND

4.1.2 Trial and failure, intolerance, or contraindication to proton pump inhibitors (e.g., omeprazole, pantoprazole, lansoprazole)

OR

4.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Dupixent		
Diagnosis Prurigo nodularis (PN)		
Approval Length	12 Month(s)*	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

Approval Criteria

- 1 Diagnosis of chronic prurigo nodularis (PN) with all of the following:
 - At least 3 months of symptoms
 - At least 20 PN lesions in total
 - Severe or very severe itch (WI-NRS score ≥ 7)

AND

2 - Prescribed by or in consultation with a Dermatologist

3 - Member is 18 years of age or older

AND

- 4 One of the following:
 - **4.1** Trial and failure of an optimized regimen of one of the following:
 - Phototherapy
 - Moderate to high potency topical corticosteroids, intralesional corticosteroids, systemic corticosteroids
 - Systemic immunosuppressive agents (e.g., cyclosporine, methotrexate, azathioprine)
 - Immunomodulator agents (e.g., thalidomide, lenalidomide)
 - Anticonvulsants (e.g., pregabalin, gabapentin)

OR

4.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.22 (6 per 28 days) for 30 days Second PA: Approve at GPI 10 for 12 months

Product Name: Dupixent			
Diagnosis Prurigo nodularis (PN)			
Approval Length	12/31/2039		
Guideline Type	Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 200 MG/1.14ML	9027302000D215	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PEN-INJECTOR 300 MG/2ML	9027302000D220	Brand

DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	9027302000E510	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 200 MG/1.14ML	9027302000E515	Brand
DUPIXENT	DUPILUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	9027302000E520	Brand

1	- Diagnosis of	chronic	prurigo	nodularis ((PN) with	all of	the	following:

- At least 3 months of symptoms
- At least 20 PN lesions in total
- Severe or very severe itch (WI-NRS score ≥ 7)

AND

2 - Prescribed by or in consultation with a Dermatologist

AND

3 - Member is 18 years of age or older

AND

- 4 One of the following:
- **4.1** Trial and failure of an optimized regimen of one of the following:
 - Phototherapy
 - Moderate to high potency topical corticosteroids, intralesional corticosteroids, systemic corticosteroids
 - Systemic immunosuppressive agents (e.g., cyclosporine, methotrexate, azathioprine)
 - Immunomodulator agents (e.g., thalidomide, lenalidomide)
 - Anticonvulsants (e.g., pregabalin, gabapentin)

OR

4.2 Continuation of prior therapy with dupilumab, verified by paid claims or medical records (e.g. chart notes)					
Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.22 (6 per 28 days) for 30 days Second PA: Approve at GPI 10 through 12/31/2039				

2. Background

Benefit/Coverage/Program Information

Severe Asthma

Exceptions to high dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier. Exceptions based on adverse effects from high dose ICS or comorbid conditions increasing long-term risks of adverse effects from high dose ICS or oral corticosteroids include:

- Cataracts in patients > 40 years of age
- Glaucoma
- Recurrent Thrush
- Dysphonia
- Growth inhibition, after evaluation by Endocrine Consult
- Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

3. Revision History

Date	Notes
4/17/2024	Update COT language

Empaveli (Pegcetacoplan)				
The Market Segment to Angle of Tooks to the stand of the Angle of the Segment				

Prior Authorization Guideline

Guideline ID	GL-129123	
Guideline Name	Empaveli (Pegcetacoplan)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Empaveli		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization		

Product Name	Generic Name	GPI	Brand/Generic
EMPAVELI	PEGCETACOPLAN SUBCUTANEOUS SOLN 1080 MG/20ML (54 MG/ML)	85804065002020	Brand

Approval Criteria

1 - Confirmed diagnosis of PNH by flow cytometry

AND 2 - Prescribed by, or in consultation with, a Hematologist or Oncologist. **AND** 3 - Low hemoglobin (≤ 9 mg/dL with symptoms of anemia), elevated lactate dehydrogenase level (LDH ≥ 1.5 X ULN) and/or number of transfusions in last year **AND** 4 - Documentation of the clinical manifestations of disease (e.g., major vascular event, transfusion dependence, renal insufficiency, disabling fatigue and/or other end organ manifestations). AND 5 - Documentation of receipt of Advisory Committee on Immunization Practices (ACIP) recommended vaccinations at least two weeks prior to therapy initiation as outlined in drug REMS program. AND 6 - Age greater than or equal to 18 AND 7 - Drug is not being used in combination with another complement inhibitor* Notes *Combination of pegcetacoplan with another agent may be considere d for circumstances where all three individual complement inhibitors failed to adequately control anemia (eculizuma b or ravulizumab) or there are signs of ongoing hemolysis (pegcetacoplan).

Product Name: Empaveli		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization		

Product Name	Generic Name	GPI	Brand/Generic
EMPAVELI	PEGCETACOPLAN SUBCUTANEOUS SOLN 1080 MG/20ML (54 MG/ML)	85804065002020	Brand

1 - Confirmed diagnosis of PNH by flow cytometry

AND

2 - Prescribed by, or in consultation with, a Hematologist or Oncologist.

AND

3 - Low hemoglobin (\leq 9 mg/dL with symptoms of anemia), elevated lactate dehydrogenase level (LDH \geq 1.5 X ULN) and/or number of transfusions in last year.

AND

4 - Documentation of the clinical manifestations of disease (e.g., major vascular event, transfusion dependence, renal insufficiency, disabling fatigue and/or other end organ manifestations).

AND

5 - Documentation of receipt of Advisory Committee on Immunization Practices (ACIP) recommended vaccinations at least two weeks prior to therapy initiation as outlined in drug REMS program.

6 - Age greater than or equal to 18

AND

7 - Drug is not being used in combination with another complement inhibitor*

AND

8 - Clinical documentation from the past 12 months of improvement or clinical stability, (e.g., improvement in hemoglobin, lactate dehydrogenase level, haptoglobin level and/or number of transfusions in the last year).

Notes	*Combination of pegcetacoplan with another agent may be considere d for circumstances where all three individual
	complement inhibitors failed to adequately control anemia (eculizuma
	b or ravulizumab) or there are signs of ongoing hemolysis (pegcetacoplan).

Product Name: Empaveli		
Approval Length	Approval Length 2 doses/week	
Guideline Type Quantity Limit		

Product Name	Generic Name	GPI	Brand/Generic
EMPAVELI	PEGCETACOPLAN SUBCUTANEOUS SOLN 1080 MG/20ML (54 MG/ML)	85804065002020	Brand

Approval Criteria

1 - Documentation of continued hemolysis (LDH levels ≥ 2X ULM) despite an adequate 2-month trial of twice weekly dosing and the prescriber provided an evidence-based rationale for using the requested dose.

2. Revision History

Date	Notes
9/11/2023	New Program

١	Enbrel (etanercept)						
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Prior Authorization Guideline

Guideline ID	GL-145316	
Guideline Name	Enbrel (etanercept)	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Enbrel					
Diagnosis		Plaque Psoriasis			
Approval Length		12 Month(s)*			
Guideline T	уре	Prior Authorization – IL and MN Pla	ns		
Product Name	Generic Name		GPI	Brand/Generic	
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML		6629003000D530	Brand	
ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML		6629003000E230	Brand	
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 25 MG/0.5ML		6629003000E525	Brand	

ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML	6629003000E530	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML	66290030002015	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

- 2 One of the following:
 - **2.1** Both of the following:
 - **2.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g. nails, scalp, genitals, or intertriginous areas)

AND

2.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

OR

2.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

AND

3 - Prescribed by or in consultation with a dermatologist

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.29 (8 every 28 days) for

90 days Second PA: Approve at GPI 10 for 12 months
Second 1 A. Approve at Of 1 to lot 12 months

Product Name: Enbrel	
Diagnosis Plaque Psoriasis	
Approval Length	12/31/2039*
Guideline Type Prior Authorization – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML	6629003000D530	Brand
ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	6629003000E230	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 25 MG/0.5ML	6629003000E525	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML	6629003000E530	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML	66290030002015	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

- 2 One of the following:
- **2.1** Both of the following:
- **2.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g. nails, scalp, genitals, or intertriginous areas)

AND

2.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

OR

2.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

AND

3 - Prescribed by or in consultation with a dermatologist

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.29 (8 every 28 days) for
	90 days Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Enbrel		
Diagnosis Psoriatic Arthritis (PsA)		
Approval Length	12/31/2039	
Guideline Type	Prior Authorization – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML	6629003000D530	Brand
ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	6629003000E230	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 25 MG/0.5ML	6629003000E525	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML	6629003000E530	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML	66290030002015	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

3.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

Product Name: Enbrel	
Diagnosis Psoriatic Arthritis (PsA)	
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO- INJECTOR 50 MG/ML	6629003000D530	Brand
ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	6629003000E230	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 25 MG/0.5ML	6629003000E525	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML	6629003000E530	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML	66290030002015	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

3.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

Product Name: Enbrel		
Diagnosis	Juvenile Idiopathic Arthritis (JIA), Moderate to Severely Active Rheumatoid Arthritis (RA)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML	6629003000D530	Brand

ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	6629003000E230	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 25 MG/0.5ML	6629003000E525	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML	6629003000E530	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML	66290030002015	Brand

- 1 Diagnosis of one of the following:
 - Moderate to severely active rheumatoid arthritis (RA)
 - Juvenile idiopathic arthritis (JIA)

AND

- 2 One of the following:
- **2.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)*
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

OR

2.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

AND

3 - Prescribed by or in consultation with a rheumatologist

Notes	* Absolute contraindications to methotrexate are pregnancy, nursing,
	alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm
	bocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Enbrel	
Diagnosis	Juvenile Idiopathic Arthritis (JIA), Moderate to Severely Active Rheumatoid Arthritis (RA)
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML	6629003000D530	Brand
ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	6629003000E230	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 25 MG/0.5ML	6629003000E525	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML	6629003000E530	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML	66290030002015	Brand

- 1 Diagnosis of one of the following:
 - Moderate to severely active rheumatoid arthritis (RA)Juvenile idiopathic arthritis (JIA)

AND

- 2 One of the following:
- 2.1 Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)*
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

OR

2.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

AND

3 - Prescribed by or in consultation with a rheumatologist

* Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu
nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Enbrel	
Diagnosis Ankylosing Spondylitis (AS)	
Approval Length 12/31/2039	
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML	6629003000D530	Brand
ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	6629003000E230	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED 662900300 SYRINGE 25 MG/0.5ML		Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML	6629003000E530	Brand
ENBREL ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML		66290030002015	Brand

Approval Criteria

1 - Diagnosis of ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a rheumatologist

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

3.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

Product Name: Enbrel	
Diagnosis Ankylosing Spondylitis (AS)	
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ENBREL SURECLICK	ETANERCEPT SUBCUTANEOUS SOLUTION AUTO-INJECTOR 50 MG/ML	6629003000D530	Brand
ENBREL MINI	ETANERCEPT SUBCUTANEOUS SOLUTION CARTRIDGE 50 MG/ML	6629003000E230	Brand
ENBREL	ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 25 MG/0.5ML	6629003000E525	Brand
ENBREL	NBREL ETANERCEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/ML		Brand
ENBREL	ETANERCEPT SUBCUTANEOUS INJ 25 MG/0.5ML	66290030002015	Brand

Approval Criteria

1 - Diagnosis of ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

OR

3.2 Continuation of prior therapy with etanercept, verified by paid claims or medical records (e.g. chart notes)

2. Revision History

Date	Notes
4/15/2024	Updated COT language

Enspryng (Satralizumab)
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Prior Authorization Guideline

Guideline ID	GL-144867
Guideline Name	Enspryng (Satralizumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	11/18/2020
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: (Enspryng				
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization		
Product Name	Generic Name		GPI	Brand/Generic
		MAB-MWGE SUBCUTANEOUS SOLN IGE 120 MG/ML	9940507040E520	Brand
Approval Criteria				

1 - Diagnosis of neuromyelitis optica spectrum disorder (NMOSD) confirmed by positive serologic test for antiaquaporin-4 (AQP4) receptor antibody

AND

2 - Prescribed by, or in consultation with, a Neurologist or other specialist in NMOSD treatment

AND

- **3** One of the following:
- 3.1 All of the following:
- 3.1.1 History of at least one NMOSD relapse in the last 12 months

AND

3.1.2 Trial and failure, contraindication or intolerance to an adequate trial of at least one of the following: rituximab, mycophenolate or azathioprine

OR

3.2 Continuation of prior therapy with satralizumab, verified by paid claims or medical records (e.g. chart notes)

Date	Notes
3/26/2024	New Program

Enzy	me Inf	nibitors	tor Ga	aucher	Disease
The bind inagrament in	edigilgusi. The file may have been record, unwest, u	r sidded. Verily that the bid points to the account for an	d trades.		

Guideline ID	GL-129253	
Guideline Name	Enzyme Inhibitors for Gaucher Disease	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Miglustat	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
MIGLUSTAT	MIGLUSTAT CAP 100 MG	82700070000120	Generic

Approval Criteria

1 - Diagnosis of type-1 Gaucher disease

AND

2 - Trial and failure, intolerance, or contraindication to enzyme replacement therapy

Product Name: Generic Miglustat				
Approval Length 12 month(s)				
Therapy Stage		Reauthorization		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
MIGLUSTAT	MIGLUSTAT CAP 100 MG		82700070000120	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Cerdelga		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CERDELGA	ELIGLUSTAT TARTRATE CAP 84 MG (BASE EQUIVALENT)	82700040600120	Brand

Approval Criteria

1 - Diagnosis of type-1 Gaucher disease

AND

2 - Trial and failure, intolerance, or contraindication to enzyme replacement therapy

AND

3 - Have been determined to be CYP2D6 extensive, intermediate, or poor metabolizers by an FDA-approved test

Product Name: Cerdelga	
Approval Length 12 month(s)	
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
CERDELGA	ELIGLUSTAT TARTRATE CAP 84 MG (BASE EQUIVALENT)	82700040600120	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Generic Miglustat	
Approval Length 12/31/2039	
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
MIGLUSTAT	MIGLUSTAT CAP 100 MG	82700070000120	Generic

Approval Criteria

1 - Diagnosis of type-1 Gaucher disease

AND

2 - Trial and failure, intolerance, or contraindication to enzyme replacement therapy

Product Name: Cerdelga		
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
CERDELGA	ELIGLUSTAT TARTRATE CAP 84 MG (BASE EQUIVALENT)	82700040600120	Brand

Approval Criteria

1 - Diagnosis of type-1 Gaucher disease

AND

2 - Trial and failure, intolerance, or contraindication to enzyme replacement therapy

AND

3 - Have been determined to be CYP2D6 extensive, intermediate, or poor metabolizers by an FDA-approved test

Date	Notes
10/25/2023	New program

Erythropoiesis-Stimulating Agents		
Commission and contains the first transfer and the first and the first and the second according		

Guideline ID	GL-129740
Guideline Name	Erythropoiesis-Stimulating Agents
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Aranesp, Mircera, Retacrit				
Approval Length		12 month(s)		
Guideline Type		Prior Authorization		
Product Name			Brand/Generic	
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 10 MCG/0.4ML		8240101510E510	Brand
ARANESP ALBUMIN FREE			8240101510E528	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 40 MCG/0.4ML		8240101510E543	Brand

ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 60 MCG/0.3ML	8240101510E552	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 100 MCG/0.5ML	8240101510E560	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 150 MCG/0.3ML	8240101510E575	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 200 MCG/0.4ML	8240101510E582	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 300 MCG/0.6ML	8240101510E588	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN PREFILLED SYRINGE 500 MCG/ML	8240101510E590	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN INJ 25 MCG/ML	82401015102010	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN INJ 40 MCG/ML	82401015102020	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN INJ 60 MCG/ML	82401015102030	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN INJ 100 MCG/ML	82401015102040	Brand
ARANESP ALBUMIN FREE	DARBEPOETIN ALFA SOLN INJ 200 MCG/ML	82401015102060	Brand
RETACRIT	EPOETIN ALFA-EPBX INJ 2000 UNIT/ML	82401020042010	Generic
RETACRIT	EPOETIN ALFA-EPBX INJ 3000 UNIT/ML	82401020042015	Generic
RETACRIT	EPOETIN ALFA-EPBX INJ 4000 UNIT/ML	82401020042020	Generic
RETACRIT	EPOETIN ALFA-EPBX INJ 10000 UNIT/ML	82401020042040	Generic
RETACRIT	EPOETIN ALFA-EPBX INJ 20000 UNIT/ML	82401020042050	Generic
RETACRIT	EPOETIN ALFA-EPBX INJ 40000 UNIT/ML	82401020042060	Brand
MIRCERA	METHOXY PEG-EPOETIN BETA SOLN PREFILLED SYR 30 MCG/0.3ML	8240104010E510	Brand
MIRCERA	METHOXY PEG-EPOETIN BETA SOLN PREFILLED SYR 50 MCG/0.3ML	8240104010E515	Brand
MIRCERA	METHOXY PEG-EPOETIN BETA SOLN PREFILLED SYR 75 MCG/0.3ML	8240104010E520	Brand

MIRCERA	METHOXY PEG-EPOETIN BETA SOLN PREFILLED SYR 100 MCG/0.3ML	8240104010E525	Brand
MIRCERA	METHOXY PEG-EPOETIN BETA SOLN PREFILLED SYR 120 MCG/0.3ML	8240104010E530	Brand
MIRCERA	METHOXY PEG-EPOETIN BETA SOLN PREFILLED SYR 150 MCG/0.3ML	8240104010E535	Brand
MIRCERA	METHOXY PEG-EPOETIN BETA SOLN PREFILLED SYR 200 MCG/0.3ML	8240104010E545	Brand

Approval Criteria

- 1 Diagnosis of one of the following:
 - Non-myeloid cancer RECEIVING chemotherapy or within 8 weeks of receiving chemotherapy where the anemia is due to the effect of chemotherapy
 - HIV infection, for zidovudine-related anemia
 - Severe autoimmune hemolytic anemia
 - Myelodysplastic syndrome
 - Anemia associated with treatment regimens for Hepatitis C if ribavirin dose reduction does not provide adequate response
 - Chronic renal failure with or without dialysis
 - Post-transplant anemia
 - Religious beliefs prohibiting blood transfusions

AND

2 - Member or family member is self-administering the medication

AND

- **3** Submission of medical records (e.g., chart notes) documenting one of the following:
 - Hemoglobin (Hgb) < 10 g/dL
 - Hematocrit (HCT) < 30%

Date	Notes

8/21/2023	2024 New Implementation

Eucrisa (crisaborole)					
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Guideline ID	GL-127846
Guideline Name	Eucrisa (crisaborole)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Eucrisa		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
EUCRISA	CRISABOROLE OINT 2%	90230025004220	Brand

Approval Criteria

- 1 Trial and failure of one of the following:
 - 1.1 Topical steroid (see background for examples)

OR

1.2 Topical calcineurin inhibitor (e.g., tacrolimus, pimecrolimus)

Product Na	Product Name: Eucrisa				
Approval Length		12 month(s)			
Therapy Stage		Reauthorization			
Guideline Type		Step Therapy - IL and MN Plans			
Product Generic Na Name		nme	GPI	Brand/Generic	
EUCRISA CRISABOROI		LE OINT 2%	90230025004220	Brand	

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Eucrisa				
Approval Length		12/31/2039		
Guideline Type		Step Therapy - All plans except IL and MN Plans		
Product Name	Generic Na	me	GPI	Brand/Generic
EUCRISA CRISABOROL		LE OINT 2%	90230025004220	Brand

Approval Criteria

- 1 Trial and failure of one of the following:
 - **1.1** Topical steroid (see background for examples)

OR

1.2 Topical calcineurin inhibitor (e.g., tacrolimus, pimecrolimus)

2. Background

Benefit/Coverage/Program Information

Examples of topical steroids

alclometasone dipropionate 0.05% cream/ointment, betamethasone dipropionate 0.05% cream/ointment/lotion/gel/spray/pump, betamethasone valerate 0.1% ointment/cream/lotion, betamethasone valerate 0.12% foam, betamethasone/propylene glycol 0.05% cream/lotion/ointment, clobetasol propionate 0.025% cream, clobetasol propionate 0.05% ointment/cream/solution/gel/foam/lotion/spray, clobetasol propionate emollient 0.05% cream/foam, clocortolone pivalate 0.1% cream, desonide 0.05% ointment/lotion/cream/foam/gel, desoximetasone 0.05% gel/cream/ointment, desoximetasone 0.25% cream/ointment/spray, diflorasone diacetate 0.05% ointment/cream, diflorasone diacetate emollient 0.05% cream, fluocinolone acetonide 0.01% solution/cream/oil, fluocinolone acetonide 0.025% ointment/cream, fluocinonide 0.05% cream/ointment/solution/gel, fluocinonide 0.1% cream, fluocinonide emollient 0.05% cream, flurandrenolide 0.025% cream, flurandrenolide 0.05% cream/lotion/ointment, fluticasone propionate 0.005% ointment, fluticasone propionate 0.05% cream/lotion/, halcinonide 0.1% cream/ointment, halobetasol propionate 0.01% lotion, halobetasol propionate 0.05% cream/ointment/lotion/foam, hydrocortisone 1% cream/ointment, hydrocortisone 2% lotion, hydrocortisone 2.5% cream/ointment/solution/lotion, hydrocortisone butyrate 0.1% solution/cream/ointment/lotion, hydrocortisone butyrate emollient 0.1% cream, hydrocortisone probutate 0.1% cream, hydrocortisone valerate 0.2% ointment/cream, mometasone furoate 0.1% cream/ointment/solution, hydrocortisone acetate/aloe vera 2% lotion, prednicarbate 0.1% ointment/cream, triamcinolone acetonide 0.025% cream/ointment/lotion, triamcinolone acetonide 0.05% ointment, triamcinolone acetonide 0.1% cream/ointment/lotion, triamcinolone acetonide 0.147mg/g aerosol, triamcinolone acetonide 0.5% cream/ointment

Date	Notes
8/25/2023	New Programs

Evrys	Evrysdi (risdiplam)					
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Guideline ID	GL-131441
Guideline Name	Evrysdi (risdiplam)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Na	Product Name: Evrysdi				
Approval Length		12 month(s)			
Therapy Stage		Initial Authorization			
Guideline Type		Prior Authorization			
Product Generic Na Name		me	GPI	Brand/Generic	
EVRYSDI RISDIPLAM F		OR SOLN 0.75 MG/ML	74706560002120	Brand	

Approval Criteria

1 - Diagnosis of spinal muscle atrophy (SMA)

AND

2 - Submission of medical records (e.g., chart notes) documenting gene mutation analysis with bi-allelic SMN1 mutations (5q-autosomal recessive point mutation/deletion) phenotype 1,2 or 3.

AND

3 - Prescribed by or in consultation with a Neurologist or other clinician with expertise in management and treatment of neuromuscular disorders

AND

4 - Member does not have advanced SMA (e.g., permanent ventilatory dependence, tracheostomy, complete limb paralysis, etc.)

AND

5 - For members less than or equal to 2 yeas of age established on nusinersen, will not continue nusinersen (Spinraza) post onasemnogene infusion

AND

6 - For members less than or equal to 2 yeas of age established on risdiplam, will not continue risdiplam (Evrysdi) post- onasemnogene infusion

Product Name: Evrysdi		
Approval Length	oproval Length 12 month(s)	
Therapy Stage	Reauthorization	

Guideline 7	Гуре	Prior Authorization		
Product Name	Generic Na	me	GPI	Brand/Generic
EVRYSDI	RISDIPLAM F	OR SOLN 0.75 MG/ML	74706560002120	Brand

Approval Criteria

1 - Diagnosis of spinal muscle atrophy (SMA)

AND

2 - Submission of medical records (e.g., chart notes) documenting gene mutation analysis with bi-allelic SMN1 mutations (5q-autosomal recessive point mutation/deletion) phenotype 1,2 or 3.

AND

3 - Prescribed by or in consultation with a Neurologist or other clinician with expertise in management and treatment of neuromuscular disorders

AND

4 - Member does not have advanced SMA (e.g., permanent ventilatory dependence, tracheostomy, complete limb paralysis, etc.)

AND

5 - For members less than or equal to 2 yeas of age established on nusinersen, will not continue nusinersen (Spinraza) post onasemnogene infusion

AND

6 - For members less than or equal to 2 yeas of age established on risdiplam, will not continue risdiplam (Evrysdi) post- onasemnogene infusion

AND

7 - Member is established on therapy

AND

- **8** Submission of medical records (e.g., chart notes) documenting both of the following:
- **8.1** Clinically significant improvement in SMA-related symptoms as evidence by an improvement, stabilization or decreased decline since previous approval

AND

8.2 Specific scale used based on age and motor function and comparison to baseline

Date	Notes
10/8/2023	2024 New Implementation

Fasenra (benralizumab)
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Guideline ID	GL-145326
Guideline Name	Fasenra (benralizumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	4/19/2023
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Fasenra				
Approval L	Approval Length 12 month(s)*			
Guideline ⁻	Guideline Type Prior Authorization - IL and MN Plans			
Product Name			GPI	Brand/Generic
FASENRA PEN	BENRALIZUM INJECTOR 30	MAB SUBCUTANEOUS SOLN AUTO-) MG/ML	4460402000D520	Brand

Approval Criteria

1 - Prescribed by or in consultation with one of the following:

- Allergist
- Immunologist
- Pulmonologist

AND

2 - Member is 12 years of age or older

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** All of the following:
 - Diagnosis of eosinophilic asthma
 - Blood eosinophil count of ≥ 150 cells/mm3
 - All other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease have been ruled out

AND

- **3.1.2** One of the following:
- **3.1.2.1** Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent ‡ ≥ 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

- **3.1.2.2** Intolerance to medium to high dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier. Exceptions based on adverse effects from medium to high dose ICS or comorbid conditions increasing long-term risks of adverse effects from high dose ICS or oral corticosteroids including at least ONE of the following:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush

- Dysphonia
- Growth inhibition, after evaluation by Endocrine Consult
- Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

OR

3.2 Continuation of prior therapy with benralizumab, verified by paid claims or medical records (e.g. chart notes)

	,
Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for 90 days
	Second PA: Approve at GPI 12 for 12 months
	‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.
	NOTE: II-5 inhibitor drugs in combination with omalizumab will be considered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms. Tezepelumab, in combination with other
	biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Na	ame: Fasenr	a		
Approval Length 12/31/2039				
Guideline Type Prior Authorization – All Plans Except IL and MN Plans			s	
Product Name	Generic Na	me	GPI	Brand/Generic
FASENRA	BENRALIZUN	MAB SUBCUTANEOUS SOLN AUTO-	4460402000D520	Brand

Approval Criteria

PEN

1 - Prescribed by or in consultation with one of the following:

INJECTOR 30 MG/ML

- Allergist
- Immunologist
- Pulmonologist

AND

2 - Member is 12 years of age or older

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** All of the following:
 - Diagnosis of eosinophilic asthma
 - Blood eosinophil count of ≥ 150 cells/mm3
 - All other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease have been ruled out

AND

- **3.1.2** One of the following:
- **3.1.2.1** Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent ‡ ≥ 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

- **3.1.2.2** Intolerance to medium to high dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier. Exceptions based on adverse effects from medium to high dose ICS or comorbid conditions increasing long-term risks of adverse effects from high dose ICS or oral corticosteroids including at least ONE of the following:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush

- Dysphonia
- Growth inhibition, after evaluation by Endocrine Consult
- Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

OR

3.2 Continuation of prior therapy with benralizumab, verified by paid claims or medical records (e.g. chart notes)

` 3	,
Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for 90 days
	Second PA: Approve at GPI 12 to 12/31/2039
	‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.
	NOTE: II-5 inhibitor drugs in combination with omalizumab will be con sidered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms. Tezepelumab, in combination with other
	biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

2. Background

Benefit/Coverage/Program Information				
Table 1. Outcome Measure values for uncontrolled asthma				
Measure	Not Well Controlled	Very Poorly Controlled		
Baseline symptoms (outside of	> 2 days/week	Throughout the day		

exacerbation)			
Nighttime awakening	1-3 times/week	≥ 4 times/week	
Interference with normal activity	Some limitation	Extremely limited	
Short acting beta agonist use for symptom control	> 2 days/week	Several times per day	
FEV1	60-80% predicted or personal best	< 60% predicted or personal best	
Asthma exacerbations requiring oral	Yes	Yes	
steroids ≥ 2 times in the past year			
Asthma Control Test (ACT)	16-19	≤ 15	

Date	Notes
4/9/2024	Guideline Update.

Febuxostat
The bit shall reagn convex its elliphysel. Therefore may have been record, or didness, shall be to print to the best prints of the conversion and insulan.

Guideline ID	GL-128129	
Guideline Name	ebuxostat	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Febuxostat		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
FEBUXOSTAT	FEBUXOSTAT TAB 40 MG	68000030000320	Generic
FEBUXOSTAT	FEBUXOSTAT TAB 80 MG	68000030000330	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting a trial and failure of 3 months of allopurinol 300 mg

Product Name: Generic Febuxostat		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FEBUXOSTAT	FEBUXOSTAT TAB 40 MG	68000030000320	Generic
FEBUXOSTAT	FEBUXOSTAT TAB 80 MG	68000030000330	Generic

Approval Criteria

1 - Submission of medical notes (e.g. chart notes) from the past 12 months that member is continuing therapy with the requested drug

Product Name: Generic Febuxostat		
Approval Length	val Length 12/31/2039	
Guideline Type Step Therapy - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
FEBUXOSTAT	FEBUXOSTAT TAB 40 MG	68000030000320	Generic
FEBUXOSTAT	FEBUXOSTAT TAB 80 MG	68000030000330	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting a trial and failure of 3 months of allopurinol 300 mg

Date	Notes
8/21/2023	New Program

Fetzin	na (levomilnacipr	an)
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Guideline ID	GL-127842	
Guideline Name	etzima (levomilnacipran)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Fetzima	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FETZIMA TITRATION PACK	LEVOMILNACIPRAN HCL CAP ER 24HR 20 & 40 MG THERAPY PACK	5818005010B620	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 20 MG (BASE EQUIVALENT)	58180050107020	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 40 MG (BASE EQUIVALENT)	58180050107040	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 80 MG (BASE EQUIVALENT)	58180050107060	Brand

FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 120 MG (BASE EQUIVALENT)	58180050107080	Brand
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Approval Criteria

- 1 Trial and failure of at least one preferred serotonin reuptake inhibitor (SSRI):
 - citalopram
 - escitalopram
 - sertraline
 - paroxetine
 - fluoxetine

AND

- **2** Trial and failure of at least one preferred Serotonin Norepinephrine Reuptake inhibitor (SNRI):
 - venlafaxine
 - duloxetine

Product Name: Fetzima	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FETZIMA TITRATION PACK	LEVOMILNACIPRAN HCL CAP ER 24HR 20 & 40 MG THERAPY PACK	5818005010B620	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 20 MG (BASE EQUIVALENT)	58180050107020	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 40 MG (BASE EQUIVALENT)	58180050107040	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 80 MG (BASE EQUIVALENT)	58180050107060	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 120 MG (BASE EQUIVALENT)	58180050107080	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Fetzima		
Approval Length	Approval Length 12/31/2039	
Guideline Type Step Therapy - All plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
FETZIMA TITRATION PACK	LEVOMILNACIPRAN HCL CAP ER 24HR 20 & 40 MG THERAPY PACK	5818005010B620	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 20 MG (BASE EQUIVALENT)	58180050107020	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 40 MG (BASE EQUIVALENT)	58180050107040	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 80 MG (BASE EQUIVALENT)	58180050107060	Brand
FETZIMA	LEVOMILNACIPRAN HCL CAP ER 24HR 120 MG (BASE EQUIVALENT)	58180050107080	Brand

Approval Criteria

- 1 Trial and failure of at least one preferred serotonin reuptake inhibitor (SSRI):
 - citalopram
 - escitalopram
 - sertraline
 - paroxetine
 - fluoxetine

AND

- **2** Trial and failure of at least one preferred Serotonin Norepinephrine Reuptake inhibitor (SNRI):
 - venlafaxine

duloxetine

Date	Notes
8/21/2023	New Program

F	Fintepla (Fenfluramine)			
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Guideline ID	GL-129617	
Guideline Name	Fintepla (Fenfluramine)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Fintepla	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FINTEPLA	FENFLURAMINE HCL ORAL SOLN 2.2 MG/ML	72600028102020	Brand

Approval Criteria

1 - Diagnosis of seizures associated with Dravet syndrome or Lennox-Gastaut syndrome

AND

2 - Prescribed by, or in consultation with, a Neurologist

AND

3 - Previous trial and failure, contraindication or intolerance to an adequate trial of cannabidiol and at least one of the following: topiramate, valproic acid, clobazam or clobazam in combination with stiripentol.

Product Name: Fintepla			
Approval Length 12 month(s)			
Therapy St	Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans			
Product Generic Name GPI Brand/Ger		Brand/Generic	

Product Name	Generic Name	GPI	Brand/Generic
FINTEPLA	FENFLURAMINE HCL ORAL SOLN 2.2 MG/ML	72600028102020	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Fintepla	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
FINTEPLA	FENFLURAMINE HCL ORAL SOLN 2.2 MG/ML	72600028102020	Brand

Approval Criteria

1 - Diagnosis of seizures associated with Dravet syndrome or Lennox-Gastaut syndrome

AND

2 - Prescribed by, or in consultation with, a Neurologist

AND

3 - Previous trial and failure, contraindication or intolerance to an adequate trial of cannabidiol and at least one of the following: topiramate, valproic acid, clobazam or clobazam in combination with stiripentol.

Date	Notes
10/6/2023	New Program

Firdapse, Ruzurgi (amifampridine)	
S have been some taken had in the same and a sea of the same and the s	

Guideline ID	GL-127692
Guideline Name	Firdapse, Ruzurgi (amifampridine)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Firdapse	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
FIRDAPSE	AMIFAMPRIDINE PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	76000012100320	Brand

Approval Criteria

1 - All of the following:

1.1 Diagnosis of Lambert-Eaton myasthenic syndrome (LEMS)

AND

1.2 Diagnosis confirmed by neurophysiology studies or a positive anti-P/Q type voltagegated calcium channel antibody test

AND

1.3 Prescribed by or in consult with an expert in the treatment of neuromuscular disorders

OR

2 - Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days), the prescriber must provide submission of medical records (e.g. chart notes) from the previous 12 months regarding the member's response to therapy with improvement or stabilization in muscle weakness compared to baseline

Product Name: Firdapse		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
FIRDAPSE	AMIFAMPRIDINE PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	76000012100320	Brand

Approval Criteria

1 - Submission of medical records (e.g. chart notes) from the previous 12 months of therapy indicating improvement or stabilization in muscle weakness compared to baseline.

Date	Notes
11/3/2023	New Program

Fycompa (perampanel)			
The birth regions in the last technic mode, count, a state she birth to provide an extraction.			

Guideline ID	GL-127845
Guideline Name	Fycompa (perampanel)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Fycompa		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FYCOMPA	PERAMPANEL TAB 2 MG	72550060000310	Brand
FYCOMPA	PERAMPANEL TAB 4 MG	72550060000320	Brand
FYCOMPA	PERAMPANEL TAB 6 MG	72550060000330	Brand
FYCOMPA	PERAMPANEL TAB 8 MG	72550060000340	Brand
FYCOMPA	PERAMPANEL TAB 10 MG	72550060000350	Brand
FYCOMPA	PERAMPANEL TAB 12 MG	72550060000360	Brand

FYCOMPA	PERAMPANEL SUSP 0.5 MG/ML	72550060001820	Brand
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- 1 Trial and failure, contraindication, or intolerance to at least TWO preferred anticonvulsants:
 - lamotrigine
 - levetiracetam
 - carbamazepine
 - valproate
 - oxcarbazepine
 - gabapentin
 - pregabalin
 - topiramate
 - phenytoin
 - zonisamide
 - primidone

Product Name: Fycompa		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FYCOMPA	PERAMPANEL TAB 2 MG	72550060000310	Brand
FYCOMPA	PERAMPANEL TAB 4 MG	72550060000320	Brand
FYCOMPA	PERAMPANEL TAB 6 MG	72550060000330	Brand
FYCOMPA	PERAMPANEL TAB 8 MG	72550060000340	Brand
FYCOMPA	PERAMPANEL TAB 10 MG	72550060000350	Brand
FYCOMPA	PERAMPANEL TAB 12 MG	72550060000360	Brand
FYCOMPA	PERAMPANEL SUSP 0.5 MG/ML	72550060001820	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Fycompa		
Approval Length 12/31/2039		
Guideline Type Step Therapy - All plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
FYCOMPA	PERAMPANEL TAB 2 MG	72550060000310	Brand
FYCOMPA	PERAMPANEL TAB 4 MG	72550060000320	Brand
FYCOMPA	PERAMPANEL TAB 6 MG	72550060000330	Brand
FYCOMPA	PERAMPANEL TAB 8 MG	72550060000340	Brand
FYCOMPA	PERAMPANEL TAB 10 MG	72550060000350	Brand
FYCOMPA	PERAMPANEL TAB 12 MG	72550060000360	Brand
FYCOMPA	PERAMPANEL SUSP 0.5 MG/ML	72550060001820	Brand

- **1** Trial and failure, contraindication, or intolerance to at least TWO preferred anticonvulsants:
 - lamotrigine
 - levetiracetam
 - carbamazepine
 - valproate
 - oxcarbazepine
 - gabapentin
 - pregabalin
 - topiramate

 - phenytoin zonisamide
 - primidone

Date	Notes
8/21/2023	New Program

Galafold (Migalastat)					
E Debisings	senet kedişiyel. Terlir vey han ker renol, v	namani, or didded. Worly that the list pulses in the no	neific ed india.		

Guideline ID	GL-129103
Guideline Name	Galafold (Migalastat)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Galafold		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
GALAFOLD	MIGALASTAT HCL CAP 123 MG (BASE EQUIVALENT)	30903650100120	Brand

Approval Criteria

1 - Diagnosis of Fabry disease

AND

2 - Submission of medical records (e.g., chart notes) of an amenable galactosidase alpha gene (GLA) variant as noted in the product labeling.

AND

3 - Member does not have severe renal impairment (eGFR

AND

4 - Member is 16 years of age or older

AND

5 - Member will not be using migalastat in combination with enzyme replacement therapy

Product Name: Galafold		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
GALAFOLD	MIGALASTAT HCL CAP 123 MG (BASE EQUIVALENT)	30903650100120	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Galafold	
Approval Length	12/31//2039

Guideline 1	Гуре	Prior Authorization - All plans excep	t IL and MN	
Product Name	Generic Na	me	GPI	Brand/Generic
GALAFOLD	MIGALASTA	THCL CAP 123 MG (BASE EQUIVALENT)	30903650100120	Brand

1 - Diagnosis of Fabry disease

AND

2 - Submission of medical records (e.g., chart notes) of an amenable galactosidase alpha gene (GLA) variant as noted in the product labeling.

AND

3 - Member does not have severe renal impairment (eGFR

AND

4 - Member is 16 years of age or older

AND

5 - Member will not be using migalastat in combination with enzyme replacement therapy

Date	Notes
9/7/2023	New Program

Gattex (Teduglutide)	
Commission of the state of the	

Guideline ID	GL-131937
Guideline Name	Gattex (Teduglutide)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Gattex		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
GATTEX	TEDUGLUTIDE (RDNA) FOR INJ KIT 5 MG	52533070006420	Brand

Approval Criteria

1 - Diagnosis of Short Bowel Syndrome

AND

2 - Prescribed by, or in consultation with, a Gastroenterologist

AND

3 - Person dependent on parenteral support

Product Name: Gattex		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization-All plans	

Product Name	Generic Name	GPI	Brand/Generic
GATTEX	TEDUGLUTIDE (RDNA) FOR INJ KIT 5 MG	52533070006420	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months a \geq 20% reduction in parenteral support requirement from baseline.

Product Name: Gattex		
Approval Length	6 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
GATTEX	TEDUGLUTIDE (RDNA) FOR INJ KIT 5 MG	52533070006420	Brand

Approval Criteria

1 - Diagnosis of Short Bowel Syndrome

AND

2 - Prescribed by, or in consultation with, a Gastroenterologist

AND

3 - Person dependent on parenteral support

Date	Notes
10/31/2023	New program

Glucagon-like Pept	tide 1 (GLP-1) Agonis
	44

Guideline ID	GL-145315
Guideline Name	Glucagon-like Peptide 1 (GLP-1) Agonist
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2012
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Byetta, Bydureon, Trulicity				
Approval Length 12/31/2039		12/31/2039		
Guideline T	уре	Prior Authorization - All plans excep	t IL and MN	
Product Name	Generic Name GPI Brand/Generic		Brand/Generic	
BYETTA	EXENATIDE SOLN PEN-INJECTOR 5 MCG/0.02ML 2717002000D220 Br		Brand	
BYETTA	EXENATIDE SOLN PEN-INJECTOR 10 MCG/0.04ML 2717002000D240		Brand	
BYDUREON BCISE	EXENATIDE EXTENDED RELEASE SUSP AUTO- INJECTOR 2 MG/0.85ML 2717002000D420 Brand		Brand	
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 0.75 MG/0.5ML 2717001500D220 Brand			

TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 1.5 MG/0.5ML	2717001500D230	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 3 MG/0.5ML	2717001500D240	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 4.5 MG/0.5ML	2717001500D250	Brand

1 - Diagnosis of diabetes mellitus

Product Name: Byetta, Bydureon, Trulicity		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
BYETTA	EXENATIDE SOLN PEN-INJECTOR 5 MCG/0.02ML	2717002000D220	Brand
BYETTA	EXENATIDE SOLN PEN-INJECTOR 10 MCG/0.04ML	2717002000D240	Brand
BYDUREON BCISE	EXENATIDE EXTENDED RELEASE SUSP AUTO- INJECTOR 2 MG/0.85ML	2717002000D420	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 0.75 MG/0.5ML	2717001500D220	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 1.5 MG/0.5ML	2717001500D230	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 3 MG/0.5ML	2717001500D240	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 4.5 MG/0.5ML	2717001500D250	Brand

Approval Criteria

1 - Diagnosis of diabetes mellitus

Product Na	Product Name: Byetta, Bydureon, Trulicity			
Approval Length		12 month(s)		
Therapy Stage		Reauthorization		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic

BYETTA	EXENATIDE SOLN PEN-INJECTOR 5 MCG/0.02ML	2717002000D220	Brand
BYETTA	EXENATIDE SOLN PEN-INJECTOR 10 MCG/0.04ML	2717002000D240	Brand
BYDUREON BCISE	EXENATIDE EXTENDED RELEASE SUSP AUTO- INJECTOR 2 MG/0.85ML	2717002000D420	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 0.75 MG/0.5ML	2717001500D220	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 1.5 MG/0.5ML	2717001500D230	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 3 MG/0.5ML	2717001500D240	Brand
TRULICITY	DULAGLUTIDE SOLN PEN-INJECTOR 4.5 MG/0.5ML	2717001500D250	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
4/15/2024	Update Reauthorization Section

GNF	RH Anta	agonist		
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Guideline ID	GL-136601
Guideline Name	GNRH Antagonist
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Myfembree				
Approval Length		2 year(s)		
Guideline Type Prior Authorization - All plans except IL and MN		t IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
MYFEMBREE	RELUGOLIX-ESTRADIOL-NORETHINDRONE ACETATE TAB 40-1-0.5 MG		24993503800320	Brand

Approval Criteria

- 1 One of the following:
- **1.1** All of the following:

1.1.1 Diagnosis of heavy menstrual bleeding (menstrual blood loss greater than 80 ml) due to uterine fibroids AND 1.1.2 Member is premenopausal AND 1.1.3 Trial and failure, intolerance, or contraindication to two of the following: Combined oral contraceptives (e.g., Aubra, Gianvi) Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena) Tranexamic acid OR 1.2 Both of the following: 1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND			
AND 1.1.3 Trial and failure, intolerance, or contraindication to two of the following: Combined oral contraceptives (e.g., Aubra, Gianvi) Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena) Tranexamic acid OR 1.2 Both of the following: 1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology			
1.1.3 Trial and failure, intolerance, or contraindication to two of the following: Combined oral contraceptives (e.g., Aubra, Gianvi) Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena) Tranexamic acid OR 1.2 Both of the following: 1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	AND		
1.1.3 Trial and failure, intolerance, or contraindication to two of the following: Combined oral contraceptives (e.g., Aubra, Gianvi) Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena) Tranexamic acid OR 1.2 Both of the following: 1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	1.1.2 Member is premenopausal		
Combined oral contraceptives (e.g., Aubra, Gianvi) Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena) Tranexamic acid OR 1.2 Both of the following: 1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	AND		
Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena) Tranexamic acid OR 1.2 Both of the following: 1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	1.1.3 Trial and failure, intolerance, or contraindication to two of the following:		
1.2 Both of the following: 1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena)		
1.2.1 Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia) AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	OR		
AND 1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	1.2 Both of the following:		
1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	· · · · · · · · · · · · · · · · · · ·		
different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs AND 2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	AND		
2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology	different continuous hormonal contraceptives in combination with prescription-strength		
Gynecology	AND		

Product Name: Myfembree	
Approval Length 12 month(s)	

Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
MYFEMBREE	RELUGOLIX-ESTRADIOL-NORETHINDRONE ACETATE TAB 40-1-0.5 MG	24993503800320	Brand

- **1** One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of heavy menstrual bleeding (menstrual blood loss greater than 80 ml) due to uterine fibroids

AND

1.1.2 Member is premenopausal

AND

- **1.1.3** Trial and failure, intolerance, or contraindication to two of the following:
 - Combined oral contraceptives (e.g., Aubra, Gianvi)
 - Levonorgestrel-releasing intrauterine device (IUD, e.g., Mirena)
 - Tranexamic acid

OR

- **1.2** Both of the following:
- **1.2.1** Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia)

AND

1.2.2 Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs

AND

2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology

Product Name: Myfembree		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
MYFEMBREE	RELUGOLIX-ESTRADIOL-NORETHINDRONE ACETATE TAB 40-1-0.5 MG	24993503800320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Orilissa		
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ORILISSA	ELAGOLIX SODIUM TAB 150 MG (BASE EQUIV)	30090030100320	Brand
ORILISSA	ELAGOLIX SODIUM TAB 200 MG (BASE EQUIV)	30090030100330	Brand

Approval Criteria

1 - Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia)

AND

2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology

AND

3 - Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs

Product Name: Orilissa	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ORILISSA	ELAGOLIX SODIUM TAB 150 MG (BASE EQUIV)	30090030100320	Brand
ORILISSA	ELAGOLIX SODIUM TAB 200 MG (BASE EQUIV)	30090030100330	Brand

Approval Criteria

1 - Diagnosis of moderate to severe pain associated with endometriosis (other than dyspareunia)

AND

2 - Prescribed by, or in consultation with, an expert in the treatment of Obstetrics and/or Gynecology

AND

3 - Trial and failure (at least 3 months), intolerance, or contraindication to at least two different continuous hormonal contraceptives in combination with prescription-strength nonsteroidal anti-inflammatory (NSAID) drugs

Product Name: Orilissa	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ORILISSA	ELAGOLIX SODIUM TAB 150 MG (BASE EQUIV)	30090030100320	Brand
ORILISSA	ELAGOLIX SODIUM TAB 200 MG (BASE EQUIV)	30090030100330	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
11/20/2023	2024 New Implementation

Hemangeol (propranolo solution 4.28 mg/mL)		
(3) Inharmonic and the first the second of t		

Guideline ID	GL-131417	
Guideline Name	Hemangeol (propranolo solution 4.28 mg/mL)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2023
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1. Criteria

Product Nam	Product Name: Hemangeol			
Approval Length 12 month(s)				
Guideline Type		Prior Authorization		
Product Name	Generic N	Name	GPI	Brand/Generic
HEMANGEOL	PROPRANOLOL HCL ORAL SOLN 4.28 MG/ML (3.75 MG/ML BASE EQUIV)		33100040102080	Brand

Approval Criteria

1 - Diagnosis of proliferating infantile hemangioma requiring systemic therapy.

AND

2 - Therapeutic failure or intolerance to the preferred propranolol solution options at an equivalent dose.

AND

3 - The prescriber provides an evidence-based clinical rationale as to why the Hemangeol product would be expected to produce superior therapeutic results

Date	Notes
10/24/2023	New Program

Н	Hemlibra (Emicizumab)			
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Guideline ID	GL-129926	
Guideline Name	Hemlibra (Emicizumab)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Hemlibra				
Approval Length		12/31/2039		
Guideline Type		Prior Authorization - All plans except IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 30 MG/ML		85105030202010	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 60 MG/0.4ML (150 MG/ML)		85105030202020	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 105 MG/0.7ML (150 MG/ML)		85105030202030	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 150 85105030202040 Brand MG/ML		Brand	

Approval Criteria
1 - Diagnosis of congenital hemophilia A
AND
2 - One of the following:
2.1 ALL of the following:
2.1.1 Hemophilia A with inhibitors to Factor VIII
AND
2.1.2 Member requires prophylaxis to prevent or reduce bleeding episodes
AND
2.1.3 One of the following:
 Not used in combination with Immune Tolerance Induction (ITI) therapy Member is currently on a bypassing agent (NovoSeven, FEIBA)
OR
2.2 BOTH of the following:
2.2.1 Hemophilia A without inhibitors
AND
2.2.2 One of the following:
Member requires prophylaxis to prevent or reduce bleeding episodes despite optimal dose/frequency of Factor VIII product

• Member is unable to administer prophylaxis based on individual patient factors (e.g., IV access, home administration, etc.)

AND

3 - Member is followed by a plan approved bleeding disorders program

Product Name: Hemlibra	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL PPO/POS Plans

Product Name	Generic Name	GPI	Brand/Generic
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 30 MG/ML	85105030202010	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 60 MG/0.4ML (150 MG/ML)	85105030202020	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 105 MG/0.7ML (150 MG/ML)	85105030202030	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 150 MG/ML	85105030202040	Brand

Approval Criteria

1 - Diagnosis of congenital hemophilia A

AND

- 2 One of the following:
 - **2.1** ALL of the following:
 - 2.1.1 Hemophilia A with inhibitors to Factor VIII

AND

2.1.2 Member requires prophylaxis to prevent or reduce bleeding episodes

AND

2.1.3 One of the following:

- Not used in combination with Immune Tolerance Induction (ITI) therapy
- Member is currently on a bypassing agent (NovoSeven, FEIBA)

OR

2.2 BOTH of the following:

2.2.1 Hemophilia A without inhibitors

AND

2.2.2 One of the following:

- Member requires prophylaxis to prevent or reduce bleeding episodes despite optimal dose/frequency of Factor VIII product
- Member is unable to administer prophylaxis based on individual patient factors (e.g., IV access, home administration, etc.)

AND

3 - Member is followed by a specialist in bleeding disorders or a bleeding disorders program

Product Name: Hemlibra				
Approval Length 12 month(s)				
Therapy St	apy Stage Initial Authorization			
Guideline Type		Prior Authorization - MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 30 MG/ML		85105030202010	Brand

HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 60 MG/0.4ML (150 MG/ML)	85105030202020	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 105 MG/0.7ML (150 MG/ML)	85105030202030	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 150 MG/ML	85105030202040	Brand

1 - Diagnosis of congenital hemophilia A

AND

- 2 One of the following:
- **2.1** ALL of the following:
- 2.1.1 Hemophilia A with inhibitors to Factor VIII

AND

2.1.2 Member requires prophylaxis to prevent or reduce bleeding episodes

AND

- **2.1.3** One of the following:
 - Not used in combination with Immune Tolerance Induction (ITI) therapy
 - Member is currently on a bypassing agent (NovoSeven, FÈIBÁ)

OR

- **2.2** BOTH of the following:
- 2.2.1 Hemophilia A without inhibitors

AND

2.2.2 One of the following:

- Member requires prophylaxis to prevent or reduce bleeding episodes despite optimal dose/frequency of Factor VIII product
- Member is unable to administer prophylaxis based on individual patient factors (e.g., IV access, home administration, etc.)

AND

3 - Member is followed by a plan approved bleeding disorders program

Product Name: Hemlibra	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 30 MG/ML	85105030202010	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 60 MG/0.4ML (150 MG/ML)	85105030202020	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 105 MG/0.7ML (150 MG/ML)	85105030202030	Brand
HEMLIBRA	EMICIZUMAB-KXWH SUBCUTANEOUS SOLN 150 MG/ML	85105030202040	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
8/14/2023	2024 New Implementation

Hepatitis C Direct Acting Antivirals		
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Guideline ID	GL-144537
Guideline Name	Hepatitis C Direct Acting Antivirals
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Brand Epclusa, Mavyret		
Diagnosis Post-Transplant		
Approval Length	pproval Length 12 months with a fill count = 2-3 fills based on drug regimen requested	
Guideline Type Prior Authorization – IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
MAVYRET	GLECAPREVIR-PIBRENTASVIR TAB 100-40 MG	12359902350320	Brand
MAVYRET	GLECAPREVIR-PIBRENTASVIR PELLET PACK 50-20 MG	12359902353020	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR TAB 200-50 MG	12359902650320	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 200-50 MG	12359902653030	Brand

EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 150-37.5 MG	12359902653020	Brand
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1 - Member is scheduled to undergo, or is status-post heart, lung, kidney or liver transplant

AND

- 2 Both of the following:
 - HCV antibody (+) donorNAT (+) donor

AND

3 - HCV-negative recipients

Notes	*Approval length: Mavyret = 12 weeks, Epclusa = 4 weeks (can exten d to 12 weeks if cannot begin on Day 0 or any interruption in treatmen t)
	** Members new to the plan who are established on therapy will have coverage under their drug benefit for the remainder of the current treat ment course. Coverage of the drug product will be extended to new m embers who have already started therapy. Duration of therapy will be determined based on the person's indication and accepted treatment course in labeled and/or guideline supported regimens. Restrictions to specific network pharmacies and participation in medication manage ment programs may apply.

Product Name: Brand Epclusa, Mavyret		
Diagnosis Post-Transplant		
Approval Length 2-3 fills based on drug regimen requested		
Guideline Type Prior Authorization – All Plans except IL and MN		
Product Generic Name GPI		Brand/Generic

Product Name	Generic Name	GPI	Brand/Generic
MAVYRET	GLECAPREVIR-PIBRENTASVIR TAB 100-40 MG	12359902350320	Brand
MAVYRET	GLECAPREVIR-PIBRENTASVIR PELLET PACK 50-20 MG	12359902353020	Brand

EPCLUSA	SOFOSBUVIR-VELPATASVIR TAB 200-50 MG	12359902650320	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 200-50 MG	12359902653030	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 150-37.5 MG	12359902653020	Brand

1 - Member is scheduled to undergo, or is status-post heart, lung, kidney or liver transplant

AND

- 2 Both of the following:
 - HCV antibody (+) donor NAT (+) donor

AND

3 - HCV-negative recipients

Notes	*Approval length: Mavyret = 12 weeks, Epclusa = 4 weeks (can exten d to 12 weeks if cannot begin on Day 0 or any interruption in treatmen t)
	** Members new to the plan who are established on therapy will have coverage under their drug benefit for the remainder of the current treat ment course. Coverage of the drug product will be extended to new m embers who have already started therapy. Duration of therapy will be determined based on the person's indication and accepted treatment course in labeled and/or guideline supported regimens. Restrictions to specific network pharmacies and participation in medication manage ment programs may apply.

Product Name: Non-Preferred Agents: Sovaldi, Viekira Pak, Zepatier				
Diagnosis Chronic Hepatitis C Virus (HCV)				
Approval Length		12 month(s)		
Guideline Type		Prior Authorization - IL and MN Plans Only		
Product Generic Name Name		me	GPI	Brand/Generic

ZEPATIER	ELBASVIR-GRAZOPREVIR TAB 50-100 MG	12359902300320	Brand
SOVALDI	SOFOSBUVIR TAB 200 MG	12353080000310	Brand
SOVALDI	SOFOSBUVIR TAB 400 MG	12353080000320	Brand
SOVALDI	SOFOSBUVIR PELLET PACK 150 MG	12353080003015	Brand
SOVALDI	SOFOSBUVIR PELLET PACK 200 MG	12353080003020	Brand
VIEKIRA PAK TAB	OMBITAS-PARITAPRE-RITON & DASAB TAB PAK 12.5-75-50 & 250 MG	1235990460B720	Brand

- **1** Both of the following:
 - Diagnosis of chronic hepatitis C infection
 - HCV infection > 6 months

AND

- 2 Submission of medical records (e.g., chart notes) documenting ALL of the following:
 - HCV genotype
 - Viral RNA levels measured within the past 3 months prior to initiating therapy
 - Ane
 - Past treatment regimens used or documented treatment naïve status
 - Extent of liver disease: non-cirrhotic, compensated cirrhosis (Child-Pugh A), decompensated cirrhosis (ChildPugh B, C)
 - Current renal function
 - NS5A RAS (if indicated to direct treatment)
 - History of liver transplant
 - History of kidney transplant
 - HIV status and therapy

AND

- 3 One of the following:
- **3.1** Contraindication or intolerance to ALL of the following preferred agents:
 - Mavyret (glecaprevir/pibrentasvir)
 - Ledipasvir/sofosbuvir (Harvoni brand)
 - Sofosbuvir/velpatasvir (Epclusa brand)

• Vosevi (sofosbuvir/velpatasvir/voxilaprevir)

OR

3.2 The requested non-preferred agent will be used in a patient population that cannot be treated with a preferred agent

Notes	*Approval length: As indicated in package labeling or hcvguidelines.or g (the shortest appropriate recommended duration will be approved) ** Members new to the plan who are established on therapy will have coverage under their drug benefit for the remainder of the current treat ment course. Coverage of the drug product will be extended to new m embers who have already started therapy. Duration of therapy will be determined based on the person's indication and accepted treatment course in labeled and/or guideline supported regimens. Restrictions to specific network pharmacies and participation in medication manage ment programs may apply.

Product Name: Non-Preferred Agents: Sovaldi, Viekira Pak, Zepatier	
Diagnosis	Chronic Hepatitis C Virus (HCV)
Approval Length	*Approval length: As indicated in package labeling or hcvguidelines.org (the shortest appropriate recommended duration will be approved)
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
ZEPATIER	ELBASVIR-GRAZOPREVIR TAB 50-100 MG	12359902300320	Brand
SOVALDI	SOFOSBUVIR TAB 200 MG	12353080000310	Brand
SOVALDI	SOFOSBUVIR TAB 400 MG	12353080000320	Brand
SOVALDI	SOFOSBUVIR PELLET PACK 150 MG	12353080003015	Brand
SOVALDI	SOFOSBUVIR PELLET PACK 200 MG	12353080003020	Brand
VIEKIRA PAK TAB	OMBITAS-PARITAPRE-RITON & DASAB TAB PAK 12.5-75-50 & 250 MG	1235990460B720	Brand

Approval Criteria

- **1** Both of the following:
 - Diagnosis of chronic hepatitis C infection

HCV infection > 6 months

AND

- 2 Submission of medical records (e.g., chart notes) documenting ALL of the following:
 - HCV genotype
 - Viral RNA levels measured within the past 3 months prior to initiating therapy
 - Age
 - Past treatment regimens used or documented treatment naïve status
 - Extent of liver disease: non-cirrhotic, compensated cirrhosis (Child-Pugh A), decompensated cirrhosis (ChildPugh B, C)
 - Current renal function
 - NS5A RAS (if indicated to direct treatment)
 - History of liver transplant
 - History of kidney transplant
 - HIV status and therapy

AND

- **3** One of the following:
- **3.1** Contraindication or intolerance to ALL of the following preferred agents:
 - Mavyret (glecaprevir/pibrentasvir)
 - Ledipasvir/sofosbuvir (Harvoni brand)
 - Sofosbuvir/velpatasvir (Epclusa brand)
 - Vosevi (sofosbuvir/velpatasvir/voxilaprevir)

OR

3.2 The requested non-preferred agent will be used in a patient population that cannot be treated with a preferred agent

*Approval length: As indicated in package labeling or hcvguidelines.or g (the shortest appropriate recommended duration will be approved) ** Members new to the plan who are established on therapy will have coverage under their drug benefit for the remainder of the current treat ment course. Coverage of the drug product will be extended to new m embers who have already started therapy. Duration of therapy will be determined based on the person's indication and accepted treatment course in labeled and/or guideline supported regimens. Restrictions to specific network pharmacies and participation in medication manage ment programs may apply.

Product Name: Preferred Agents: Brand Epclusa, Brand Harvoni, Brand Harvoni Pak, Mavyret, Vosevi		
Diagnosis	Chronic Hepatitis C Virus (HCV)	
Approval Length 12 month(s)		
Guideline Type	Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
MAVYRET	GLECAPREVIR-PIBRENTASVIR TAB 100-40 MG	12359902350320	Brand
MAVYRET	GLECAPREVIR-PIBRENTASVIR PELLET PACK 50-20 MG	12359902353020	Brand
VOSEVI	SOFOSBUVIR-VELPATASVIR-VOXILAPREVIR TAB 400-100-100 MG	12359903800330	Brand
HARVONI	LEDIPASVIR-SOFOSBUVIR TAB 90-400 MG	12359902400320	Generic
EPCLUSA	SOFOSBUVIR-VELPATASVIR TAB 200-50 MG	12359902650320	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 200-50 MG	12359902653030	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 150-37.5 MG	12359902653020	Brand
HARVONI	LEDIPASVIR-SOFOSBUVIR TAB 45-200 MG	12359902400310	Brand
HARVONI	LEDIPASVIR-SOFOSBUVIR PELLET PACK 33.75-150 MG	12359902403006	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR TAB 400-100 MG	12359902650330	Generic

- **1** Both of the following:
 - Diagnosis of chronic hepatitis C infection
 - HCV infection > 6 months

AND

- **2** Submission of medical records (e.g., chart notes) documenting ALL of the following:

 - HCV genotype Viral RNA levels measured within the past 3 months prior to initiating therapy
 - Age

- Past treatment regimens used or documented treatment naïve status
- Extent of liver disease: non-cirrhotic, compensated cirrhosis (Child-Pugh A), decompensated cirrhosis (ChildPugh B, C)
- Current renal function
- NS5A RAS (if indicated to direct treatment)
- History of liver transplant
- History of kidney transplant
- HIV status and therapy

Notes	*Approval length: As indicated in package labeling or hcvguidelines.or g (the shortest appropriate recommended duration will be approved) ** Members new to the plan who are established on therapy will have coverage under their drug benefit for the remainder of the current treat ment course. Coverage of the drug product will be extended to new m embers who have already started therapy. Duration of therapy will be determined based on the person's indication and accepted treatment course in labeled and/or guideline supported regimens. Restrictions to specific network pharmacies and participation in medication manage ment programs may apply.

Product Name: Preferred Agents: Brand Epclusa, Brand Harvoni, Brand Harvoni Pak, Mavyret, Vosevi	
Diagnosis	Chronic Hepatitis C Virus (HCV)
Approval Length	*Approval length: As indicated in package labeling or hcvguidelines.org (the shortest appropriate recommended duration will be approved)
Guideline Type	Prior Authorization-All Plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
MAVYRET	GLECAPREVIR-PIBRENTASVIR TAB 100-40 MG	12359902350320	Brand
MAVYRET	GLECAPREVIR-PIBRENTASVIR PELLET PACK 50-20 MG	12359902353020	Brand
VOSEVI	SOFOSBUVIR-VELPATASVIR-VOXILAPREVIR TAB 400-100-100 MG	12359903800330	Brand
HARVONI	LEDIPASVIR-SOFOSBUVIR TAB 90-400 MG	12359902400320	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR TAB 200-50 MG	12359902650320	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 200-50 MG	12359902653030	Brand
EPCLUSA	SOFOSBUVIR-VELPATASVIR PELLET PACK 150-37.5 MG	12359902653020	Brand
HARVONI	LEDIPASVIR-SOFOSBUVIR TAB 45-200 MG	12359902400310	Brand

HARVONI	LEDIPASVIR-SOFOSBUVIR PELLET PACK 33.75-150 MG	12359902403006	Brand
EPCLUSA	PCLUSA SOFOSBUVIR-VELPATASVIR TAB 400-100 MG 12359902650330 Generic		Generic

- **1** Both of the following:
 - Diagnosis of chronic hepatitis C infection
 - HCV infection > 6 months

AND

- 2 Submission of medical records (e.g., chart notes) documenting ALL of the following:
 - HCV genotype
 - Viral RNA levels measured within the past 3 months prior to initiating therapy
 - Age
 - Past treatment regimens used or documented treatment naïve status
 - Extent of liver disease: non-cirrhotic, compensated cirrhosis (Child-Pugh A), decompensated cirrhosis (ChildPugh B, C)
 - Current renal function
 - NS5A RAS (if indicated to direct treatment)
 - History of liver transplant
 - History of kidney transplant
 - HIV status and therapy

g (the shortest appropriate recommended duration will be approved)		
coverage under their drug benefit for the remainder of the current treatment course. Coverage of the drug product will be extended to new nembers who have already started therapy. Duration of therapy will be determined based on the person's indication and accepted treatment course in labeled and/or guideline supported regimens. Restrictions to	Notes	*** Members new to the plan who are established on therapy will have coverage under their drug benefit for the remainder of the current treat ment course. Coverage of the drug product will be extended to new m embers who have already started therapy. Duration of therapy will be determined based on the person's indication and accepted treatment course in labeled and/or guideline supported regimens. Restrictions to specific network pharmacies and participation in medication manage

Date	Notes

3/18/2024	Updated product name

Hereditary Angioedema (HAE) Medicati				
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Prior Authorization Guideline

Guideline ID	GL-145339	
Guideline Name	Hereditary Angioedema (HAE) Medications	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	1/17/2018
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Berinert, generic icatibant, Ruconest				
Diagnosis Treatment of Acute Attacks		Treatment of Acute Attacks		
Approval Length		12/31/2039		
Therapy Stage Initial Authorization				
Guideline Type		Prior Authorization - All Plans Except IL and MN Plans		
Product Name	Generic Name C1 ESTERASE INHIBITOR (HUMAN) FOR IV INJ KIT 500 UNIT		GPI	Brand/Generic
BERINERT			85802022006420	Brand
RUCONEST	C1 ESTERAS INJ 2100 UN	SE INHIBITOR (RECOMBINANT) FOR IV	85802022102130	Brand

ICATIBANT ACETATE SUBCUTANEOUS SOLN PREF SYR 30 MG/3ML	8582004010E520	Generic
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1 - Diagnosis of Hereditary Angioedema (HAE)

AND

- 2 One of the following:
- 2.1 Low C4 AND low C1 inhibitor level or function

OR

- **2.2** Both of the following:
 - Normal C1 inhibitor level with a family history of HAE
 - High dose antihistamines did not control symptoms

AND

3 - Prescribed by or in consultation with an allergist or other provider experienced in the treatment of HAE

AND

4 - All medications that may cause angioedema have been discontinued (e.g., ACE inhibitors, estrogens, ARBs)

Product Name: Cinryze	
Diagnosis	Long-Term Prevention/Prophylaxis
Approval Length 12/31/2039	
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
CINRYZE	C1 ESTERASE INHIBITOR (HUMAN) FOR IV INJ 500 UNIT	85802022002120	Brand

1 - Diagnosis of Hereditary Angioedema (HAE)

AND

2 - Prescribed by or in consultation with an allergist or other provider experienced in the treatment of HAE

AND

- 3 One of the following:
 - **3.1** All of the following:
 - **3.1.1** One of the following:
 - History of ≥ 2 attacks per month
 - Symptoms are moderate to severe

AND

- **3.1.2** One of the following:
- 3.1.2.1 Low C4 AND low C1 inhibitor level or function

OR

- **3.1.2.2** Both of the following:
 - Normal C1 inhibitor level with a family history of HAE
 - High dose antihistamines did not control symptoms

AND

3.1.3 All medications that may cause angioedema have been discontinued (e.g., ACE inhibitors, estrogens, ARBs)

AND

- **3.1.4** Trial and failure (defined as no reduction in frequency of attacks or severity of attacks), contraindication, or intolerance to BOTH of the following:
 - Haegarda
 - Takhzyro

AND

- **3.1.5** One of the following:
 - Trial and failure (defined as no reduction in frequency of attacks or severity of attacks), contraindication, or intolerance to Orladeyo
 - Member is between 6 and 12 years of age

OR

3.2 Continuation of prior therapy with Cinryze, verified by paid claims or medical records (e.g. chart notes)

Product Name: Berinert, generic icatibant, Ruconest	
Diagnosis	Treatment of Acute Attacks
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
BERINERT	C1 ESTERASE INHIBITOR (HUMAN) FOR IV INJ KIT 500 UNIT	85802022006420	Brand

	RUCONEST	C1 ESTERASE INHIBITOR (RECOMBINANT) FOR IV INJ 2100 UNIT	85802022102130	Brand
ı	ICATIBANT ACETATE	ICATIBANT ACETATE SUBCUTANEOUS SOLN PREF SYR 30 MG/3ML	8582004010E520	Generic

1 - Diagnosis of Hereditary Angioedema (HAE)

AND

- 2 One of the following:
- 2.1 Low C4 AND low C1 inhibitor level or function

OR

- **2.2** Both of the following:
 - Normal C1 inhibitor level with a family history of HAE
 - High dose antihistamines did not control symptoms

AND

3 - Prescribed by or in consultation with an allergist or other provider experienced in the treatment of HAE

AND

4 - All medications that may cause angioedema have been discontinued (e.g., ACE inhibitors, estrogens, ARBs)

AND

5 - Requested medication will not be used in combination with other approved treatments for acute attacks

Product Name: Cinryze	Product Name: Cinryze		
Diagnosis	Long-Term Prevention/Prophylaxis		
Approval Length	12 month(s)		
Guideline Type	Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
CINRYZE	C1 ESTERASE INHIBITOR (HUMAN) FOR IV INJ 500 UNIT	85802022002120	Brand

1 - Diagnosis of Hereditary Angioedema (HAE)

AND

2 - Prescribed by or in consultation with an allergist or other provider experienced in the treatment of HAE

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** One of the following:
 - History of ≥ 2 attacks per month
 - Symptoms are moderate to severe

AND

- **3.1.2** One of the following:
- **3.1.2.1** Low C4 AND low C1 inhibitor level or function

OR

3.1.2.2 Both of the following:

- Normal C1 inhibitor level with a family history of HAE
- High dose antihistamines did not control symptoms

AND

3.1.3 All medications that may cause angioedema have been discontinued (e.g., ACE inhibitors, estrogens, ARBs)

AND

- **3.1.4** Trial and failure (defined as no reduction in frequency of attacks or severity of attacks), contraindication, or intolerance to BOTH of the following:
 - Haegarda
 - Takhzyro

AND

- **3.1.5** One of the following:
 - Trial and failure (defined as no reduction in frequency of attacks or severity of attacks), contraindication, or intolerance to Orladevo
 - Member is between 6 and 12 years of age

OR

3.2 Continuation of prior therapy with Cinryze, verified by paid claims or medical records (e.g. chart notes)

Product Name: Berinert, generic icatibant, Ruconest		
Diagnosis Treatment of Acute Attacks		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
BERINERT	C1 ESTERASE INHIBITOR (HUMAN) FOR IV INJ KIT 500 UNIT	85802022006420	Brand
RUCONEST	C1 ESTERASE INHIBITOR (RECOMBINANT) FOR IV INJ 2100 UNIT	85802022102130	Brand
ICATIBANT ACETATE	ICATIBANT ACETATE SUBCUTANEOUS SOLN PREF SYR 30 MG/3ML	8582004010E520	Generic

1 - For members new to plan* (as evidenced by coverage effective date of less than or equal to 90 days)**: Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member has experienced positive clinical response to therapy with the requested drug

OR

2 - For members requesting renewal (reauthorization): Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member has experienced positive clinical response to therapy with the requested drug

Notes	*Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on therapy, will have coverage under their drug benefit for the remainder of the curren t treatment course. Restrictions to specific network pharmacies and p articipation in medication management programs may apply.
	**Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufact urer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

Product Name: Berinert, generic icatibant, Ruconest		
Diagnosis	iagnosis Treatment of Acute Attacks	
Approval Length	12/31/2039	
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
BERINERT	C1 ESTERASE INHIBITOR (HUMAN) FOR IV INJ KIT 500 UNIT	85802022006420	Brand
RUCONEST	C1 ESTERASE INHIBITOR (RECOMBINANT) FOR IV INJ 2100 UNIT	85802022102130	Brand
ICATIBANT ACETATE	ICATIBANT ACETATE SUBCUTANEOUS SOLN PREF SYR 30 MG/3ML	8582004010E520	Generic

1 - For members new to plan* (as evidenced by coverage effective date of less than or equal to 90 days)**: Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member has experienced positive clinical response to therapy with the requested drug

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Notes	*Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on therapy, will have coverage under their drug benefit for the remainder of the curren t treatment course. Restrictions to specific network pharmacies and p articipation in medication management programs may apply.
	**Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufact urer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

Product Name: Haegarda, Orladeyo, Takhzyro				
Diagnosis Long-Term Prevention/Prophylaxis				
Approval Length 6 month(s)				
Therapy Sta	age	Initial Authorization		
Guideline T	уре	Prior Authorization - All Plans Except IL and MN Plans		
Product Name	Generic Na	Generic Name		Brand/Generic
HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 2000 UNIT		85802022002130	Brand
HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 3000 UNIT		85802022002140	Brand
TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 150 MG/ML		8584204020E510	Brand

TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 300 MG/2ML (150 MG/ML)	8584204020E520	Brand
TAKHZYRO	LANADELUMAB-FLYO INJ 300 MG/2ML (150 MG/ML)	85842040202020	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 110 MG	85840010200120	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 150 MG	85840010200130	Brand

1 - Diagnosis of Hereditary Angioedema (HAE)

AND

- 2 One of the following:
 - History of ≥ 2 attacks per month
 - Symptoms are moderate to severe

AND

- 3 One of the following:
- 3.1 Low C4 AND low C1 inhibitor level or function

OR

- **3.2** Both of the following:
 - Normal C1 inhibitor level with a family history of HAE
 - High dose antihistamines did not control symptoms

AND

4 - Prescribed by or in consultation with an allergist or other provider experienced in the treatment of HAE

AND

5 - All medications that may cause angioedema have been discontinued (e.g., ACE inhibitors, estrogens, ARBs)

AND

6 - Requested medication will not be used in combination with other approved HAE prevention treatments

Product Name: Haegarda, Orladeyo, Takhzyro		
Diagnosis Long-Term Prevention/Prophylaxis		
Approval Length	6 month(s)	
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 2000 UNIT	85802022002130	Brand
HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 3000 UNIT	85802022002140	Brand
TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 150 MG/ML	8584204020E510	Brand
TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 300 MG/2ML (150 MG/ML)	8584204020E520	Brand
TAKHZYRO	LANADELUMAB-FLYO INJ 300 MG/2ML (150 MG/ML)	85842040202020	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 110 MG	85840010200120	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 150 MG	85840010200130	Brand

Approval Criteria

1 - For members new to plan* (as evidenced by coverage effective date of less than or equal to 90 days)**: Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member has experienced positive clinical response to therapy with the requested drug

OR

- 2 For members requesting renewal (reauthorization), ALL of the following:
- **2.1** Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member has experienced positive clinical response to therapy with the requested drug

AND

2.2 For Takhzyro and Orladeyo requests ONLY: Submission of medical records (e.g., chart notes) supporting member has experienced no attacks during the preceding 12 months

AND

2.3 For Haegarda requests ONLY: Confirmation there are no weight changes warranting different quantity limits

different quartity fiffits	unicioni quantity ininio			
Notes	*Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on therapy, will have coverage under their drug benefit for the remainder of the curren t treatment course. Restrictions to specific network pharmacies and p articipation in medication management programs may apply.			
	**Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufact urer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies			

Product Name: Haegarda, Orladeyo, Takhzyro				
Diagnosis		Long-Term Prevention/Prophylaxis		
Approval Le	Approval Length 12 month(s)			
Therapy Sta	herapy Stage Initial Authorization			
Guideline Type Prior Authorization - IL and MN Plans				
Product Generic Name		GPI	Brand/Generic	

HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 2000 UNIT	85802022002130	Brand
HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 3000 UNIT	85802022002140	Brand
TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 150 MG/ML	8584204020E510	Brand
TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 300 MG/2ML (150 MG/ML)	8584204020E520	Brand
TAKHZYRO	LANADELUMAB-FLYO INJ 300 MG/2ML (150 MG/ML)	85842040202020	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 110 MG	85840010200120	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 150 MG	85840010200130	Brand

1 - Diagnosis of Hereditary Angioedema (HAE)

AND

- 2 One of the following:
 - History of ≥ 2 attacks per month
 - Symptoms are moderate to severe

AND

- 3 One of the following:
- 3.1 Low C4 AND low C1 inhibitor level or function

OR

- **3.2** Both of the following:
 - Normal C1 inhibitor level with a family history of HAE
 - High dose antihistamines did not control symptoms

AND

4 - Prescribed by or in consultation with an allergist or other provider experienced in the treatment of HAE

AND

5 - All medications that may cause angioedema have been discontinued (e.g., ACE inhibitors, estrogens, ARBs)

AND

6 - Requested medication will not be used in combination with other approved HAE prevention treatments

Product Name: Haegarda, Orladeyo, Takhzyro		
Diagnosis Long-Term Prevention/Prophylaxis		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 2000 UNIT	85802022002130	Brand
HAEGARDA	C1 ESTERASE INHIBITOR (HUMAN) FOR SUBCUTANEOUS INJ 3000 UNIT	85802022002140	Brand
TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 150 MG/ML	8584204020E510	Brand
TAKHZYRO	LANADELUMAB-FLYO SOLN PREF SYRINGE 300 MG/2ML (150 MG/ML)	8584204020E520	Brand
TAKHZYRO	LANADELUMAB-FLYO INJ 300 MG/2ML (150 MG/ML)	85842040202020	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 110 MG	85840010200120	Brand
ORLADEYO	BEROTRALSTAT HCL CAP 150 MG	85840010200130	Brand

Approval Criteria

1 - For members new to plan* (as evidenced by coverage effective date of less than or equal to 90 days)**: Submission of medical records (e.g., chart notes) documenting that within the

past 12 months the member has experienced positive clinical response to therapy with the requested drug

OR

- 2 For members requesting renewal (reauthorization), ALL of the following:
- **2.1** Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member has experienced positive clinical response to therapy with the requested drug

AND

2.2 For Takhzyro and Orladeyo requests ONLY: Submission of medical records (e.g., chart notes) supporting member has experienced no attacks during the preceding 12 months

AND

2.3 For Haegarda requests ONLY: Confirmation there are no weight changes warranting different quantity limits

Notes	*Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on therapy, will have coverage under their drug benefit for the remainder of the curren t treatment course. Restrictions to specific network pharmacies and p articipation in medication management programs may apply.
	**Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufact urer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

2. Revision History

Date	Notes
4/9/2024	Guideline Update.

Hetlioz (tasimelteon)			

Prior Authorization Guideline

Guideline ID	GL-131133
Guideline Name	Hetlioz (tasimelteon)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic tasimelteon, Hetlioz LQ			
Approval Length 12/31/2039			
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans			ns

Product Name	Generic Name	GPI	Brand/Generic
TASIMELTEON	TASIMELTEON CAPSULE 20 MG	60250070000130	Generic
HETLIOZ LQ	TASIMELTEON ORAL SUSP 4 MG/ML	60250070001820	Brand

Approval Criteria

- 1 One of the following:
 - **1.1** Both of the following:

1.1.1 Diagnosis of Smith-Magenis syndrome

AND

1.1.2 Trial and failure, contraindication, or intolerance to 3 months of melatonin

OR

- **1.2** All of the following:
- **1.2.1** Diagnosis of a non-24-hour sleep-wake disorder

AND

1.2.2 Member is completely blind

AND

1.2.3 Trial and failure, contraindication, or intolerance to 3 months of generic ramelteon

AND

1.2.4 Prescribed by, or in consultation with a sleep specialist

Product Name: Generic tasimelteon, Hetlioz LQ		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
TASIMELTEON	TASIMELTEON CAPSULE 20 MG	60250070000130	Generic
HETLIOZ LQ	TASIMELTEON ORAL SUSP 4 MG/ML	60250070001820	Brand

Approval Criteria 1 - One of the following: **1.1** Both of the following: 1.1.1 Diagnosis of Smith-Magenis syndrome AND **1.1.2** Trial and failure, contraindication, or intolerance to 3 months of melatonin OR **1.2** All of the following: **1.2.1** Diagnosis of a non-24-hour sleep-wake disorder **AND** 1.2.2 Member is completely blind **AND 1.2.3** Trial and failure, contraindication, or intolerance to 3 months of generic ramelteon AND **1.2.4** Prescribed by, or in consultation with a sleep specialist Notes *Members new to the plan (as evidenced by coverage effective date o f less than or equal to 90 days) must meet the initial criteria for covera ge

Product N	Name:	Generic i	tasimel	teon, I	Heti	lioz L	_C	Į
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Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TASIMELTEON	TASIMELTEON CAPSULE 20 MG	60250070000130	Generic
HETLIOZ LQ	TASIMELTEON ORAL SUSP 4 MG/ML	60250070001820	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Notes	*Members new to the plan (as evidenced by coverage effective date o
	f less than or equal to 90 days) must meet the initial criteria for covera
	ge

2. Revision History

Date	Notes
10/6/2023	2024 New Implementation

Human Chorionic Gonadotropin (Novarel, Pregnyl equivalents) and Clomiphene (Clomid) Prior Auth					
(a) The Manufalling areas to delighter. The day to be seen read, commerce and the state plants the covered and delicts.					

Prior Authorization Guideline

Guideline ID	GL-145618	
	Human Chorionic Gonadotropin (Novarel, Pregnyl equivalents) and Clomiphene (Clomid) Prior Auth	
Formulary	Quartz	

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Brand Clomid, generic clomiphene			
Approval Length	12/31/2039		
Guideline Type Prior Authorization - All Plans Except IL and MN Plans*			S*
Double to the Committee of the Committee			D

Product Name	Generic Name	GPI	Brand/Generic
CLOMID	CLOMIPHENE CITRATE TAB 50 MG	30066030100305	Brand
Clomiphene	CLOMIPHENE CITRATE TAB 50 MG	30066030100305	Generic

Approval Criteria

1 - All of the following:

1.1 Diagnosis of hypogonadism not seeking fertility treatment

AND

1.2 Submission of medical records (e.g., chart notes) documenting two low morning testosterone levels (or within 3 hours of waking for shift workers) including the normal ranges for the laboratory

AND

1.3 Submission of medical records (e.g., chart notes) documenting symptoms due to low testosterone other than sexual dysfunction or decreased libido

OR

2 - Submission of medical records (e.g., chart notes) documenting reduction of the uterine lining or fibroid size prior to surgical procedures, unrelated to infertility (e.g., endometriosis, uterine fibroids)

Notes	*Coverage of clomiphene for use in infertility is limited to members who have the artificial
	insemination rider attached to their benefit and coverage is limited to the duration and cost share amounts as defined in
	the rider.

Product Name: Brand Clomid, generic clomiphene	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
CLOMID	CLOMIPHENE CITRATE TAB 50 MG	30066030100305	Brand
Clomiphene	CLOMIPHENE CITRATE TAB 50 MG	30066030100305	Generic

Approval Criteria

1 - All of the following:	
1.1 Diagnosis of hypo	gonadism not seeking fertility treatment
	AND
	dical records (e.g., chart notes) documenting two low morning within 3 hours of waking for shift workers) including the normal ranges
	AND
	dical records (e.g., chart notes) documenting symptoms due to low sexual dysfunction or decreased libido
	OR
	cal records (e.g., chart notes) documenting reduction of the uterine or to surgical procedures, unrelated to infertility (e.g., endometriosis,
	OR
3 - For Illinois Plans ON	NLY: Both of the following:
3.1 Member has Quartz plan issued in the state of Illinois	
	AND
3.2 Provider attests patient has infertility coverage as outlined in Illinois Insurance Code 215 ILCS 5/356m	
Notes	*Coverage of clomiphene for use in infertility for MN plans is limited to members who have the artificial insemination rider attached to their benefit and coverage is limited to the duration and cost share amounts as defined in the rider.

Product Name: Brand Clomid, generic clomiphene		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans*		

Product Name	Generic Name	GPI	Brand/Generic
CLOMID	CLOMIPHENE CITRATE TAB 50 MG	30066030100305	Brand
Clomiphene	CLOMIPHENE CITRATE TAB 50 MG	30066030100305	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

*Coverage of clomiphene for use in infertility for MN plans is limited to members who have the artificial insemination rider attached to their benefit and coverage is limited to the duration and cost share amounts as defined in
he duration and cost share amounts as defined in
the rider.

Product Name: Brand Pregnyl, Brand Novarel, Brand Choionic gonadotropin		
Approval Length 12/31/2039		
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans*	

Product Name	Generic Name	GPI	Brand/Generic
NOVAREL	CHORIONIC GONADOTROPIN FOR IM INJ 5000 UNIT	30062020002130	Brand
NOVAREL	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand
PREGNYL W/DILUENT BENZYL ALCOHOL/NACL	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand
CHORIONIC GONADOTROPIN	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand

Approval Criteria

1 - Diagnosis of hypogonadism not seeking fertility treatment

AND

2 - Submission of medical records (e.g., chart notes) documenting two low morning testosterone levels (or within 3 hours of waking for shift workers) including the normal ranges for the laboratory

AND

3 - Submission of medical records (e.g., chart notes) documenting symptoms due to low testosterone other than sexual dysfunction or decreased libido

AND

4 - Trial and failure, contraindication or intolerance to clomiphene

AND

5 - The drug is being self-administered by the individual and not by a health care professional

*Coverage of chorionic gonadotropin for the treatment of hypogonadis m is limited to the prescription drug benefit and not covered on the medical benefit. Chorionic gonadotropin is covered on the medical benefit without restriction for indications that are not excluded from coverage based by the plan be nefits (e.g. treatment of prepuberal cryptorchidism). Diagnoses such as the treatment of infertility are excluded from coverage
uded from coverage.

Product Name: Brand Pregnyl, Brand Novarel, Brand Choionic gonadotropin			
Approval Length	Approval Length 12 month(s)		
Therapy Stage Initial Authorization			
Guideline Type	Prior Authorization - IL and MN Plans*		

Product Name	Generic Name	GPI	Brand/Generic
NOVAREL	CHORIONIC GONADOTROPIN FOR IM INJ 5000 UNIT	30062020002130	Brand

NOVAREL	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand
PREGNYL W/DILUENT BENZYL ALCOHOL/NACL	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand
CHORIONIC GONADOTROPIN	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand

- **1** All of the following must be met:
- **1.1** Diagnosis of hypogonadism not seeking fertility treatment

AND

1.2 Submission of medical records (e.g., chart notes) documenting two low morning testosterone levels (or within 3 hours of waking for shift workers) including the normal ranges for the laboratory

AND

1.3 Submission of medical records (e.g., chart notes) documenting symptoms due to low testosterone other than sexual dysfunction or decreased libido

AND

1.4 Trial and failure, contraindication or intolerance to clomiphene

AND

1.5 The drug is being self-administered by the individual and not by a health care professional

OR

- 2 For Illinois Plans Only: Both of the following:
- 2.1 Member has Quartz plan issued in the state of Illinois

AND

2.2 Provider attests patient has infertility coverage as outlined in Illinois Insurance Code 215 ILCS 5/356m

Notes	*Coverage of chorionic gonadotropin for the treatment of hypogonadis m is limited to the prescription drug benefit and not covered on the medical benefit. Chorionic gonadotropin is covered on the medical benefit without restriction for indications that are not excluded from coverage based by the plan be nefits (e.g. treatment of prepuberal cryptorchidism). Diagnoses such as the treatment of infertility are excluded from coverage. *Members new to the plan (as evidenced by coverage effective date of
	*Members new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) must meet initial criteria for coverage

Product Name: Brand Pregnyl, Brand Novarel, Brand Choionic gonadotropin		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans*		

Product Name	Generic Name	GPI	Brand/Generic
NOVAREL	CHORIONIC GONADOTROPIN FOR IM INJ 5000 UNIT	30062020002130	Brand
NOVAREL	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand
PREGNYL W/DILUENT BENZYL ALCOHOL/NACL	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand
CHORIONIC GONADOTROPIN	CHORIONIC GONADOTROPIN FOR IM INJ 10000 UNIT	30062020002140	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Notes	*Coverage of chorionic gonadotropin for the treatment of hypogonadis m is limited to the prescription drug benefit and not covered on the medical benefit. Chorionic gonadotropin is covered on the medical benefit without restriction for indications that are not excluded from coverage based by the plan be nefits (e.g. treatment of prepuberal cryptorchidism). Diagnoses such as the treatment of infertility are excluded.
	cryptorchidism). Diagnoses such as the treatment of infertility are excluded from coverage.
	*Members new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) must meet initial criteria for coverage

2. Revision History

Date	Notes
4/11/2024	Updated IL Specific criteria

Hydrocodone ER					
(a) The latest separate ballon, but the most count of the cold separate contributions.					

Prior Authorization Guideline

Guideline ID	GL-127837
Guideline Name	Hydrocodone ER
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Hydrocodone ER		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 10 MG	65100030106910	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 15 MG	65100030106915	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 20 MG	65100030106920	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 30 MG	65100030106930	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 40 MG	65100030106940	Generic

HYDROCODONE BITARTRATE ER		65100030106950	Generic
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- 1 Trial and failure of at least 2 of the following preferred long-acting opioids:
 - morphine ERT (generic of MS Contin)
 - morphine ERC (generic of Kadian)
 Oxycodone ER (Oxycontin)

OR

2 - For Minnesota Plans step therapy does not apply if member has stage four metastatic cancer and the requested drug is being used to treat cancer-related pain

Product Name: Generic Hydrocodone ER		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 10 MG	65100030106910	Generic
HYDROCODONE BITARTRATE ER		65100030106915	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 20 MG	65100030106920	Generic
HYDROCODONE BITARTRATE ER		65100030106930	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 40 MG	65100030106940	Generic
HYDROCODONE BITARTRATE ER		65100030106950	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Generic Hydrocodone ER		
Approval Length 12/31/2039		
Guideline Type Step Therapy - All plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
HYDROCODONE HYDROCODONE BITARTRATE CAP ER 12HR 10 BITARTRATE ER MG		65100030106910	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 15 MG	65100030106915	Generic
HYDROCODONE BITARTRATE ER		65100030106920	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 30 MG	65100030106930	Generic
HYDROCODONE BITARTRATE ER	HYDROCODONE BITARTRATE CAP ER 12HR 40 MG	65100030106940	Generic
HYDROCODONE BITARTRATE ER		65100030106950	Generic

- 1 Trial and failure of at least 2 of the following preferred long-acting opioids:
 - morphine ERT (generic of MS Contin) morphine ERC (generic of Kadian) Oxycodone ER (Oxycontin)

2. Revision History

Date	Notes
8/25/2023	New Program

Inb	rija (Le	v odopa	inhalat	ion po	wder)
The historings on	ennt kraligiajust. Trafiis nay haas kann mesal, seram	i, or dichest. Verily that has bits pointers that connectifie a	of looker.		

Prior Authorization Guideline

Guideline ID	GL-129635	
Guideline Name	Inbrija (Levodopa inhalation powder)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Inbrija		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
INBRIJA	LEVODOPA INHAL POWDER CAP 42 MG	73200040000160	Brand

Approval Criteria

1 - Diagnosis of Parkinson's disease

AND

2 - Prescribed by, or in consultation with, a Neurologist

AND

3 - Current treatment with combination of long-acting and short-acting carbidopa/levodopa

AND

4 - Person experiencing intermittent "off" episodes despite adequate trial with appropriate dosage adjustment with carbidopa/levodopa

Product Name: Inbrija				
Approval Length 12 month(s)				
Therapy St	herapy Stage Reauthorization			
Guideline Type Prior Authorization - IL and MN Plans				
Product Generic Name GPI Bran		Brand/Generic		

Product Name	Generic Name	GPI	Brand/Generic
INBRIJA	LEVODOPA INHAL POWDER CAP 42 MG	73200040000160	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Inbrija				
Approval Length 12/31/2039				
Therapy St	age	Initial Authorization		
Guideline Type Prior Authorization - All plans except IL and MN				
Product Generic Name GI		GPI	Brand/Generic	

INBRIJA	LEVODOPA INHAL POWDER CAP 42 MG	73200040000160	Brand
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1 - Diagnosis of Parkinson's disease

AND

2 - Prescribed by, or in consultation with, a Neurologist

AND

3 - Current treatment with combination of long-acting and short-acting carbidopa/levodopa

AND

4 - Person experiencing intermittent "off" episodes despite adequate trial with appropriate dosage adjustment with carbidopa/levodopa

2. Revision History

Date	Notes
10/6/2023	New Program

	Increlex (mecasermin)		
	Control trapperson straighter. This has been created another and soft below to provide an extraple of the control and the cont		

Prior Authorization Guideline

Guideline ID	GL-129115
Guideline Name	Increlex (mecasermin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Increlex				
Approval Length Guideline Type		12/31/2039		
		Prior Authorization - ALL Plans Except IL and MN Plans*		
Product Name	Generic Na	me	GPI	Brand/Generic
INCRELEX	MECASERMI	N INJ 40 MG/4ML (10 MG/ML)	30160045002020	Brand

Approval Criteria

- 1 One of the following:
 - **1.1** Diagnosis of one of the following:

- Primary insulin-like growth factor deficiency (IGFD)
- Low insulin-like growth factor-1 (IGF-1) levels
- Growth hormone deletion with neutralizing antibodies to growth hormone

OR

1.2 Submission of medical records (e.g., chart notes) documenting lack of response to growth hormone following therapeutic trial of somatropin (for members with growth hormone deficiency)

AND

2 - Member is less than 18 years of age

AND

3 - Member has confirmed open epiphyses

AND

4 - Prescribed by or in consultation with a pediatric endocrinologist

Notes	*Increlex is not indicated to treat secondary IGFD due to GH deficienc y, malnutrition, hypothyroidism or other
	causes *Increlex is not covered for treatment of idiopathic short stature
	*Increlex is not a substitute for growth hormone (somatropin)

Product Name: Increlex	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
INCRELEX	MECASERMIN INJ 40 MG/4ML (10 MG/ML)	30160045002020	Brand

- 1 One of the following:
- **1.1** Diagnosis of one of the following:
 - Primary insulin-like growth factor deficiency (IGFD)
 - Low insulin-like growth factor-1 (IGF-1) levels
 - Growth hormone deletion with neutralizing antibodies to growth hormone

OR

1.2 Submission of medical records (e.g., chart notes) documenting lack of response to growth hormone following therapeutic trial of somatropin (for members with growth hormone deficiency)

AND

2 - Member is less than 18 years of age

AND

3 - Member has confirmed open epiphyses

AND

4 - Prescribed by or in consultation with a pediatric endocrinologist

·	-
	*Increlex is not indicated to treat secondary IGFD due to GH deficienc y, malnutrition, hypothyroidism or other
	causes
	*Increlex is not covered for treatment of idiopathic short stature *Increlex is not a substitute for growth hormone (somatropin)

Product Name: Increlex	
Approval Length	12 month(s)

Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
INCRELEX	MECASERMIN INJ 40 MG/4ML (10 MG/ML)	30160045002020	Brand

1	- (One	e of	the	fol	low	ing	٠

- **1.1** Diagnosis of one of the following:
 - Primary insulin-like growth factor deficiency (IGFD)
 - Low insulin-like growth factor-1 (IGF-1) levels
 - Growth hormone deletion with neutralizing antibodies to growth hormone

OR

1.2 Submission of medical records (e.g., chart notes) documenting lack of response to growth hormone following therapeutic trial of somatropin (for members with growth hormone deficiency)

AND

2 - Member is less than 18 years of age

AND

3 - Member has confirmed open epiphyses

AND

4 - Prescribed by or in consultation with a pediatric endocrinologist

5 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months, the member remains on therapy				
Notes	*Increlex is not indicated to treat secondary IGFD due to GH deficienc y, malnutrition, hypothyroidism or other causes *Increlex is not covered for treatment of idiopathic short stature *Increlex is not a substitute for growth hormone (somatropin)			

2. Revision History

Date	Notes
7/31/2023	2024 New Implementation

I	Ingrez	za (va	lbena	zine)		
-	The Inhael Image current has displayed. Th	efic ney have have nessel, vecamed, or dele	ed. Verily that he liek points in the convenille.	ed Inadios.		

Prior Authorization Guideline

Guideline ID	GL-130583	
Guideline Name	Ingrezza (valbenazine)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Ingrezza				
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
INGREZZA VALBENAZINE TOSYLATE CAP THERAPY PACK 40 MG (7) & 80 MG (21)		6238008020B220	Brand	

INGREZZA	VALBENAZINE TOSYLATE CAP 40 MG (BASE EQUIV)	62380080200120	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 60 MG (BASE EQUIV)	62380080200130	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 80 MG (BASE EQUIV)	62380080200140	Brand

1 - Diagnosis of tardive dyskinesia (TD)

AND

- **2** Prescribed by or in consultation with one of the following:
 - Neurologist
 - Psychiatrist
 - Specialist in the treatment of TD

AND

- 3 One of the following:
- 3.1 Symptoms persist despite stopping the dopamine receptor blocking drug that caused TD

OR

3.2 Submission of medical records (e.g., chart notes) documenting evidence-based clinical rationale why discontinuation of the drug is not a treatment option for the person based on their diagnosis and previous treatment history

AND

4 - Trial and failure, contraindication, or intolerance to clonazepam

5 - (For patients whose primary symptomology is tardive dystonia ONLY): Trial and failure, contraindication, or intolerance to trihexyphenidyl

Product Name: Ingrezza		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
INGREZZA	VALBENAZINE TOSYLATE CAP THERAPY PACK 40 MG (7) & 80 MG (21)	6238008020B220	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 40 MG (BASE EQUIV)	62380080200120	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 60 MG (BASE EQUIV)	62380080200130	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 80 MG (BASE EQUIV)	62380080200140	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Ingrezza		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
INGREZZA	VALBENAZINE TOSYLATE CAP THERAPY PACK 40 MG (7) & 80 MG (21)	6238008020B220	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 40 MG (BASE EQUIV)	62380080200120	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 60 MG (BASE EQUIV)	62380080200130	Brand
INGREZZA	VALBENAZINE TOSYLATE CAP 80 MG (BASE EQUIV)	62380080200140	Brand

Approval Criteria

1 - Diagnosis of tardive dyskinesia (TD)

- 2 Prescribed by or in consultation with one of the following:
 - Neurologist
 - Psychiatrist
 - Specialist in the treatment of TD

AND

- **3** One of the following:
- 3.1 Symptoms persist despite stopping the dopamine receptor blocking drug that caused TD

OR

3.2 Submission of medical records (e.g., chart notes) documenting evidence-based clinical rationale why discontinuation of the drug is not a treatment option for the person based on their diagnosis and previous treatment history

AND

4 - Trial and failure, contraindication, or intolerance to clonazepam

AND

5 - (For patients whose primary symptomology is tardive dystonia ONLY): Trial and failure, contraindication, or intolerance to trihexyphenidyl

2. Revision History

Date	Notes
8/16/2023	2024 New Implementation

Inhaled Bronchodilators for Chronic Obstructive Pulmonary Dise			
	(3) Thinkings are in display has been used count or date that has been been counted as earth.		

Prior Authorization Guideline

Guideline ID GL-129738		
Guideline Name	Inhaled Bronchodilators for Chronic Obstructive Pulmonary Disease	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Stiolto Respimat	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
	TIOTROPIUM BR-OLODATEROL INHAL AERO SOLN 2.5-2.5 MCG/ACT	44209902923420	Brand

Approval Criteria

1 - Diagnosis of chronic obstructive pulmonary disease (COPD)

2 - Trial and failure, contraindication or intolerance to use of umeclidinium/vilanterol (Anoro Ellipta)

Product Name: Stiolto Respimat	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
STIOLTO RESPIMAT	TIOTROPIUM BR-OLODATEROL INHAL AERO SOLN 2.5-2.5 MCG/ACT	44209902923420	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Stiolto Respimat	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
	TIOTROPIUM BR-OLODATEROL INHAL AERO SOLN 2.5-2.5 MCG/ACT	44209902923420	Brand

Approval Criteria

1 - Diagnosis of chronic obstructive pulmonary disease (COPD)

2 - Trial and failure, contraindication or intolerance to use of umeclidinium/vilanterol (Anoro Ellipta)

2. Revision History

Date	Notes
10/6/2023	New Program

In	haled	Cortico	steroi	d Step	thera	ру
E Person	Design current to oliquiques. The file may have be	en reconst, unament, or deleted. Verily that the list ye	in the constituted basin.			

Prior Authorization Guideline

Guideline ID	GL-143611
Guideline Name	Inhaled Corticosteroid Step therapy
Formulary	Quartz

Guideline Note:

Effective Date:	4/1/2024
P&T Approval Date:	2/15/2022
P&T Revision Date:	7/18/2023

1. Criteria

HFA

HFA

ASMANEX

SUSPENSION 100 MCG/ACT

SUSPENSION 200 MCG/ACT

MOMETASONE FUROATE INHAL AEROSOL

Product Name: Asmanex, Asmanex HFA				
Approval Len	gth	12 month(s)		
Therapy Stag	erapy Stage Initial Authorization			
Guideline Ty	Гуре Step Therapy - IL and MN Plans			
Product Name	Generic Name		GPI	Brand/Generic
ASMANEX HFA	MOMETASONE FUROATE INHAL AEROSOL SUSPENSION 50 MCG/ACT		44400036203210	Brand
ASMANEX	MOMETAS	ONE FUROATE INHAL AEROSOL	44400036203220	Brand

Brand

44400036203230

ASMANEX TWISTHALER 30 METERED DOSES	MOMETASONE FUROATE INHAL POWD 110 MCG/ACT (BREATH ACTIVATED)	44400036208010	Brand
ASMANEX TWISTHALER 14 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 60 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 120 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 30 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand

- **1** Trial and failure, intolerance, or contraindication to one of the following:

 - an inhaled fluticasone propionate product an inhaled fluticasone furoate product

Product Name: Asmanex, Asmanex HFA		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ASMANEX HFA	MOMETASONE FUROATE INHAL AEROSOL SUSPENSION 50 MCG/ACT	44400036203210	Brand
ASMANEX HFA	MOMETASONE FUROATE INHAL AEROSOL SUSPENSION 100 MCG/ACT	44400036203220	Brand
ASMANEX HFA	MOMETASONE FUROATE INHAL AEROSOL SUSPENSION 200 MCG/ACT	44400036203230	Brand
ASMANEX TWISTHALER	MOMETASONE FUROATE INHAL POWD 110 MCG/ACT (BREATH ACTIVATED)	44400036208010	Brand

30 METERED DOSES			
ASMANEX TWISTHALER 14 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 60 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 120 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 30 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand

1 - Submission of medical notes (e.g. chart notes) from the past 12 months that member is continuing therapy with the requested drug

Product Name: Asmanex, Asmanex HFA		
Approval Length	12/31/2039	
Guideline Type Step Therapy - All plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
ASMANEX HFA	MOMETASONE FUROATE INHAL AEROSOL SUSPENSION 50 MCG/ACT	44400036203210	Brand
ASMANEX HFA	MOMETASONE FUROATE INHAL AEROSOL SUSPENSION 100 MCG/ACT	44400036203220	Brand
ASMANEX HFA	MOMETASONE FUROATE INHAL AEROSOL SUSPENSION 200 MCG/ACT	44400036203230	Brand
ASMANEX TWISTHALER 30 METERED DOSES	MOMETASONE FUROATE INHAL POWD 110 MCG/ACT (BREATH ACTIVATED)	44400036208010	Brand
ASMANEX TWISTHALER 14 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand

ASMANEX TWISTHALER 60 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 120 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand
ASMANEX TWISTHALER 30 METERED DOSES	MOMETASONE FUROATE INHAL POWD 220 MCG/ACT (BREATH ACTIVATED)	44400036208020	Brand

- **1** Trial and failure, intolerance, or contraindication to one of the following:
 - an inhaled fluticasone propionate product an inhaled fluticasone furoate product

2. Revision History

Date	Notes
2/28/2024	New Program

Injectable Calcitonin Gene-Related Peptide (CGRP) Inhibit				
	(g Milating our habite. Note in hack much count a loss set to the his part to earth and and			

Prior Authorization Guideline

Guideline ID GL-144870		
Guideline Name	e Injectable Calcitonin Gene-Related Peptide (CGRP) Inhibitors	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	5/16/2018
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Aimovig, Emgality				
Diagnosis		Preventative Treatment of Migraine		
Approval Length		12/31/2039*		
Guideline ¹	Туре	Prior Authorization - ALL Plans Except IL and MN Plans		
Product Generic Nar Name AIMOVIG ERENUMAB-AINJECTOR 70		me	GPI	Brand/Generic
		AOOE SUBCUTANEOUS SOLN AUTO-) MG/ML	6770202010D520	Brand
AIMOVIG	AIMOVIG ERENUMAB-AOOE SUBCUTANEOUS SOLN AUTO-INJECTOR 140 MG/ML		6770202010D540	Brand
		MAB-GNLM SUBCUTANEOUS SOLN TOR 120 MG/ML	6770203530D520	Brand

EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 100 MG/ML	6770203530E515	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 120 MG/ML	6770203530E520	Brand

1 - Submission of medical records (e.g., chart notes) documenting member has greater than or equal to 4 migraine days per month with headaches that are disabling (e.g., unable to work/attend school, unable to participate in activities of daily living [ADLs], moderate to severe MIDAS score)

AND

2 - Trial and failure, intolerance, or contraindication to 2 generic preventive migraine medications (e.g., anti-hypertensives, antiepileptics, antidepressants, botulinum toxin)

Notes	*For new starts to galcanezumab therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.08 (2 every 28 days) for 30 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Aimovig, Emgality	
Diagnosis Preventative Treatment of Migraine	
Approval Length	12 months*
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
AIMOVIG	ERENUMAB-AOOE SUBCUTANEOUS SOLN AUTO-INJECTOR 70 MG/ML	6770202010D520	Brand
AIMOVIG	ERENUMAB-AOOE SUBCUTANEOUS SOLN AUTO- INJECTOR 140 MG/ML	6770202010D540	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN AUTO-INJECTOR 120 MG/ML	6770203530D520	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 100 MG/ML	6770203530E515	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 120 MG/ML	6770203530E520	Brand

1 - Submission of medical records (e.g., chart notes) documenting member has greater than or equal to 4 migraine days per month with headaches that are disabling (e.g., unable to work/attend school, unable to participate in activities of daily living [ADLs], moderate to severe MIDAS score)

AND

2 - Trial and failure, intolerance, or contraindication to 2 generic preventive migraine medications (e.g., anti-hypertensives, antiepileptics, antidepressants, botulinum toxin)

Notes	*For new starts to galcanezumab therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.08 (2 every 28 days) for
	30 days
	Second PA: Approve at GPI 10 for 12 months

Product Name: Aimovig, Emgality		
Diagnosis	Preventative Treatment of Migraine	
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans*	

Product Name	Generic Name	GPI	Brand/Generic
AIMOVIG	ERENUMAB-AOOE SUBCUTANEOUS SOLN AUTO-INJECTOR 70 MG/ML	6770202010D520	Brand
AIMOVIG	ERENUMAB-AOOE SUBCUTANEOUS SOLN AUTO- INJECTOR 140 MG/ML	6770202010D540	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN AUTO-INJECTOR 120 MG/ML	6770203530D520	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 100 MG/ML	6770203530E515	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 120 MG/ML	6770203530E520	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the previous 12 months the member is showing a response to therapy (e.g., symptom improvement such as decreased frequency or severity of headaches from baseline, reduced cluster headache frequency, improved ability to participate in therapies/ADLs, improved MIDAS score, less acute medication use, fewer ER/UC visits for migraine, ability to return to work/school)

Product Name: Ajovy		
Diagnosis	Preventative Treatment of Migraine	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - ALL Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
AJOVY	FREMANEZUMAB-VFRM SUBCUTANEOUS SOLN AUTO-INJ 225 MG/1.5ML	6770203020D520	Brand
AJOVY	FREMANEZUMAB-VFRM SUBCUTANEOUS SOLN PREF SYR 225 MG/1.5ML	6770203020E520	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting member has greater than or equal to 4 migraine days per month with headaches that are disabling (e.g., unable to work/attend school, unable to participate in activities of daily living [ADLs], moderate to severe MIDAS score)

AND

2 - Trial and failure, contraindication or intolerance to 2 generic preventive migraine medications (e.g., anti-hypertensives, antiepileptics, antidepressants, botulinum toxin)

- **3** Trial and failure, contraindication or intolerance to both of the following:
 - Aimovig
 - Emgality

Product Name: Ajovy		
Diagnosis	Preventative Treatment of Migraine	
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans*	

Product Name	Generic Name	GPI	Brand/Generic
AJOVY	FREMANEZUMAB-VFRM SUBCUTANEOUS SOLN AUTO-INJ 225 MG/1.5ML	6770203020D520	Brand
AJOVY	FREMANEZUMAB-VFRM SUBCUTANEOUS SOLN PREF SYR 225 MG/1.5ML	6770203020E520	Brand

1 - Submission of medical records (e.g., chart notes) documenting member has greater than or equal to 4 migraine days per month with headaches that are disabling (e.g., unable to work/attend school, unable to participate in activities of daily living [ADLs], moderate to severe MIDAS score)

AND

2 - Trial and failure, contraindication or intolerance to 2 generic preventive migraine medications (e.g., anti-hypertensives, antiepileptics, antidepressants, botulinum toxin)

- 3 Trial and failure, contraindication or intolerance to both of the following:
 - Aimovig
 - Emgality

Product Name: Ajovy		
Diagnosis	Preventative Treatment of Migraine	
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	

Guideline Type Prior Authorization - IL and MN Plans*		าร*		
Product Name	Generic Name		GPI	Brand/Generic
AJOVY	FREMANEZUMAB-VFRM SUBCUTANEOUS SOLN AUTO-INJ 225 MG/1.5ML		6770203020D520	Brand
AJOVY	AJOVY FREMANEZUMAB-VFRM SUBCUTANEOUS SOLN PREF SYR 225 MG/1.5ML		6770203020E520	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the previous 12 months the member is showing a response to therapy (e.g., symptom improvement such as decreased frequency or severity of headaches from baseline, reduced cluster headache frequency, improved ability to participate in therapies/ADLs, improved MIDAS score, less acute medication use, fewer ER/UC visits for migraine, ability to return to work/school)

Product Name: Emgality		
Diagnosis Episodic Cluster Headache		
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - ALL Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN AUTO-INJECTOR 120 MG/ML	6770203530D520	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 100 MG/ML	6770203530E515	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 120 MG/ML	6770203530E520	Brand

Approval Criteria

1 - Diagnosis of cluster headaches that are not rebound headaches due to medication overuse

AND

2 - Patient is 18 years of age or older

Product Name: Emgality		
Diagnosis Episodic Cluster Headache		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans*		

Product Name	Generic Name	GPI	Brand/Generic
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN AUTO-INJECTOR 120 MG/ML	6770203530D520	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 100 MG/ML	6770203530E515	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 120 MG/ML	6770203530E520	Brand

1 - Diagnosis of cluster headaches that are not rebound headaches due to medication overuse

AND

2 - Patient is 18 years of age or older

Product Name: Emgality		
Diagnosis Episodic Cluster Headache		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans*		

Product Name	Generic Name	GPI	Brand/Generic
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN AUTO-INJECTOR 120 MG/ML	6770203530D520	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 100 MG/ML	6770203530E515	Brand
EMGALITY	GALCANEZUMAB-GNLM SUBCUTANEOUS SOLN PREFILLED SYR 120 MG/ML	6770203530E520	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the previous 12 months the member is showing a response to therapy (e.g., symptom improvement such as decreased frequency or severity of headaches from baseline, reduced cluster headache frequency, improved ability to participate in therapies/ADLs, improved MIDAS score, less acute medication use, fewer ER/UC visits for migraine, ability to return to work/school)

2. Revision History

Date	Notes
3/26/2024	New Guideline

Interferons
The Mark Programment for English, The Bits may have been record, wreward, or deleter law for the bits problem that convenils and leaders.

Prior Authorization Guideline

Guideline ID	GL-130130	
Guideline Name	Interferons	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Alferon N				
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization		
Product Generic Name		GPI	Brand/Generic	
ALFERON INTERFERON		N ALFA-N3 INJ 5000000 UNIT/ML	21700060302020	Brand

1 - Diagnosis of external genital or perianal warts

AND

2 - Must be self-administered or administered by family member or caretaker

Product Name: Alferon N		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization		

Product Name	Generic Name	GPI	Brand/Generic
ALFERON N	INTERFERON ALFA-N3 INJ 5000000 UNIT/ML	21700060302020	Brand

Approval Criteria

1 - Diagnosis of external genital or perianal warts

AND

2 - Must be self-administered or administered by family member or caretaker

AND

3 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member demonstrates positive clinical response to therapy

Product Name: Actimmune	
Approval Length	12 month(s)

Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
ACTIMMUNE	INTERFERON GAMMA-1B INJ 100 MCG/0.5ML (2000000 UNIT/0.5ML)	21700060702020	Brand

- 1 Diagnosis of ONE of the following:
 - Chronic granulomatous disease
 - Congenital malignant osteopetrosis

AND

2 - Must be self-administered or administered by family member or caretaker

Product Name: Actimmune		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization		

Product Name	Generic Name	GPI	Brand/Generic
ACTIMMUNE	INTERFERON GAMMA-1B INJ 100 MCG/0.5ML (2000000 UNIT/0.5ML)	21700060702020	Brand

Approval Criteria

- 1 Diagnosis of ONE of the following:
 - Chronic granulomatous disease
 - Congenital malignant osteopetrosis

2 - Must be self-administered or administered by family member or caretaker

AND

3 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member demonstrates positive clinical response to therapy

2. Revision History

Date	Notes
8/15/2023	2024 New Implementation

Itraconazole/Onychomycosis
The behaviory control algorithm is the first and control or seem to the first his possible control and

Prior Authorization Guideline

Guideline ID	GL-145447
Guideline Name	Itraconazole/Onychomycosis
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	4/17/2013
P&T Revision Date:	7/18/2023

Note:

For systemic infections only: Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Itraconazole (generic Sporanox)				
Diagnosis		Onychomycosis		
Approval Length	1	4 month(s)		
Guideline Type Prior Authorization – All plans except IL and MN				
Product Generic Name		GPI	Brand/Generic	

ITRACONAZOLE	ITRACONAZOLE CAP 100 MG	11407035000120	Generic
ITRACONAZOLE	ITRACONAZOLE ORAL SOLN 10 MG/ML	11407035002020	Generic

1 - Diagnosis of fungal or mold infection confirmed by culture or positive KOH test

AND

- 2 Submission of medical records (e.g., chart notes) documenting one of the following:
 - functional disability due to onychomycosis
 - peripheral vascular disease
 - diabetes
 - immunosuppressed or immunocompromised state
 - history of recurrent cellulitis

AND

3 - Trial and failure, contraindication, or intolerance to terbinafine therapy

Product Name: Itraconazole (generic Sporanox)		
Diagnosis	Onychomycosis	
Approval Length	12 month(s) with a fill count = 4	
Guideline Type	Prior Authorization- IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ITRACONAZOLE	ITRACONAZOLE CAP 100 MG	11407035000120	Generic
ITRACONAZOLE	ITRACONAZOLE ORAL SOLN 10 MG/ML	11407035002020	Generic

Approval Criteria

1 - Diagnosis of fungal or mold infection confirmed by culture or positive KOH test

- 2 Submission of medical records (e.g., chart notes) documenting one of the following:
 - functional disability due to onychomycosis
 - peripheral vascular disease
 - diabetes
 - immunosuppressed or immunocompromised state
 - history of recurrent cellulitis

AND

3 - Trial and failure, contraindication, or intolerance to terbinafine therapy

Product Name: Jublia, Brand Kerydin, Generic tavaborole		
Diagnosis	Onychomycosis	
Approval Length	6 month(s)	
Guideline Type	Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
JUBLIA	EFINACONAZOLE SOLN 10%	90154037002020	Brand
KERYDIN	TAVABOROLE SOLN 5%	90156080002010	Brand
TAVABOROLE SOLN 5%	TAVABOROLE SOLN 5%	90156080002010	Generic

Approval Criteria

1 - Diagnosis of fungal or mold infection confirmed by culture or positive KOH test

- 2 Submission of medical records (e.g., chart notes) documenting one of the following:
 - functional disability due to onychomycosis
 - peripheral vascular disease

- diabetes
- immunosuppressed or immunocompromised state
- history of recurrent cellulitis

- **3** Trial and failure, contraindication, or intolerance to both of the following:
 - Oral terbinafine
 - Oral itraconazole

Product Name: Jublia, Brand Kerydin, Generic tavaborole		
Diagnosis Onychomycosis		
Approval Length 12 month(s) with a fill count = 6		
Guideline Type Prior Authorization – IL and MN plans		

Product Name	Generic Name	GPI	Brand/Generic
JUBLIA	EFINACONAZOLE SOLN 10%	90154037002020	Brand
KERYDIN	TAVABOROLE SOLN 5%	90156080002010	Brand
TAVABOROLE	TAVABOROLE SOLN 5%	90156080002010	Generic

Approval Criteria

1 - Diagnosis of fungal or mold infection confirmed by culture or positive KOH test

- **2** Submission of medical records (e.g., chart notes) documenting one of the following:
 - functional disability due to onychomycosis
 - peripheral vascular disease
 - diabetes
 - immunosuppressed or immunocompromised state
 - history of recurrent cellulitis

- **3** Trial and failure, contraindication, or intolerance to both of the following:
 - Oral terbinafine
 - Oral itraconazole

Product Name: Tolsura		
Diagnosis Onychomycosis		
Approval Length 12 month(s) with fill count = 6		
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Name		Brand/Generic	
TOLSURA	ITRACONAZOLE CAP 65 MG	11407035000113	Brand	

Approval Criteria

1 - Diagnosis of fungal or mold infection confirmed by culture or positive KOH test

AND

- **2** Submission of medical records (e.g., chart notes) documenting one of the following:
 - functional disability due to onychomycosis
 - peripheral vascular disease
 - diabetes
 - immunosuppressed or immunocompromised state
 - history of recurrent cellulitis

AND

3 - Trial and failure, contraindication, or intolerance to use of generic itraconazole (Sporanox equivalent)

Product Name: Tolsura	
Diagnosis	Onychomycosis
Approval Length 6 month(s)	
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TOLSURA	ITRACONAZOLE CAP 65 MG	11407035000113	Brand

1 - Diagnosis of fungal or mold infection confirmed by culture or positive KOH test

AND

- 2 Submission of medical records (e.g., chart notes) documenting one of the following:
 - functional disability due to onychomycosis
 - peripheral vascular disease
 - diabetes
 - immunosuppressed or immunocompromised state
 - history of recurrent cellulitis

ITRACONAZOLE ITRACONAZOLE CAP 100 MG

AND

3 - Trial and failure, contraindication, or intolerance to use of generic itraconazole (Sporanox equivalent)

Product Name: Itraconazole (generic Sporanox)					
Diagnosis		Systemic Infections			
Approval Length		12 month(s)			
Therapy Stage		Initial Authorization			
Guideline Type Product Generi Name		Prior Authorization			
		ic Name	GPI	Brand/Generic	

Generic

11407035000120

ITRACONAZOLE	ITRACONAZOLE ORAL SOLN 10 MG/ML	11407035002020	Generic
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1 - Diagnosis of Blastomycosis, Histoplasmosis, Aspergillosis or other verified systemic fungal infection susceptible to itraconazole

OR

2 - (Illinois plans only): The drug is being used for the long-term treatment of tick-borne disease

Product Name: Tolsura		
Diagnosis	Systemic Infections	
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
TOLSURA	ITRACONAZOLE CAP 65 MG	11407035000113	Brand

Approval Criteria

- 1 One of the following:
- **1.1** Diagnosis of Blastomycosis, Histoplasmosis, Aspergillosis or other verified systemic fungal infection susceptible to itraconazole

OR

1.2 (Illinois Plans Only): The drug is being used for the long-term treatment of tick-borne disease

2 - Trial and failure, contraindication, or intolerance to use of generic itraconazole (Sporanox equivalent)

Product Name: Itraconazole (generic Sporanox), Tolsura		
Diagnosis	Systemic Infections	
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic	
ITRACONAZOLE	ITRACONAZOLE CAP 100 MG	11407035000120	Generic	
ITRACONAZOLE	ITRACONAZOLE ORAL SOLN 10 MG/ML	11407035002020	Generic	
TOLSURA	ITRACONAZOLE CAP 65 MG	11407035000113	Brand	

Approval Criteria

1 - New to the plan (within the past 90 days and submission of medical records (e.g., chart notes) documenting that within the past 12 months treatment of the condition is ongoing

OR

- **2** BOTH of the following:
- **2.1** ONE of the following:
 - Diagnosis of Blastomycosis, Histoplasmosis, Aspergillosis or other verified systemic fungal infection susceptible to itraconazole
 - (Illinois plans only): The drug is being used for the long-term treatment of tick-borne disease

AND

2.2 Submission of medical records (e.g., chart notes) documenting that within the past 12 months treatment of the condition is ongoing

2. Revision History

Date	Notes
4/5/2024	Guideline Update.

•	Juxtapid (lomitapide)									
	Debadage	ument to display set. The fil	a may have been necreal, where	al, or deleted. Verify that the life yes	the terroriffs and institu-					

Prior Authorization Guideline

Guideline ID	GL-136594
Guideline Name	Juxtapid (lomitapide)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

1. Criteria

Product Name: Juxtapid				
Approval Length 12 month(s)				
Therapy Stage Initial Authorization				
Guideline Type		Prior Authorization – All Plans		
Product Name	Generic Name		GPI	Brand/Generic
JUXTAPID	LOMITAPIDE MESYLATE CAP 5 MG (BASE EQUIV)		39480050200120	Brand
JUXTAPID	LOMITAPIDE MESYLATE CAP 10 MG (BASE EQUIV)		39480050200130	Brand

JUXTAPID	LOMITAPIDE MESYLATE CAP 20 MG (BASE EQUIV)	39480050200140	Brand
JUXTAPID	LOMITAPIDE MESYLATE CAP 30 MG (BASE EQUIV)	39480050200150	Brand

- **1** Diagnosis of homozygous familial hypercholesteremia (HoFH) with one of the following:
 - Clinical diagnosis (LDL-C greater than 500 mg/dL with xanthomas or family history of both parents with LDL-C levels greater than 250 mg/dL)
 - Genetic verification of HoFH

AND

2 - Prescribed by, or in consultation with, a Cardiologist or other specialist in the treatment of congenital lipid disorders

AND

3 - LDL-C level is greater than 70 mg/dL

AND

4 - Trial and failure, contraindication, or intolerance to a proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor unless there is genetic verification of receptor negative (null-null mutation) HoFH

Product Name: Juxtapid	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
JUXTAPID	LOMITAPIDE MESYLATE CAP 5 MG (BASE EQUIV)	39480050200120	Brand
JUXTAPID	LOMITAPIDE MESYLATE CAP 10 MG (BASE EQUIV)	39480050200130	Brand

JUXTAPID	LOMITAPIDE MESYLATE CAP 20 MG (BASE EQUIV)	39480050200140	Brand
JUXTAPID	LOMITAPIDE MESYLATE CAP 30 MG (BASE EQUIV)	39480050200150	Brand

- **1** Diagnosis of homozygous familial hypercholesteremia (HoFH) with one of the following:
 - Clinical diagnosis (LDL-C greater than 500 mg/dL with xanthomas or family history of both parents with LDL-C levels greater than 250 mg/dL)
 - Genetic verification of HoFH

AND

2 - Prescribed by, or in consultation with, a Cardiologist or other specialist in the treatment of congenital lipid disorders

AND

3 - Submission of medical records (e.g., chart notes) documenting a clinically meaningful (at least 10%) reduction in LDL-C from baseline

Product Name: Juxtapid		
Approval Length 12/31/2039		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
JUXTAPID	LOMITAPIDE MESYLATE CAP 5 MG (BASE EQUIV)	39480050200120	Brand
JUXTAPID	LOMITAPIDE MESYLATE CAP 10 MG (BASE EQUIV)	39480050200130	Brand
JUXTAPID	LOMITAPIDE MESYLATE CAP 20 MG (BASE EQUIV)	39480050200140	Brand
JUXTAPID	LOMITAPIDE MESYLATE CAP 30 MG (BASE EQUIV)	39480050200150	Brand

Approval Criteria

- 1 Diagnosis of homozygous familial hypercholesteremia (HoFH) with one of the following:
 - Clinical diagnosis (LDL-C greater than 500 mg/dL with xanthomas or family history of both parents with LDL-C levels greater than 250 mg/dL)
 - Genetic verification of HoFH

AND

2 - Prescribed by, or in consultation with, a Cardiologist or other specialist in the treatment of congenital lipid disorders

AND

3 - Submission of medical records (e.g., chart notes) documenting a clinically meaningful (at least 10%) reduction in LDL-C from baseline

Date	Notes
11/20/2023	2024 New Implementation

Jynarque (Tolvaptan)				
(2) International analysis for the fact that county and a state of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and analysis of the fact to provide an extended and an extended an extended and an extended and an extended an extended and an extended and an extended an extended and an extended an extended and an extended and an extended an extended and an extended an extended an extended and an extended an extended and an extended an extended and an extended and an extended an extended and an extended an extended and an extended and an extended an extended and an extended and an extended and an extended an extended and an extended and an extended an extended and an extended an extended and an extended and an extended and an extended an extended and an exte				

Guideline ID	GL-144542
Guideline Name	Jynarque (Tolvaptan)
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Brand Jynarque, Jynarque PAK, generic tolvaptan			
Approval Length 12 month(s)			
Therapy Stage Initial Authorization			
Guideline Type Prior Authorization			

Product Name	Generic Name	GPI	Brand/Generic
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 15 MG	3045406000B710	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 30 & 15 MG	3045406000B720	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 45 & 15 MG	3045406000B725	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 60 & 30 MG	3045406000B735	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 90 & 30 MG	3045406000B745	Brand
TOLVAPTAN	TOLVAPTAN TAB 15 MG	30454060000320	Generic

JYNARQUE	TOLVAPTAN TAB 15 MG	30454060000320	Brand
TOLVAPTAN	TOLVAPTAN TAB 30 MG	30454060000330	Generic
JYNARQUE	TOLVAPTAN TAB 30 MG	30454060000330	Brand

1 - Diagnosis of autosomal dominant polycystic kidney disease (ADPKD)

AND

2 - Prescribed by, or on the recommendation of, a Nephrologist or other expert in kidney disease

AND

3 - Age greater than or equal to 18 years

AND

4 - Estimated glomerular filtration rate ≥ 25 ml/min

Product Name: Brand Jynarque, Jynarque PAK, generic tolvaptan			
Approval Length 12 month(s)			
Therapy Stage	Reauthorization		
Guideline Type Prior Authorization			

Product Name	Generic Name	GPI	Brand/Generic
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 15 MG	3045406000B710	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 30 & 15 MG	3045406000B720	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 45 & 15 MG	3045406000B725	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 60 & 30 MG	3045406000B735	Brand
JYNARQUE	TOLVAPTAN TAB THERAPY PACK 90 & 30 MG	3045406000B745	Brand

TOLVAPTAN	TOLVAPTAN TAB 15 MG	30454060000320	Generic
JYNARQUE	TOLVAPTAN TAB 15 MG	30454060000320	Brand
TOLVAPTAN	TOLVAPTAN TAB 30 MG	30454060000330	Generic
JYNARQUE	TOLVAPTAN TAB 30 MG	30454060000330	Brand

1 - Submission of medical records (e.g., chart notes) documenting that current laboratory values for liver and kidneys remain within acceptable treatment ranges

Date	Notes
3/18/2024	Updated product name

Kerendia (finerenone)
The behavioring arrows to engagine. This long have been consider an older that it is probe the promotion of consideration of the consid

Guideline ID	GL-129742
Guideline Name	Kerendia (finerenone)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Kerendia				
Approval Length		12 month(s)		
Guideline Type Prior Authorization - IL and MN Plans Only				
Product Name	Generic Na	me	GPI	Brand/Generic
KERENDIA	FINERENONE TAB 10 MG 30354030000310 Brand		Brand	
KERENDIA	FINERENONE TAB 20 MG 30354030000320 Brand		Brand	

Approval Criteria

1 - Diagnosis of chronic kidney disease (CKD) associated with type 2 diabetes

AND

- 2 Diagnosis is confirmed by one of the following:
 - Urine albumin creatinine ratio (UACR) between 30 mg/g and 300 mg/g and estimated glomerular filtration rate (eGFR) of 25 to 60 mL/min
 - UACR > 300 mg/g and eGFR of 25 to 75 mL/min

AND

3 - Serum potassium level ≤ 5 mEq/L

AND

- **4** Trial and failure of a maximally tolerated dose, contraindication or intolerance to both of the following:
 - Ace-inhibitor (ACE-I) or angiotensin receptor blocker (ARB)
 - Sodium-glucose cotransporter-2 (SGLT2) drug that has an indication for CKD benefit (e.g., dapagliflozin, canagliflozin, empagliflozin)

AND

5 - UACR remains above 30 mg/g despite use of ACE-I/ARB and SGLT2 (unless contraindicated or not tolerated)

Product Name: Kerendia			
Approval Length 12/31/2039			
Therapy Stage	Initial Authorization		
Guideline Type	Prior Authorization - All plans excep	t IL and MN	
Product Generic Na	ame	GPI	Brand/Generic

Product Name	Generic Name	GPI	Brand/Generic
KERENDIA	FINERENONE TAB 10 MG	30354030000310	Brand
KERENDIA	FINERENONE TAB 20 MG	30354030000320	Brand

1 - Diagnosis of chronic kidney disease (CKD) associated with type 2 diabetes

AND

- **2** Diagnosis is confirmed by one of the following:
 - Urine albumin creatinine ratio (UACR) between 30 mg/g and 300 mg/g and estimated glomerular filtration rate (eGFR) of 25 to 60 mL/min
 - UACR > 300 mg/g and eGFR of 25 to 75 mL/min

AND

3 - Serum potassium level ≤ 5 mEq/L

AND

- **4** Trial and failure of a maximally tolerated dose, contraindication or intolerance to both of the following:
 - Ace-inhibitor (ACE-I) or angiotensin receptor blocker (ARB)
 - Sodium-glucose cotransporter-2 (SGLT2) drug that has an indication for CKD benefit (e.g., dapagliflozin, canagliflozin, empagliflozin)

AND

5 - UACR remains above 30 mg/g despite use of ACE-I/ARB and SGLT2 (unless contraindicated or not tolerated)

Product Name: Kerendia	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN plans

Product Name	Generic Name	GPI	Brand/Generic
KERENDIA	FINERENONE TAB 10 MG	30354030000310	Brand
KERENDIA	FINERENONE TAB 20 MG	30354030000320	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
10/31/2023	2024 New Implementation

Ketorolac Injection
The bandway warm budgue. The lost bands most, waste, a ballet likely later bit years to consider an auto-

Guideline ID	GL-132775
Guideline Name	Ketorolac Injection
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Ketorlac Injection	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Quantity Limit - Applies to IL and MN plans only	

Product Name	Generic Name	GPI	Brand/Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE INJ 15 MG/ML	66100037102015	Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE IM INJ 60 MG/2ML (30 MG/ML)	66100037102071	Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE INJ 30 MG/ML	66100037102030	Generic

Approval Criteria

- 1 Submission of medical records (e.g., chart notes) documenting that the member does not have one of the following:

 - reduced kidney function history of gastrointestinal ulcers/bleeds

Product Name: Ketorla	Product Name: Ketorlac Injection	
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type Quantity Limit - Applies to IL and MN plans only		

Product Name	Generic Name	GPI	Brand/Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE INJ 15 MG/ML	66100037102015	Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE IM INJ 60 MG/2ML (30 MG/ML)	66100037102071	Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE INJ 30 MG/ML	66100037102030	Generic

1 - Submission of medical records (e.g., chart notes) from the past 12 months that the member is having a positive response to therapy

Product Name: Ketorlac Injection	
Approval Length 12/31/2039	
Guideline Type Quantity Limit - Applies to all plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE INJ 15 MG/ML	66100037102015	Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE IM INJ 60 MG/2ML (30 MG/ML)	66100037102071	Generic
KETOROLAC TROMETHAMINE	KETOROLAC TROMETHAMINE INJ 30 MG/ML	66100037102030	Generic

Approval Criteria

- 1 Submission of medical records (e.g., chart notes) documenting that the member does not have one of the following:

 - reduced kidney function history of gastrointestinal ulcers/bleeds

Date	Notes
10/31/2023	New Program

K	Keveyis (Dichlorphenamide)		
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Guideline ID	GL-131972
Guideline Name	Keveyis (Dichlorphenamide)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Dichlorphenamide	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
DICHLORPHENAMIDE	DICHLORPHENAMIDE TAB 50 MG	37100020000305	Generic

Approval Criteria

1 - Diagnosis of primary hyperkalemic periodic paralysis, primary hypokalemic periodic paralysis, or related variants

AND

2 - Age greater than or equal to 18

Product Name: Generic Dichlorphenamide			
Approval Length	12 month(s)		
Therapy Stage	Reauthorization		
Guideline Type	Prior Authorization-IL and MN Plans Only		
Product Name	Generic Name	GPI	Brand/Generic
DICHLORPHENAMIDE	DICHLORPHENAMIDE TAB 50 MG	37100020000305	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Ge	neric Dichlorphenamide		
Approval Length	12/31/2039		
Guideline Type	Prior Authorization-All p	lans except IL and MN	
Product Namo	Conorio Namo	GBI	Brand/Ganaria

Product Name	Generic Name	GPI	Brand/Generic
DICHLORPHENAMIDE	DICHLORPHENAMIDE TAB 50 MG	37100020000305	Generic

Approval Criteria

 ${\bf 1}$ - Diagnosis of primary hyperkalemic periodic paralysis, primary hypokalemic periodic paralysis, or related variants

AND

2 - Age greater than or equal to 18

Date	Notes
10/31/2023	New program

	Kineret (anakinra)	
[The Standard Group control to High, and "Stand Stand Standard," or desired Standard	_

Guideline ID	GL-145449
Guideline Name	Kineret (anakinra)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	1/16/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Kineret		
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis (RA), Juvenile Idiopathic Arthritis (JIA)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization – All Plans except IL and MN Plans	

KINERET ANAKINRA SUBCUTANEOUS SOLN PREFILLED 6626001000E52	
SYRINGE 100 MG/0.67ML 002000 1000E32	0 Brand

Approval Criteria
1 - Diagnosis of one of the following:
 Moderate to severely active rheumatoid arthritis (RA) Juvenile idiopathic arthritis (JIA)
AND
2 - Prescribed by or in consultation with a rheumatologist
AND
3 - One of the following:
3.1 All of the following:
3.1.1 Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 Methotrexate (MTX)* Leflunomide Hydroxychloroquine Sulfasalazine
AND
3.1.2 Both of the following:
3.1.2.1 Trial and failure, contraindication, or intolerance to TWO of the following:
 Certolizumab Etanercept Adalimumab (biosimilars or Humira) Upadacitinib Golimumab Tofacitinib/ER
AND

3.1.2.2 Trial and failure, contraindication, or intolerance to BOTH of the following:

- Tocilizumab
- Abatacept

OR

3.2 Continuation of prior therapy with anakinra, verified by paid claims or medical records (e.g. chart notes)

Notes	*Absolute contraindications to methotrexate are pregnancy, nursing, a
	Icoholism, alcoholic liver disease or other chronic liver disease, immun
	odeficiency syndromes, bone marrow hyperplasia, leukopenia, thromb
	ocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Kineret		
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis (RA), Juvenile Idiopathic Arthritis (JIA)	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
KINERET	ANAKINRA SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	6626001000E520	Brand

Approval Criteria

- 1 Diagnosis of one of the following:
 - Moderate to severely active rheumatoid arthritis (RA)
 - Juvenile idiopathic arthritis (JIA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - One of the following: **3.1** All of the following: 3.1.1 Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following: Methotrexate (MTX)* Leflunomide • Hydroxychloroquine Sulfasalazine **AND 3.1.2** Both of the following: 3.1.2.1 Trial and failure, contraindication, or intolerance to TWO of the following: Certolizumab Etanercept Adalimumab (biosimilars or Humira) Upadacitinib Golimumab Tofacitinib/ER **AND** 3.1.2.2 Trial and failure, contraindication, or intolerance to BOTH of the following: **Tocilizumab** Abatacept OR

3.2 Continuation of prior therapy with anakinra, verified by paid claims or medical records (e.g. chart notes)

Notes	*Absolute contraindications to methotrexate are pregnancy, nursing, a
	Icoholism, alcoholic liver disease or other chronic liver disease, immun
	odeficiency syndromes, bone marrow hyperplasia, leukopenia, thromb
	ocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Kineret			
Diagnosis	Cryopyrin Associated Periodic Syndromes (CAPS)		
Approval Length	Approval Length 12/31/2039		
Guideline Type Prior Authorization – All Plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
KINERET	ANAKINRA SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	6626001000E520	Brand

1 - Diagnosis of cryopyrin associated periodic syndromes (CAPS) including Muckle Wells, neonatal-onset multisystemic inflammatory disorder, familial cold autoinflammatory syndrome, or other periodic syndromes

AND

2 - Prescribed by or in consultation with a rheumatologist

Product Name: Kineret		
Diagnosis	Cryopyrin Associated Periodic Syndromes (CAPS)	
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
KINERET	ANAKINRA SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	6626001000E520	Brand

Approval Criteria

1 - Diagnosis of cryopyrin associated periodic syndromes (CAPS) including Muckle Wells, neonatal-onset multisystemic inflammatory disorder, familial cold autoinflammatory syndrome, or other periodic syndromes

AND

2 - Prescribed by or in consultation with a rheumatologist

Product Name: Kineret		
Diagnosis	Systemic Juvenile Arthritis, Adult-Onset Still's Disease	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
KINERET	ANAKINRA SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	6626001000E520	Brand

Approval Criteria

1 - Diagnosis of systemic juvenile arthritis or adult-onset Still's disease

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to ONE of the following for 3 months:
 - corticosteroids
 - methotrexate
 - nonsteroidal anti-inflammatory drugs (NSAIDs)

OR

3.2 Continuation of prior therapy with anakinra, verified by paid claims or medical records (e.g. chart notes)

Product Name: Kineret		
Diagnosis	Systemic Juvenile Arthritis, Adult-Onset Still's Disease	
Approval Length	12 month(s)	
Guideline Type Prior Authorization – IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
KINERET	ANAKINRA SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	6626001000E520	Brand

Approval Criteria

1 - Diagnosis of systemic juvenile arthritis or adult-onset Still's disease

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to ONE of the following for 3 months:
 - corticosteroids
 - methotrexate
 - nonsteroidal anti-inflammatory drugs (NSAIDs)

OR

3.2 Continuation of prior therapy with anakinra, verified by paid claims or medical records (e.g. chart notes)

Product Name: Kineret			
Diagnosis	Cryopyrin Associated Periodic Syndromes (CAPS)		
Approval Length	12 month(s)		
Therapy Stage	Reauthorization		
Guideline Type	Prior Authorization – IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
KINERET	ANAKINRA SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/0.67ML	6626001000E520	Brand

1 - Prescriber provides clinical documentation from the previous 12 months that describes the member's response as stable disease or improvement seen on therapy

Date	Notes
4/5/2024	Guideline Update.

K	uvan	(sap	ropte	rin)		
. ~	triced image current he sliquiques. The	rik ney have been moved, sonamed, or	states. Welly that he list points the m	mentile and leaders.		

Guideline ID	GL-131589
Guideline Name	Kuvan (sapropterin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

1. Criteria

Product Name: Generic sapropterin				
Approval Length 2 month(s)				
Therapy Stage Initial Authorization				
Guideline Type		Prior Authorization - All plans except IL and MN		
Product Name	Ge	neric Name	GPI	Brand/Generic
SAPROPTERIN DIHYDROCHLORIDE		PROPTERIN DIHYDROCHLORIDE TAB MG	30908565100320	Generic

	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 100 MG	30908565103020	Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 500 MG	30908565103040	Generic

1 - Diagnosis of phenylketonuria (PKU)

AND

2 - Used in conjunction with a phenylalanine (Phe) restricted diet

AND

3 - Member is not on concurrent pegvaliase therapy

Product Name: Generic sapropterin		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE TAB 100 MG	30908565100320	Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 100 MG	30908565103020	Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 500 MG	30908565103040	Generic

Approval Criteria

1 - Diagnosis of phenylketonuria (PKU)

AND

2 - Used in conjunction with a phenylalanine (Phe) restricted diet

AND

3 - Member is not on concurrent pegvaliase therapy

Product Name: Generic sapropterin		
Diagnosis	After 2 month initial fill	
Approval Length	12 month(s)	
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE TAB 100 MG	30908565100320	Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 100 MG	30908565103020	Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 500 MG	30908565103040	Generic

Approval Criteria

 ${\bf 1}$ - Clinical documentation of a 30% or more reduction in Phe levels from baseline on sapropterin treatment

AND

2 - Used in conjunction with a phenylalanine (Phe) restricted diet

AND

3 - Member will continue to have blood Phe levels measured periodically during treatment

AND

4 - Member is not on concurrent pegvaliase therapy

Product Name: Generic sapropterin	
Diagnosis	Continuation of Coverage
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - All Plans

Product Name	Generic Name	GPI	Brand/Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE TAB 100 MG	30908565100320	Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 100 MG	30908565103020	Generic
SAPROPTERIN DIHYDROCHLORIDE	SAPROPTERIN DIHYDROCHLORIDE POWDER PACKET 500 MG	30908565103040	Generic

Approval Criteria

1 - Used in conjunction with a phenylalanine (Phe) restricted diet

AND

2 - Member will continue to have blood Phe levels measured periodically during treatment

AND

3 - Member is not on concurrent pegvaliase therapy

Date	Notes
10/27/2023	2024 New Implementation

Lescol (Fluvastatin), Lescol XL (Fluvastatin)		
(3) buildings and halpes hall as before that exact a man hall belong path to make a rhado.		

Guideline ID	GL-144572
Guideline Name	Lescol (Fluvastatin), Lescol XL (Fluvastatin XR)
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Generic: Fluvastatin, Fluvastatin XR		
Approval Length 12 month(s)		
Therapy Stage	nerapy Stage Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
FLUVASTATIN SODIUM ER	FLUVASTATIN SODIUM TAB ER 24 HR 80 MG (BASE EQUIVALENT)	39400030107530	Generic
FLUVASTATIN	FLUVASTATIN SODIUM CAP 20 MG (BASE EQUIVALENT)	39400030100120	Generic
FLUVASTATIN	FLUVASTATIN SODIUM CAP 40 MG (BASE EQUIVALENT)	39400030100140	Generic

Approval Criteria	
•	ntraindication or intolerance to all generic preferred statins n, pravastatin, rosuvastatin and simvastatin)
Notes	Quartz members aged 45-75 who meet the Lescol (Fluvastatin), Lesc ol XL (Fluvastatin XR) guideline, are eligible to receive the medication at a zero-dollar copay, upon approval.

Product Name: Generic: Fluvastatin, Fluvastatin XR		
Approval Length	pproval Length 12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
FLUVASTATIN SODIUM ER	FLUVASTATIN SODIUM TAB ER 24 HR 80 MG (BASE EQUIVALENT)	39400030107530	Generic
FLUVASTATIN	FLUVASTATIN SODIUM CAP 20 MG (BASE EQUIVALENT)	39400030100120	Generic
FLUVASTATIN	FLUVASTATIN SODIUM CAP 40 MG (BASE EQUIVALENT)	39400030100140	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Notes	Quartz members aged 45-75 who meet the Lescol (Fluvastatin), Lesc
	ol XL (Fluvastatin XR) guideline, are eligible to receive the medication
	at a zero-dollar copay, upon approval.

Product Name: Generic: Fluvastatin, Fluvastatin XR					
Approval Length		12/31/2039			
Guideline Type		Prior Authorization-All plans except IL and MN			
Product Generic Name		Name	GPI	Brand/Generic	
FLUVASTATIN SODIUM ER	FLUVASTATIN SODIUM TAB ER 24 HR 80 MG (BASE EQUIVALENT)		39400030107530	Generic	
FLUVASTATIN	UVASTATIN FLUVASTATIN SODIUM CAP 20 MG (BASE EQUIVALENT)		39400030100120	Generic	

FLUVASTATIN	JVASTATIN FLUVASTATIN SODIUM CAP 40 MG (BASI EQUIVALENT)		39400030100140	Generic	
Approval Criteria					
1 - Trial and failure, contraindication or intolerance to all generic preferred statins (atorvastatin, lovastatin, pravastatin, rosuvastatin and simvastatin)					
Notes	Quartz members aged 45-75 who meet the Lescol (Fluvastatin), Lesc ol XL (Fluvastatin XR) guideline, are eligible to receive the medication at a zero-dollar copay, upon approval.				

Date	Notes
3/18/2024	Updated notes section

Leukine (Sargramostim)				
The Market integrations for fingless. The firm to be here mored, was and, or defined that produce the control that of trades.				

Guideline ID	GL-136712
Guideline Name	Leukine (Sargramostim)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Na	Product Name: Leukine			
Approval Length		12 month(s)		
Guideline Type		Prior Authorization		
Product Generic Na Name LEUKINE SARGRAMOS		me	GPI	Brand/Generic
		STIM LYOPHILIZED FOR INJ 250 MCG	82402050002120	Brand

Approval Criteria

- 1 One of the following:
 - **1.1** Trial and failure, contraindication, or intolerance to tbo-filgrastim (i.e. Granix)

OR

- **1.2** Both of the following:
- 1.2.1 Diagnosis if neuroblastoma

AND

1.2.2 Used in combination with naxitamab (Danyelza)

OR

1.3 Minnesota plans only: The person has stage four metastatic cancer and the requested drug is being used as supportive care for their cancer treatment.

Date	Notes
11/27/2023	Criteria updated

Leuprolide daily injection				
(2) International analysis for the first terminal counts and and both to proceed an analysis and the counts are considered as a count of the counts and the counts are counts are counts and the counts are considered as a count of the counts are counts are considered as a count of the count of the counts are considered as a count of the counts are considered as a count of the count of the counts are considered as a count of the co				

Guideline ID	GL-132743
Guideline Name	Leuprolide daily injection
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Nam	e: Leupro	ide Injection		
Approval Length		12/31/2039		
Guideline Type		Prior Authorization - All plans except MN Plans		
Product Name	Generic N	Name	GPI	Brand/Generic
LEUPROLIDE ACETATE	LEUPROLI	DE ACETATE INJ KIT 5 MG/ML	21405010106407	Generic

Approval Criteria

1 - The injections will be self-administered

AND

2 - Use is for a diagnosis other than infertility (e.g.,. prostate cancer, endometriosis, dysmenorrhea, etc.)

Product Name: Leuprolide Injection		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
LEUPROLIDE ACETATE	LEUPROLIDE ACETATE INJ KIT 5 MG/ML	21405010106407	Generic

Approval Criteria

1 - The injections will be self-administered

AND

2 - Use is for a diagnosis other than infertility (e.g.,. prostate cancer, endometriosis, dysmenorrhea, etc.)

Product Name: Leuprolide Injection		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
LEUPROLIDE ACETATE	LEUPROLIDE ACETATE INJ KIT 5 MG/ML	21405010106407	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
10/31/2023	2024 New Implementation

L	Levemir (insulin detemir)				
•	and the state of t	_			

Guideline ID	GL-129856	
Guideline Name	Levemir (insulin detemir)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Levemir	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
LEVEMIR FLEXPEN	INSULIN DETEMIR SOLN PEN-INJECTOR 100 UNIT/ML	2710400600D220	Brand
LEVEMIR	INSULIN DETEMIR INJ 100 UNIT/ML	27104006002020	Brand

Approval Criteria

1 - Both of the following:

- Member is currently pregnant
- Diagnosis of gestational diabetes

AND

- 2 Prescribed by or in consultation with one of the following:
 - Endocrinologist
 - Diabetes specialist

AND

- **3** One of the following:
- **3.1** Member cannot meet their glycemic goals despite an adequate trial of insulin isophane (NPH) including:
 - Dose escalation, unless dose increases cannot be tolerated due to nocturnal hypoglycemia or at least one severe low blood sugar event (requiring assistance from another) or would not be appropriate given the person's self-monitoring blood glucose profile
 - Submission of medical records (e.g., chart notes) documenting use of a specific adherence intervention deployed by a health care professional if nonadherence is evident

OR

3.2 Member is intolerant to insulin isophane (NPH)

Product Name: Levemir	
Approval Length 12 month(s)	
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
LEVEMIR FLEXPEN	INSULIN DETEMIR SOLN PEN-INJECTOR 100 UNIT/ML	2710400600D220	Brand

LEVEMIR	INSULIN DETEMIR INJ 100 UNIT/ML	27104006002020	Brand
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1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Levemir	
Approval Length 12/31/2039	
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
LEVEMIR FLEXPEN	INSULIN DETEMIR SOLN PEN-INJECTOR 100 UNIT/ML	2710400600D220	Brand
LEVEMIR	INSULIN DETEMIR INJ 100 UNIT/ML	27104006002020	Brand

Approval Criteria

- **1** Both of the following:
 - Member is currently pregnant
 - Diagnosis of gestational diabetes

AND

- 2 Prescribed by or in consultation with one of the following:
 - Endocrinologist
 - Diabetes specialist

AND

- 3 One of the following:
- **3.1** Member cannot meet their glycemic goals despite adequate trials of insulin isophane (NPH) including:

- Dose escalation, unless dose increases cannot be tolerated due to nocturnal hypoglycemia or at least one severe low blood sugar event (requiring assistance from another) or would not be appropriate given the person's self-monitoring blood glucose profile
- Submission of medical records (e.g., chart notes) documenting use of a specific adherence intervention deployed by a health care provider if nonadherence is evident

OR

3.2 Member is intolerant to insulin isophane (NPH)

Date	Notes
10/12/2023	2024 New Implementation

Livmarli (maralixibat)		
The State Designer of the State of States and county a State		

Guideline ID	GL-135578
Guideline Name	Livmarli (maralixibat)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Livmarli	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
LIVMARLI	MARALIXIBAT CHLORIDE ORAL SOLN 9.5 MG/ML	52350050102020	Brand

Approval Criteria

1 - Diagnosis of Alagille syndrome (ALGS)

AND

2 - Molecular genetic testing confirms mutations in the JAG1 or NOTCH2 gene

AND

- **3** One of the following:
 - Total serum bile acid greater than 3x the upper limit of normal (ULN)
 - Conjugated bilirubin greater than 1 mg/dL
 - Fat soluble vitamin deficiency otherwise unexplainable
 - Gamma-glutamyl transpeptidase (GGT) greater than 3x ULN

AND

4 - Member is experiencing moderate to severe cholestatic pruritus

AND

5 - Member has not had a liver transplant or decompensated liver disease

AND

- **6** Trial and failure, contraindication or intolerance to TWO of the following medications for pruritis:
 - Ursodeoxycholic acid (e.g., Ursodiol)
 - Antihistamines (e.g., diphenhydramine, hydroxyzine)
 - Rifampin
 - Bile acid sequestrants (e.g., Questran, Colestid, Welchol)

AND

7 - Prescribed by or in consultation with one of the following:

 Hepatologist Expert in the treatment of cholestasis 			
Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies. **For members new to plan (as evidenced by coverage effective date of less than or equal to 90 days) prescriber provides submission of me dical records (e.g., chart notes) documenting that from the previous 1 2 months, member demonstrates an improvement or stabilization in pr		

Product Name: Livmarli				
Approval Length		12 month(s)		
Therapy Stage		Reauthorization		
Guideline Type		Prior Authorization		
Product Name	Generic Na	me	GPI	Brand/Generic
LIVMARLI	MARALIXIBAT CHLORIDE ORAL SOLN 9.5 MG/ML		52350050102020	Brand

uritus and the member is tolerating therapy.

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months of therapy, member is experiencing improvement or stabilization in pruritus compared to baseline (e.g., change in member reported pruritus, change in sleeping habits due to itch) and the member is tolerating therapy (e.g., does not have chronic diarrhea requiring ongoing intravenous fluids, bile acid reduction)

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies. **For members new to plan (as evidenced by coverage effective date of less than or equal to 90 days) prescriber provides submission of me dical records (e.g., chart notes) documenting that from the previous 1 2 months, member demonstrates an improvement or stabilization in pruritus and the member is tolerating therapy.
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Date	Notes
10/30/2023	2024 New Implementation

Livtencity (maribavir)					
(g) the state of t					

Guideline ID	GL-129857
Guideline Name	Livtencity (maribavir)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Livtencity				
Approval Length		16 Week(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization		
Product Name	Generic N	ame	GPI	Brand/Generic
LIVTENCITY	ICITY MARIBAVIR TAB 200 MG		12200050000320	Brand

1 - Diagnosis of cytomegalovirus (CMV) infection based on clinical history and laboratory testing

AND

2 - History of stem cell or solid organ transplant

AND

- **3** Prescribed by or in consultation with one of the following:
 - Hematologist
 - Oncologist
 - Infectious Disease Specialist
 - Transplant Specialist

AND

4 - Submission of medical records (e.g., chart notes) documenting baseline viral load prior to initiating therapy

AND

- **5** Trial and failure, contraindication, or intolerance to one of the following:
 - Ganciclovir
 - Valganciclovir
 - Cidofovir
 - Foscarnet

*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsore d free drug program, provider samples, and/or
vouchers.

Product Name: Livtencity	
Approval Length	16 Week(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
LIVTENCITY	MARIBAVIR TAB 200 MG	12200050000320	Brand

1 - Submission of medical records (e.g., chart notes) supporting treatment response and evidence-based clinical rationale for use beyond 16 weeks of therapy

OR

2 - Members new to coverage (as evidenced by coverage effective date of less than or equal to 90 days) who are established on therapy will have coverage under their drug benefit for the remainder of the current treatment course (to a maximum of 16 weeks)

*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsore d free drug program, provider samples, and/or vouchers.
vouchers.

Date	Notes
8/21/2023	2024 New Implementation

L	Lupkynis (voclosporin)		
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Guideline ID	GL-132812
Guideline Name	Lupkynis (voclosporin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Lupkynis		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN plans only		

Product Name	Generic Name	GPI	Brand/Generic
LUPKYNIS	VOCLOSPORIN CAP 7.9 MG	99402080000120	Brand

Approval Criteria

1 - Diagnosis of biopsy-proven lupus nephritis

AND

- 2 Prescribed by or in consultation with one of the following:
 - Nephrologist
 - Rheumatologist
 - specialist in the treatment of lupus nephritis

AND

3 - Trial and failure, contraindication, or intolerance to concurrent use of mycophenolate with corticosteroids

AND

4 - Requested drug will be used in combination with mycophenolate and corticosteroids unless contraindicated or not tolerated

AND

5 - Requested drug will not be used in combination with cyclophosphamide

Product Name: Lupkynis	
Approval Length 6 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization - All plans except IL and MN	

Produ Name	et Generic Name	GPI	Brand/Generic
LUPKY	VOCLOSPORIN CAP 7.9 MG	99402080000120	Brand

Approval Criteria

1 - Diagnosis of biopsy-proven lupus nephritis

AND

- 2 Prescribed by or in consultation with one of the following:
 - Nephrologist
 - Rheumatologist
 - · specialist in the treatment of lupus nephritis

AND

3 - Trial and failure, contraindication, or intolerance to concurrent use of mycophenolate with corticosteroids

AND

4 - Requested drug will be used in combination with mycophenolate and corticosteroids unless contraindicated or not tolerated

AND

5 - Requested drug will not be used in combination with cyclophosphamide

Product Name: Lupkynis	
Approval Length 12 month(s)	
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - All plans	

Product Name	Generic Name	GPI	Brand/Generic
LUPKYNIS	VOCLOSPORIN CAP 7.9 MG	99402080000120	Brand

Approval Criteria

1 - Patient has demonstrated a positive response to therapy

Date	Notes
11/1/2023	New Program

Ν	Mucosal Protectants		
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Guideline ID	GL-137862
Guideline Name	Mucosal Protectants
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Mugard, Episil, Oramagicrx				
Approval Length 12 month(s)				
Guideline Type Prior Authorization				
Product Name	Generic	Name	GPI	Brand/Generic
MUGARD	*ORAL WOUND CARE PRODUCTS - LIQUID RINSE***		88502050000900	Brand
EPISIL	*ORAL WOUND CARE PRODUCTS - LIQUID PUMP***		88502050000950	Brand

Approval Criteria

ORAMAGICRX *ORAL WOUND CARE PRODUCTS - FOR SUSP RINSE***

Brand

88502050001900

1 - Diagnosis of grade 3 or 4 mucositis due to chemotherapy and/or radiation (i.e. oral mucositis: severe pain, interferes with oral intake, oral ulceration, difficulty with speech and swallowing)

AND

- 2 Both of the following:
- **2.1** Trial and failure or intolerance to ONE of any moisturizing salivation agents:
 - Rinses / Mouthwashes (i.e. OTC sodium/sodium bicarbonate, OTC Biotene, OTC Oasis)
 - Gel, Spray (i.e. OTC Biotene, OTC Mouthkote)

AND

2.2 Trial and failure or intolerance to ONE Muco-protectant with or without anesthetic agent [i.e., Mylanta, magic mouthwash (lidocaine/diphenhydramine/Mylanta)]

Product Name: Prothelial, Orafate, Silatrix		
Approval Length 12 month(s)		
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
PROTHELIAL	*SUCRALFATE-MALATE PASTE 10%***	88502002804410	Brand
ORAFATE	*SUCRALFATE-MALATE PASTE 10%***	88502002804410	Brand
SILATRIX	*SUCRALFATE-MALATE GEL 10%***	88502002804010	Brand

Approval Criteria

1 - Diagnosis of grade 3 or 4 mucositis due to chemotherapy and/or radiation (i.e. oral mucositis: severe pain, interferes with oral intake, oral ulceration, difficulty with speech and swallowing)

AND

- 2 Both of the following:
- **2.1** Trial and failure or intolerance to ONE of any moisturizing salivation agents:
 - Rinses / Mouthwashes (i.e. OTC sodium/sodium bicarbonate, OTC Biotene, OTC Oasis)
 - Gel, Spray (i.e. OTC Biotene, OTC Mouthkote)

AND

2.2 Trial and failure or intolerance to ONE Muco-protectant with or without anesthetic agent [i.e., Mylanta, magic mouthwash (lidocaine/diphenhydramine/Mylanta)]

AND

3 - Trial and failure, contraindication or intolerance to ONE bioadhesive gel (i.e., Gelclair, Oramagic Rx, Mugard or Episil)

Date	Notes
12/15/2023	Update

Multiple Sclerosis	
The links at longs current to displayed. The fire way have been record, received, or debad. Verily that the links points to be convertile and deaders.	
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Guideline ID	GL-145312
Guideline Name	Multiple Sclerosis
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	7/1/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Avonex, Extavia, Generic dimethyl fumarate, Generic fingolimod, Generic glatiramer acetate, Generic teriflunomide, Glatopa, Plegridy, Rebif				
Approval Length		12/31/2039		
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans		ns		
Product Name	Gene	Generic Name GPI Brand/Generic		
DIMETHYL FUMARATE	DIMETHYL FUMARATE CAPSULE DELAYED RELEASE 120 MG		62405525006520	Generic
DIMETHYL FUMARATE		HYL FUMARATE CAPSULE DELAYED SE 240 MG	62405525006540	Generic
FINGOLIMOD	FINGOLIMOD HCL CAP 0.5 MG (BASE EQUIV)		62407025100120	Generic
GLATOPA		RAMER ACETATE SOLN PREFILLED GE 20 MG/ML	6240003010E520	Generic

GLATIRAMER ACETATE	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 20 MG/ML	6240003010E520	Generic
GLATOPA	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 40 MG/ML	6240003010E540	Generic
GLATIRAMER ACETATE	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 40 MG/ML	6240003010E540	Generic
AVONEX PEN	INTERFERON BETA-1A IM AUTO-INJECTOR KIT 30 MCG/0.5ML	6240306045F530	Brand
AVONEX	INTERFERON BETA-1A IM PREFILLED SYRINGE KIT 30 MCG/0.5ML	6240306045F830	Brand
REBIF REBIDOSE	INTERFERON BETA-1A SOLN AUTO-INJ 22 MCG/0.5ML	6240306045D520	Brand
REBIF REBIDOSE	INTERFERON BETA-1A SOLN AUTO-INJ 44 MCG/0.5ML	6240306045D540	Brand
REBIF REBIDOSE TITRATION PACK	INTERFERON BETA-1A AUTO-INJ 6X8.8 MCG/0.2ML & 6X22 MCG/0.5ML	6240306045D560	Brand
REBIF	INTERFERON BETA-1A SOLN PREF SYR 22 MCG/0.5ML	6240306045E520	Brand
REBIF	INTERFERON BETA-1A SOLN PREF SYR 44 MCG/0.5ML	6240306045E540	Brand
REBIF TITRATION PACK	INTERFERON BETA-1A PREF SYR 6X8.8 MCG/0.2ML & 6X22 MCG/0.5ML	6240306045E560	Brand
EXTAVIA	INTERFERON BETA-1B FOR INJ KIT 0.3 MG	62403060506420	Brand
PLEGRIDY	PEGINTERFERON BETA-1A SOLN PEN- INJECTOR 125 MCG/0.5ML	6240307530D220	Brand
PLEGRIDY STARTER PACK	PEGINTERFERON BETA-1A SOLN PEN-INJ 63 & 94 MCG/0.5ML PACK	6240307530D250	Brand
PLEGRIDY	PEGINTERFERON BETA-1A SOLN PREFILLED SYRINGE 125 MCG/0.5ML	6240307530E520	Brand
PLEGRIDY	PEGINTERFERON BETA-1A IM SOLN PREFILLED SYR 125 MCG/0.5ML	6240307530E521	Brand
PLEGRIDY STARTER PACK	PEGINTERFERON BETA-1A SOLN PREF SYR 63 & 94 MCG/0.5ML PACK	6240307530E550	Brand
TERIFLUNOMIDE	TERIFLUNOMIDE TAB 7 MG	62404070000320	Generic
TERIFLUNOMIDE	TERIFLUNOMIDE TAB 14 MG	62404070000330	Generic
DIMETHYL FUM CAP STARTER	DIMETHYL FUM CAP STARTER	6240552500B320	Generic
DIMETHYL FUMARATE STARTERPACK	DIMETHYL FUMARATE CAPSULE DR STARTER PACK 120 MG & 240 MG	6240552500B320	Generic

- 1 One of the following:
- **1.1** Diagnosis of one of the following relapsing forms of multiple sclerosis:
 - Relapsing-Remitting
 - Active secondary progressive
 - Relapsing-progressive

OR

1.2 Diagnosis of Clinically Isolated Syndrome (CIS) with a high probability of developing Clinically Definite MS (CDMS) (i.e. greater than or equal to 3 T2 white matter lesions or greater than or equal to 2 GdE lesions on MRI)

AND

2 - Prescribed by or in consultation with a neurologist or other expert in the treatment of multiple sclerosis

Product Name: Avonex, Extavia, Generic dimethyl fumarate, Generic fingolimod, Generic glatiramer acetate, Generic teriflunomide, Glatopa, Plegridy, Rebif		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans*	

Product Name	Generic Name	GPI	Brand/Generic
DIMETHYL FUMARATE	DIMETHYL FUMARATE CAPSULE DELAYED RELEASE 120 MG	62405525006520	Generic
DIMETHYL FUMARATE	DIMETHYL FUMARATE CAPSULE DELAYED RELEASE 240 MG	62405525006540	Generic
FINGOLIMOD	FINGOLIMOD HCL CAP 0.5 MG (BASE EQUIV)	62407025100120	Generic
GLATOPA	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 20 MG/ML	6240003010E520	Generic
GLATIRAMER ACETATE	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 20 MG/ML	6240003010E520	Generic
GLATOPA	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 40 MG/ML	6240003010E540	Generic
GLATIRAMER ACETATE	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 40 MG/ML	6240003010E540	Generic

AVONEX PEN	INTERFERON BETA-1A IM AUTO-INJECTOR KIT 30 MCG/0.5ML	6240306045F530	Brand
AVONEX	INTERFERON BETA-1A IM PREFILLED SYRINGE KIT 30 MCG/0.5ML	6240306045F830	Brand
REBIF REBIDOSE	INTERFERON BETA-1A SOLN AUTO-INJ 22 MCG/0.5ML	6240306045D520	Brand
REBIF REBIDOSE	INTERFERON BETA-1A SOLN AUTO-INJ 44 MCG/0.5ML	6240306045D540	Brand
REBIF REBIDOSE TITRATION PACK	INTERFERON BETA-1A AUTO-INJ 6X8.8 MCG/0.2ML & 6X22 MCG/0.5ML	6240306045D560	Brand
REBIF	INTERFERON BETA-1A SOLN PREF SYR 22 MCG/0.5ML	6240306045E520	Brand
REBIF	INTERFERON BETA-1A SOLN PREF SYR 44 MCG/0.5ML	6240306045E540	Brand
REBIF TITRATION PACK	INTERFERON BETA-1A PREF SYR 6X8.8 MCG/0.2ML & 6X22 MCG/0.5ML	6240306045E560	Brand
EXTAVIA	INTERFERON BETA-1B FOR INJ KIT 0.3 MG	62403060506420	Brand
PLEGRIDY	PEGINTERFERON BETA-1A SOLN PEN- INJECTOR 125 MCG/0.5ML	6240307530D220	Brand
PLEGRIDY STARTER PACK	PEGINTERFERON BETA-1A SOLN PEN-INJ 63 & 94 MCG/0.5ML PACK	6240307530D250	Brand
PLEGRIDY	PEGINTERFERON BETA-1A SOLN PREFILLED SYRINGE 125 MCG/0.5ML	6240307530E520	Brand
PLEGRIDY	PEGINTERFERON BETA-1A IM SOLN PREFILLED SYR 125 MCG/0.5ML	6240307530E521	Brand
PLEGRIDY STARTER PACK	PEGINTERFERON BETA-1A SOLN PREF SYR 63 & 94 MCG/0.5ML PACK	6240307530E550	Brand
TERIFLUNOMIDE	TERIFLUNOMIDE TAB 7 MG	62404070000320	Generic
TERIFLUNOMIDE	TERIFLUNOMIDE TAB 14 MG	62404070000330	Generic
DIMETHYL FUM CAP STARTER	DIMETHYL FUM CAP STARTER	6240552500B320	Generic
DIMETHYL FUMARATE STARTERPACK	DIMETHYL FUMARATE CAPSULE DR STARTER PACK 120 MG & 240 MG	6240552500B320	Generic

- **1** One of the following:
 - **1.1** Diagnosis of one of the following relapsing forms of multiple sclerosis:

- Relapsing-Remitting
- Active secondary progressive
- Relapsing-progressive

OR

1.2 Diagnosis of Clinically Isolated Syndrome (CIS) with a high probability of developing Clinically Definite MS (CDMS) (i.e. greater than or equal to 3 T2 white matter lesions or greater than or equal to 2 GdE lesions on MRI)

AND

2 - Prescribed by or in consultation with a neurologist or other expert in the treatment of multiple sclerosis

Product Name: Avonex, Extavia, Generic dimethyl fumarate, Generic fingolimod, Generic glatiramer acetate, Generic teriflunomide, Glatopa, Plegridy, Rebif		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans*	

Product Name	Generic Name	GPI	Brand/Generic
DIMETHYL FUMARATE	DIMETHYL FUMARATE CAPSULE DELAYED RELEASE 120 MG	62405525006520	Generic
DIMETHYL FUMARATE	DIMETHYL FUMARATE CAPSULE DELAYED RELEASE 240 MG	62405525006540	Generic
FINGOLIMOD	FINGOLIMOD HCL CAP 0.5 MG (BASE EQUIV)	62407025100120	Generic
GLATOPA	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 20 MG/ML	6240003010E520	Generic
GLATIRAMER ACETATE	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 20 MG/ML	6240003010E520	Generic
GLATOPA	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 40 MG/ML	6240003010E540	Generic
GLATIRAMER ACETATE	GLATIRAMER ACETATE SOLN PREFILLED SYRINGE 40 MG/ML	6240003010E540	Generic
AVONEX PEN	INTERFERON BETA-1A IM AUTO-INJECTOR KIT 30 MCG/0.5ML	6240306045F530	Brand
AVONEX	INTERFERON BETA-1A IM PREFILLED SYRINGE KIT 30 MCG/0.5ML	6240306045F830	Brand

REBIF REBIDOSE	INTERFERON BETA-1A SOLN AUTO-INJ 22 MCG/0.5ML	6240306045D520	Brand
REBIF REBIDOSE	INTERFERON BETA-1A SOLN AUTO-INJ 44 MCG/0.5ML	6240306045D540	Brand
REBIF REBIDOSE TITRATION PACK	INTERFERON BETA-1A AUTO-INJ 6X8.8 MCG/0.2ML & 6X22 MCG/0.5ML	6240306045D560	Brand
REBIF	INTERFERON BETA-1A SOLN PREF SYR 22 MCG/0.5ML	6240306045E520	Brand
REBIF	INTERFERON BETA-1A SOLN PREF SYR 44 MCG/0.5ML	6240306045E540	Brand
REBIF TITRATION PACK	INTERFERON BETA-1A PREF SYR 6X8.8 MCG/0.2ML & 6X22 MCG/0.5ML	6240306045E560	Brand
EXTAVIA	INTERFERON BETA-1B FOR INJ KIT 0.3 MG	62403060506420	Brand
PLEGRIDY	PEGINTERFERON BETA-1A SOLN PEN- INJECTOR 125 MCG/0.5ML	6240307530D220	Brand
PLEGRIDY STARTER PACK	PEGINTERFERON BETA-1A SOLN PEN-INJ 63 & 94 MCG/0.5ML PACK	6240307530D250	Brand
PLEGRIDY	PEGINTERFERON BETA-1A SOLN PREFILLED SYRINGE 125 MCG/0.5ML	6240307530E520	Brand
PLEGRIDY	PEGINTERFERON BETA-1A IM SOLN PREFILLED SYR 125 MCG/0.5ML	6240307530E521	Brand
PLEGRIDY STARTER PACK	PEGINTERFERON BETA-1A SOLN PREF SYR 63 & 94 MCG/0.5ML PACK	6240307530E550	Brand
TERIFLUNOMIDE	TERIFLUNOMIDE TAB 7 MG	62404070000320	Generic
TERIFLUNOMIDE	TERIFLUNOMIDE TAB 14 MG	62404070000330	Generic
DIMETHYL FUM CAP STARTER	DIMETHYL FUM CAP STARTER	6240552500B320	Generic
DIMETHYL FUMARATE STARTERPACK	DIMETHYL FUMARATE CAPSULE DR STARTER PACK 120 MG & 240 MG	6240552500B320	Generic

- **1** Submission of medical records (e.g., chart notes) by the treating neurologist documenting that within the past 12 months member has BOTH of the following:
 - Relapsing form of multiple sclerosis or Clinically Isolated Syndrome (CIS) Member is established on therapy

Product Name: Kesimpta, Mavenclad		
Approval Length	12/31/2039*	
Guideline Type	Prior Authorization - ALL Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
KESIMPTA	OFATUMUMAB SOLN AUTO-INJECTOR 20 MG/0.4ML	6240506500D520	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (4 TABS)	6240101500B718	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (5 TABS)	6240101500B722	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (6 TABS)	6240101500B726	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (7 TABS)	6240101500B732	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (8 TABS)	6240101500B736	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (9 TABS)	6240101500B740	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (10 TABS)	6240101500B744	Brand

- 1 One of the following:
- **1.1** Diagnosis of one of the following relapsing forms of multiple sclerosis:
 - Relapsing-Remitting
 - Active secondary progressive
 - Relapsing-progressive

OR

1.2 Diagnosis of Clinically Isolated Syndrome (CIS) with a high probability of developing Clinically Definite MS (CDMS) (i.e. greater than or equal to 3 T2 white matter lesions or greater than or equal to 2 GdE lesions on MRI)

AND

- 2 One of the following:
 - **2.1** One of the following:

- **2.1.1** Trial and failure (i.e., acute relapse or new lesion formation) while on one of the following:
 - dimethyl fumarate
 - fingolimod

OR

- **2.1.2** Contraindication, intolerance, or the inability to take BOTH of the following:
 - dimethyl fumarate
 - fingolimod

OR

2.2 Continuation of prior therapy with the requested disease modifying drug, verified by paid claims or medical records (e.g. chart notes)

AND

3 - Prescribed by or in consultation with a neurologist or other expert in the treatment of multiple sclerosis

Notes	*For new starts to ofatumumab therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve with a MDD of 0.05 (1.2ml every 28 days) for 30 da
	ys
	Second PA: Approve through 12/31/2039

Product Name: Kesimpta, Mavenclad		
Approval Length	12 Month(s)*	
Guideline Type	Prior Authorization - IL and MN Plans*	

Product Name	Generic Name	GPI	Brand/Generic
KESIMPTA	OFATUMUMAB SOLN AUTO-INJECTOR 20 MG/0.4ML	6240506500D520	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (4 TABS)	6240101500B718	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (5 TABS)	6240101500B722	Brand

MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (6 TABS)	6240101500B726	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (7 TABS)	6240101500B732	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (8 TABS)	6240101500B736	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (9 TABS)	6240101500B740	Brand
MAVENCLAD	CLADRIBINE TAB THERAPY PACK 10 MG (10 TABS)	6240101500B744	Brand

- 1 One of the following:
- **1.1** Diagnosis of one of the following relapsing forms of multiple sclerosis:
 - Relapsing-Remitting
 - Active secondary progressive
 - Relapsing-progressive

OR

1.2 Diagnosis of Clinically Isolated Syndrome (CIS) with a high probability of developing Clinically Definite MS (CDMS) (i.e. greater than or equal to 3 T2 white matter lesions or greater than or equal to 2 GdE lesions on MRI)

AND

- 2 One of the following:
 - **2.1** One of the following:
- **2.1.1** Trial and failure (i.e., acute relapse or new lesion formation) while on one of the following:
 - dimethyl fumarate
 - fingolimod

OR

2.1.2 Contraindication, intolerance, or the inability to take BOTH of the following:

- dimethyl fumarate
- fingolimod

OR

2.2 Continuation of prior therapy with the requested disease modifying drug, verified by paid claims or medical records (e.g. chart notes)

AND

3 - Prescribed by or in consultation with a neurologist or other expert in the treatment of multiple sclerosis

For new starts to ofatumumab therapy: Enter 2 PAs as follows with the same start date: First PA: Approve with a MDD of 0.05 (1.2ml every 28 days) for 30 days
Second PA: Approve for 12 months*

Date	Notes
4/15/2024	Update COT language

Myalept (Metreleptin)		
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Guideline ID	GL-129645
Guideline Name	Myalept (Metreleptin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2023
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1. Criteria

Product Name: Myalept	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic	
MYALEPT	METRELEPTIN FOR SUBCUTANEOUS INJ 11.3 MG	30906050002120	Brand	

Approval Criteria

1 - Diagnosis of congenital or acquired generalized lipodystrophy

AND

2 - Have experienced metabolic changes (e.g. increased triglycerides or fasting blood glucoses) despite an adequate trial of dietary modification

AND

3 - Failure, intolerance, or contraindication to metformin

AND

4 - Failure, intolerance, or contraindication to at least one statin medication (e.g. atorvastatin, rosuvastatin)

Product Name: Myalept	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
MYALEPT	METRELEPTIN FOR SUBCUTANEOUS INJ 11.3 MG	30906050002120	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Na	Product Name: Myalept			
Approval L	ength	12/31/2039		
Guideline Type		Prior Authorization - All plans except IL and MN		
Product Generic Na Name		me	GPI	Brand/Generic

MYALEPT	METRELEPTIN FOR SUBCUTANEOUS INJ 11.3 MG	30906050002120	Brand
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1 - Diagnosis of congenital or acquired generalized lipodystrophy

AND

2 - Have experienced metabolic changes (e.g. increased triglycerides or fasting blood glucoses) despite an adequate trial of dietary modification

AND

3 - Failure, intolerance, or contraindication to metformin

AND

4 - Failure, intolerance, or contraindication to at least one statin medication (e.g. atorvastatin, rosuvastatin)

Date	Notes
10/6/2023	New program

Myrbetriq (mirabegron)	
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Guideline ID	GL-127843
Guideline Name	Myrbetriq (mirabegron)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Myrbetriq		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
MYRBETRIQ	MIRABEGRON GRANULES FOR ORAL EXTENDED RELEASE SUSP 8 MG/ML	0+20000000220	
MYRBETRIQ	MIRABEGRON TAB ER 24 HR 25 MG	54200050007520	Brand
MYRBETRIQ	MIRABEGRON TAB ER 24 HR 50 MG	54200050007530	Brand

Approval Criteria

- **1** Trial and failure to one of the following:
 - trospium
 - oxybutynin
 - solifenacin
 - tolterodine
 - darifenacin
 - fesoterodine

Product Name: Myrbetriq		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
MYRBETRIQ	MIRABEGRON GRANULES FOR ORAL EXTENDED RELEASE SUSP 8 MG/ML	5420005000G220	Brand
MYRBETRIQ	MIRABEGRON TAB ER 24 HR 25 MG	54200050007520	Brand
MYRBETRIQ	MIRABEGRON TAB ER 24 HR 50 MG	54200050007530	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Myrbetriq			
Approval Length	Approval Length 12/31/2039		
Guideline Type Step Therapy - All Plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
	MIRABEGRON GRANULES FOR ORAL EXTENDED RELEASE SUSP 8 MG/ML	5420005000G220	Brand
MYRBETRIQ	MIRABEGRON TAB ER 24 HR 25 MG	54200050007520	Brand
MYRBETRIQ	MIRABEGRON TAB ER 24 HR 50 MG	54200050007530	Brand

- **1** Trial and failure to one of the following:

 - trospium oxybutynin solifenacin

 - tolterodine
 - darifenacin
 - fesoterodine

Date	Notes
8/25/2023	New Program

New Indication Administrative Guide					
S histories and the same and th					

Guideline ID	GL-135282	
Guideline Name	New Indication Administrative Guideline	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

1.1 Both of the following:

Diagnosis		Drugs with a prior authorization requirement for which a guideline is unavailable, OR new FDA-approved indications which are not addressed in the existing drug-specific prior authorization guideline		
Approval Length		12 month(s)		
Guideline Type		Administrative		
Product Name Gene		ric Name	GPI	Brand/Generic
Approval Criteria				
1 - One of the following:				

1.1.1 Prescribed medication is being used for a Food and Drug Administration (FDA)-approved indication

AND

- **1.1.2** Both of the following:
- **1.1.2.1** All components of the FDA approved indication are met (e.g., concomitant use, previous therapy requirements, age limitations, testing requirements, etc.)

AND

1.1.2.2 Prescribed medication will be used at a dose which is within FDA recommendations

OR

1.2 Meets the off-label administrative guideline criteria

AND

2 - (For nonpreferred medications only) Trial and failure or intolerance, or contraindication to at least 1 preferred alternative for the same indication if available

Date	Notes
11/27/2023	New Program

Non-formulary Exceptions Administrative Guidel							
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Guideline ID	GL-145455		
Guideline Name	Non-formulary Exceptions Administrative Guideline		
Formulary	Quartz		

Guideline Note:

Effective Date:	5/1/2024
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1. Criteria

Product Name: Non-formulary drugs					
Approval Length		12 month(s)			
Guideline Type		Administrative			
Product Name	Gene	ric Name	GPI	Brand/Generic	

Approval Criteria

- 1 Both of the following:
- **1.1** One of the following:
- **1.1.1** Provider attests that it is medically necessary for the individual to receive that specific contraceptive

OR

- **1.1.2** Both of the following:
- **1.1.2.1** One of the following:
- **1.1.2.1.1** Paid claims or submission of medical records (e.g. chart notes) show there have been adequate trials of ALL appropriate therapeutic alternatives and there was (a) inadequate clinical response, (b) inappropriate clinical response, (c) intolerance or (d) allergy to those medications

OR

1.1.2.1.2 An exception to the formulary may be considered when ALL appropriate therapeutic alternatives have not been tried and there is submission of medical records (e.g. chart notes) that documents that ALL appropriate therapeutic alternatives will be (a) ineffective, (b) less effective or (c) will result in adverse effects

OR

- **1.1.2.1.3** An exception to the formulary may be considered when it is a situation that it is not clinically appropriate to have adequate trials of ALL therapeutic alternatives, such as the individual has complex medical conditions, would be subject to prolonged pain, or there is a risk of severe or significant adverse medical outcomes if there is significant delay in treating the condition AND one of the following were tried:
 - At least four formulary alternatives in the same drug class as the requested medication (if the formulary includes an alternative dose form or salt of the requested nonformulary drug, one of the alternatives MUST be the formulary formulation of the requested drug (e.g. bupropion SR or XL must be an alternative for Forfivo XL))
 - If there are not four formulary alternatives in the same drug class, at least four formulary alternatives from three different drug classes (if available) when it is appropriate under the standards of acceptable medical practice for the treatment of the diagnosis to trial medications with different mechanisms of action
 - No formulary alternative is appropriate to treat the patient's condition based on mechanism of action or lack of data to support use based on standards outlined in the off-label use administrative guideline

AND

1.1.2.2 When there are prior authorization criteria for the drug class or therapeutic alternatives, an exception to the formulary should take into consideration those criteria and should not be less stringent for the non-formulary drug.

AND

- **1.2** One of the following:
- 1.2.1 Requested drug is FDA-approved for the condition being treated

OR

1.2.2 If requested for an off-label indication, the off-label guideline approval criteria have been met.

OR

2 - For Illinois Plans only: Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on nonpreferred therapy, will have coverage under their drug benefit with submission of medical records (e.g., chart notes) documenting symptom improvement or disease stability

Product Name: Non-formulary drugs				
Approval Length		3 month(s)		
Guideline Type		Administrative		
Product Name Generic Name GPI Brand/Ger		Brand/Generic		

Approval Criteria

- **1** All of the following:
- **1.1** Requested non-formulary drug is a brand with a generic available (MSC-O)

AND

1.2 The generic formulation (MSC-Y) of the requested drug is on a covered tier on the formulary

AND

1.3 The generic formulation (MSC-Y) of the requested drug is on a nationwide shortage verified by the Food and Drug Administration (FDA) website

Date	Notes
4/5/2024	Guideline Update.

N	on-Prefe	erred	Iopical	Steroid	S	
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Guideline ID	GL-145457
Guideline Name	Non-Preferred Topical Steroids
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	6/15/2016
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: AMCINONIDE OINT 0.1%, SERNIVO, IMPOYZ, CLOCORTOLONE PIVALATE CREAM 0.1%, DESONIDE CREAM 0.05%, VERDESO, DESOXIMETASONE SPRAY 0.25%, DESOXIMETASONE CREAM 0.05%, DESOXIMETASONE CREAM 0.25% DESOXIMETASONE GEL 0.05%, DESOXIMETASONE OINT 0.05%, DESOXIMETASONE OINT 0.25%, DIFLORASONE DIACETATE CREAM 0.05%, DIFLORASONE DIACETATE OINT 0.05%, APEXICON E, FLUOCINONIDE CREAM 0.1%, FLURANDRENOLIDE CREAM 0.05%, FLURANDRENOLIDE LOTION 0.05%, HALCINONIDE CREAM 0.1%, HALOG, ULTRAVATE, TEXACORT, HYDROCORTISONE BUTYRATE SOLN 0.1%, HYDROCORTISONE BUTYRATE LOTION 0.1%, HYDROCORTISONE BUTYRATE CREAM 0.1%, HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%, HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%, PANDEL, HYDROCORTISONE VALERATE OINT 0.2%, HYDROCORTISONE VALERATE CREAM 0.2%, TRIAMCINOLONE ACETONIDE AEROSOL SOLN 0.147 MG/GM, AMCINONIDE CREAM 0.1%, BRYHALI, HALOBETASOL PROPIONATE FOAM 0.05%

Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
AMCINONIDE	AMCINONIDE OINT 0.1%	90550010004205	Generic
SERNIVO	BETAMETHASONE DIPROPIONATE SPRAY EMULSION 0.05% (BASE EQUIV)	90550020001620	Brand
IMPOYZ	CLOBETASOL PROPIONATE CREAM 0.025%	90550025103703	Brand
CLOCORTOLONE PIVALATE	CLOCORTOLONE PIVALATE CREAM 0.1%	90550030103705	Generic
DESONIDE	DESONIDE CREAM 0.05%	90550035003705	Generic
VERDESO	DESONIDE FOAM 0.05%	90550035003920	Brand
DESOXIMETASONE	DESOXIMETASONE SPRAY 0.25%	90550040000910	Generic
DESOXIMETASONE	DESOXIMETASONE CREAM 0.05%	90550040003705	Generic
DESOXIMETASONE	DESOXIMETASONE CREAM 0.25%	90550040003710	Generic
DESOXIMETASONE	DESOXIMETASONE GEL 0.05%	90550040004005	Generic
DESOXIMETASONE	DESOXIMETASONE OINT 0.05%	90550040004203	Generic
DESOXIMETASONE	DESOXIMETASONE OINT 0.25%	90550040004205	Generic
DIFLORASONE DIACETATE	DIFLORASONE DIACETATE CREAM 0.05%	90550050103705	Generic
DIFLORASONE DIACETATE	DIFLORASONE DIACETATE OINT 0.05%	90550050104205	Generic
APEXICON E	DIFLORASONE DIACETATE EMOLLIENT BASE CREAM 0.05%	90550050153705	Brand
FLUOCINONIDE	FLUOCINONIDE CREAM 0.1%	90550060003710	Generic
FLURANDRENOLIDE	FLURANDRENOLIDE CREAM 0.05%	90550065003710	Generic
FLURANDRENOLIDE	FLURANDRENOLIDE LOTION 0.05%	90550065004105	Generic
HALCINONIDE	HALCINONIDE CREAM 0.1%	90550070003710	Generic
HALOG	HALCINONIDE OINT 0.1%	90550070004205	Brand
ULTRAVATE	HALOBETASOL PROPIONATE LOTION 0.05%	90550073104110	Brand
TEXACORT	HYDROCORTISONE SOLN 2.5%	90550075002020	Brand
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE SOLN 0.1%	90550075302020	Generic
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE CREAM 0.1%	90550075303705	Generic

HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE LOTION 0.1%	90550075304120	Generic
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE OINT 0.1%	90550075304205	Generic
HYDROCORTISONE BUTYRATE (LIPID)	HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%	90550075323705	Generic
HYDROCORTISONE BUTYRATE (LIPOPHILIC)	HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%	90550075323705	Generic
PANDEL	HYDROCORTISONE PROBUTATE CREAM 0.1%	90550075273720	Brand
HYDROCORTISONE VALERATE	HYDROCORTISONE VALERATE OINT 0.2%	90550075204205	Generic
HYDROCORTISONE VALERATE	HYDROCORTISONE VALERATE CREAM 0.2%	90550075203705	Generic
TRIAMCINOLONE ACETONIDE	TRIAMCINOLONE ACETONIDE AEROSOL SOLN 0.147 MG/GM	90550085103400	Generic
AMCINONIDE	AMCINONIDE CREAM 0.1%	90550010003705	Generic
BRYHALI	HALOBETASOL PROPIONATE LOTION 0.01%	90550073104105	Brand
HALOBETASOL PROPIONATE	HALOBETASOL PROPIONATE FOAM 0.05%	90550073103920	Generic

1 - Trial and failure, contraindication, or intolerance of a preferred topical steroid in comparable potency and/or formulation

Product Name: AMCINONIDE OINT 0.1%, SERNIVO, IMPOYZ, CLOCORTOLONE PIVALATE CREAM 0.1%, DESONIDE CREAM 0.05%, VERDESO, DESOXIMETASONE SPRAY 0.25%, DESOXIMETASONE CREAM 0.05%, DESOXIMETASONE CREAM 0.25% DESOXIMETASONE GEL 0.05%, DESOXIMETASONE OINT 0.05%, DESOXIMETASONE OINT 0.05%, DIFLORASONE DIACETATE CREAM 0.05%, DIFLORASONE DIACETATE OINT 0.05%, APEXICON E, FLUOCINONIDE CREAM 0.1%, FLURANDRENOLIDE CREAM 0.05%, FLURANDRENOLIDE LOTION 0.05%, HALCINONIDE CREAM 0.1%, HALOG, ULTRAVATE, TEXACORT, HYDROCORTISONE BUTYRATE SOLN 0.1%, HYDROCORTISONE BUTYRATE CREAM 0.1%, HYDROCORTISONE BUTYRATE LOTION 0.1%, HYDROCORTISONE BUTYRATE OINT 0.1%, HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%, HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%, PANDEL, HYDROCORTISONE VALERATE OINT 0.2%, HYDROCORTISONE VALERATE CREAM 0.2%, TRIAMCINOLONE ACETONIDE AEROSOL SOLN 0.147 MG/GM, AMCINONIDE CREAM 0.1%, BRYHALI, HALOBETASOL PROPIONATE FOAM 0.05%

Approval Length 12 month(s)

Therapy Stage	Reauthorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
AMCINONIDE	AMCINONIDE OINT 0.1%	90550010004205	Generic
SERNIVO	BETAMETHASONE DIPROPIONATE SPRAY EMULSION 0.05% (BASE EQUIV)	90550020001620	Brand
IMPOYZ	CLOBETASOL PROPIONATE CREAM 0.025%	90550025103703	Brand
CLOCORTOLONE PIVALATE	CLOCORTOLONE PIVALATE CREAM 0.1%	90550030103705	Generic
DESONIDE	DESONIDE CREAM 0.05%	90550035003705	Generic
VERDESO	DESONIDE FOAM 0.05%	90550035003920	Brand
DESOXIMETASONE	DESOXIMETASONE SPRAY 0.25%	90550040000910	Generic
DESOXIMETASONE	DESOXIMETASONE CREAM 0.05%	90550040003705	Generic
DESOXIMETASONE	DESOXIMETASONE CREAM 0.25%	90550040003710	Generic
DESOXIMETASONE	DESOXIMETASONE GEL 0.05%	90550040004005	Generic
DESOXIMETASONE	DESOXIMETASONE OINT 0.05%	90550040004203	Generic
DESOXIMETASONE	DESOXIMETASONE OINT 0.25%	90550040004205	Generic
DIFLORASONE DIACETATE	DIFLORASONE DIACETATE CREAM 0.05%	90550050103705	Generic
DIFLORASONE DIACETATE	DIFLORASONE DIACETATE OINT 0.05%	90550050104205	Generic
APEXICON E	DIFLORASONE DIACETATE EMOLLIENT BASE CREAM 0.05%	90550050153705	Brand
FLUOCINONIDE	FLUOCINONIDE CREAM 0.1%	90550060003710	Generic
FLURANDRENOLIDE	FLURANDRENOLIDE CREAM 0.05%	90550065003710	Generic
FLURANDRENOLIDE	FLURANDRENOLIDE LOTION 0.05%	90550065004105	Generic
HALCINONIDE	HALCINONIDE CREAM 0.1%	90550070003710	Generic
HALOG	HALCINONIDE OINT 0.1%	90550070004205	Brand
ULTRAVATE	HALOBETASOL PROPIONATE LOTION 0.05%	90550073104110	Brand
TEXACORT	HYDROCORTISONE SOLN 2.5%	90550075002020	Brand
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE SOLN 0.1%	90550075302020	Generic
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE CREAM 0.1%	90550075303705	Generic
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE LOTION 0.1%	90550075304120	Generic

HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE OINT 0.1%	90550075304205	Generic
HYDROCORTISONE BUTYRATE (LIPID)	HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%	90550075323705	Generic
HYDROCORTISONE BUTYRATE (LIPOPHILIC)	HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%	90550075323705	Generic
PANDEL	HYDROCORTISONE PROBUTATE CREAM 0.1%	90550075273720	Brand
HYDROCORTISONE VALERATE	HYDROCORTISONE VALERATE OINT 0.2%	90550075204205	Generic
HYDROCORTISONE VALERATE	HYDROCORTISONE VALERATE CREAM 0.2%	90550075203705	Generic
TRIAMCINOLONE ACETONIDE	TRIAMCINOLONE ACETONIDE AEROSOL SOLN 0.147 MG/GM	90550085103400	Generic
AMCINONIDE	AMCINONIDE CREAM 0.1%	90550010003705	Generic
HALOBETASOL PROPIONATE	HALOBETASOL PROPIONATE FOAM 0.05%	90550073103920	Generic
BRYHALI	HALOBETASOL PROPIONATE LOTION 0.01%	90550073104105	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Notes	*Members new to the plan (as evidenced by coverage effective date o
	f less than or equal to 90 days) must meet the initial criteria for covera
	ge

Product Name: AMCINONIDE OINT 0.1%, SERNIVO, IMPOYZ, CLOCORTOLONE PIVALATE CREAM 0.1%, DESONIDE CREAM 0.05%, VERDESO, DESOXIMETASONE SPRAY 0.25%, DESOXIMETASONE CREAM 0.05%, DESOXIMETASONE CREAM 0.25% DESOXIMETASONE GEL 0.05%, DESOXIMETASONE OINT 0.05%, DESOXIMETASONE OINT 0.25%, DIFLORASONE DIACETATE CREAM 0.05%, DIFLORASONE DIACETATE OINT 0.05%, APEXICON E, FLUOCINONIDE CREAM 0.1%, FLURANDRENOLIDE CREAM 0.05%, FLURANDRENOLIDE LOTION 0.05%, HALCINONIDE CREAM 0.1%, HALOG, ULTRAVATE, TEXACORT, HYDROCORTISONE BUTYRATE SOLN 0.1%, HYDROCORTISONE BUTYRATE CREAM 0.1%, HYDROCORTISONE BUTYRATE LOTION 0.1%, HYDROCORTISONE BUTYRATE OINT 0.1%, HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%, HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%, PANDEL, HYDROCORTISONE VALERATE OINT 0.2%, HYDROCORTISONE VALERATE CREAM 0.2%, TRIAMCINOLONE ACETONIDE AEROSOL SOLN 0.147 MG/GM, AMCINONIDE CREAM 0.1%, BRYHALI, HALOBETASOL PROPIONATE FOAM 0.05%

Approval Length	12/31/2039
Guideline Type	Prior Authorization-All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
AMCINONIDE	AMCINONIDE OINT 0.1%	90550010004205	Generic
SERNIVO	BETAMETHASONE DIPROPIONATE SPRAY EMULSION 0.05% (BASE EQUIV)	90550020001620	Brand
IMPOYZ	CLOBETASOL PROPIONATE CREAM 0.025%	90550025103703	Brand
CLOCORTOLONE PIVALATE	CLOCORTOLONE PIVALATE CREAM 0.1%	90550030103705	Generic
DESONIDE	DESONIDE CREAM 0.05%	90550035003705	Generic
VERDESO	DESONIDE FOAM 0.05%	90550035003920	Brand
DESOXIMETASONE	DESOXIMETASONE SPRAY 0.25%	90550040000910	Generic
DESOXIMETASONE	DESOXIMETASONE CREAM 0.05%	90550040003705	Generic
DESOXIMETASONE	DESOXIMETASONE CREAM 0.25%	90550040003710	Generic
DESOXIMETASONE	DESOXIMETASONE GEL 0.05%	90550040004005	Generic
DESOXIMETASONE	DESOXIMETASONE OINT 0.05%	90550040004203	Generic
DESOXIMETASONE	DESOXIMETASONE OINT 0.25%	90550040004205	Generic
DIFLORASONE DIACETATE	DIFLORASONE DIACETATE CREAM 0.05%	90550050103705	Generic
DIFLORASONE DIACETATE	DIFLORASONE DIACETATE OINT 0.05%	90550050104205	Generic
APEXICON E	DIFLORASONE DIACETATE EMOLLIENT BASE CREAM 0.05%	90550050153705	Brand
FLUOCINONIDE	FLUOCINONIDE CREAM 0.1%	90550060003710	Generic
FLURANDRENOLIDE	FLURANDRENOLIDE CREAM 0.05%	90550065003710	Generic
FLURANDRENOLIDE	FLURANDRENOLIDE LOTION 0.05%	90550065004105	Generic
HALCINONIDE	HALCINONIDE CREAM 0.1%	90550070003710	Generic
HALOG	HALCINONIDE OINT 0.1%	90550070004205	Brand
ULTRAVATE	HALOBETASOL PROPIONATE LOTION 0.05%	90550073104110	Brand
TEXACORT	HYDROCORTISONE SOLN 2.5%	90550075002020	Brand
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE SOLN 0.1%	90550075302020	Generic
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE CREAM 0.1%	90550075303705	Generic
HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE LOTION 0.1%	90550075304120	Generic

HYDROCORTISONE BUTYRATE	HYDROCORTISONE BUTYRATE OINT 0.1%	90550075304205	Generic
HYDROCORTISONE BUTYRATE (LIPID)	HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%	90550075323705	Generic
HYDROCORTISONE BUTYRATE (LIPOPHILIC)	HYDROCORTISONE BUTYRATE HYDROPHILIC LIPO BASE CREAM 0.1%	90550075323705	Generic
PANDEL	HYDROCORTISONE PROBUTATE CREAM 0.1%	90550075273720	Brand
HYDROCORTISONE VALERATE	HYDROCORTISONE VALERATE OINT 0.2%	90550075204205	Generic
HYDROCORTISONE VALERATE	HYDROCORTISONE VALERATE CREAM 0.2%	90550075203705	Generic
TRIAMCINOLONE ACETONIDE	TRIAMCINOLONE ACETONIDE AEROSOL SOLN 0.147 MG/GM	90550085103400	Generic
AMCINONIDE	AMCINONIDE CREAM 0.1%	90550010003705	Generic
BRYHALI	HALOBETASOL PROPIONATE LOTION 0.01%	90550073104105	Brand
HALOBETASOL PROPIONATE	HALOBETASOL PROPIONATE FOAM 0.05%	90550073103920	Generic

1 - Trial and failure, contraindication, or intolerance of a preferred topical steroid in comparable potency and/or formulation

Date	Notes
4/15/2024	Guideline Update.

Non-Sedating Antihistamine			
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Guideline ID	GL-129167
Guideline Name	Non-Sedating Antihistamine
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

1. Criteria

Product Name: Generic desloratadine, Clarinex D (desloratadine/pseudoephedrine)				
Diagnosis Allergic Rhinitis				
Approval Length		12/31/2039		
Guideline Type Prior Authorization - All Plans except IL and		t IL and MN Plans	3	
Product Name	Gene	ric Name	GPI	Brand/Generic
DESLORATADINE	DESLO	DRATADINE TAB 5 MG	41550021000320	Generic
CLARINEX-D 12 HOUR		DRATADINE & PSEUDOEPHEDRINE TAB HR 2.5-120 MG	43993002627420	Brand

DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 2.5 MG	41550021007210	Generic
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 5 MG	41550021007220	Generic

1 - Diagnosis of allergic rhinitis

AND

- **2** Trial and failure, contraindication, or intolerance to ALL of the following over the counter (OTC) agents:
 - Cetirizine
 - Fexofenadine
 - Levocetirizine
 - Loratadine

AND

3 - Trial and failure, contraindication, or intolerance to one nasal steroid* (e.g., fluticasone)

Notes	*Note: The nasal steroid criterion does not apply in the case of predict
	able situational exposures where nasal steroids would not be the best
	clinical choice or for children 12 years of age or younger.

Product Name: Generic desloratadine, Clarinex D (desloratadine/pseudoephedrine)		
Diagnosis	Allergic Rhinitis	
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DESLORATADINE	DESLORATADINE TAB 5 MG	41550021000320	Generic
CLARINEX-D 12 HOUR	DESLORATADINE & PSEUDOEPHEDRINE TAB ER 12HR 2.5-120 MG	43993002627420	Brand
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 2.5 MG	41550021007210	Generic

DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 5 MG	41550021007220	Generic
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1 - Diagnosis of allergic rhinitis

AND

- **2** Trial and failure, contraindication, or intolerance to ALL of the following over the counter (OTC) agents:
 - Cetirizine
 - Fexofenadine
 - Levocetirizine
 - Loratadine

AND

3 - Trial and failure, contraindication, or intolerance to one nasal steroid* (e.g., fluticasone)

Notes	*Note: The nasal steroid criterion does not apply in the case of predict
	able situational exposures where nasal steroids would not be the best
	clinical choice or for children 12 years of age or younger.

Product Name: Generic	Product Name: Generic desloratadine, Clarinex D (desloratadine/pseudoephedrine)	
Diagnosis	Diagnosis Urticarial Disease	
Approval Length	12/31/2039	
Guideline Type Prior Authorization - All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
DESLORATADINE	DESLORATADINE TAB 5 MG	41550021000320	Generic
CLARINEX-D 12 HOUR	DESLORATADINE & PSEUDOEPHEDRINE TAB ER 12HR 2.5-120 MG	43993002627420	Brand
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 2.5 MG	41550021007210	Generic
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 5 MG	41550021007220	Generic

1 - Diagnosis of urticarial disease

AND

- **2** Trial and failure, contraindication, or intolerance to ALL of the following over the counter (OTC) agents:
 - Cetirizine
 - Fexofenadine
 - Levocetirizine
 - Loratadine

Product Name: Generic	Product Name: Generic desloratadine, Clarinex D (desloratadine/pseudoephedrine)	
Diagnosis Urticarial Disease		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DESLORATADINE	DESLORATADINE TAB 5 MG	41550021000320	Generic
CLARINEX-D 12 HOUR	DESLORATADINE & PSEUDOEPHEDRINE TAB ER 12HR 2.5-120 MG	43993002627420	Brand
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 2.5 MG	41550021007210	Generic
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 5 MG	41550021007220	Generic

Approval Criteria

1 - Diagnosis of urticarial disease

AND

- **2** Trial and failure, contraindication, or intolerance to ALL of the following over the counter (OTC) agents:
 - Cetirizine
 - Fexofenadine
 - Levocetirizine
 - Loratadine

Product Name: Generic	Product Name: Generic desloratadine, Clarinex D (desloratadine/pseudoephedrine)	
Diagnosis All Indications		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DESLORATADINE	DESLORATADINE TAB 5 MG	41550021000320	Generic
CLARINEX-D 12 HOUR	DESLORATADINE & PSEUDOEPHEDRINE TAB ER 12HR 2.5-120 MG	43993002627420	Brand
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 2.5 MG	41550021007210	Generic
DESLORATADINE ODT	DESLORATADINE TAB ORALLY DISINTEGRATING 5 MG	41550021007220	Generic

1 - Prescriber provides clinical documentation from the past 12 months that the member is continuing therapy on the requested drug

Date	Notes
9/27/2023	2024 New Implementation

Non-solid Dosage Forms	S
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Guideline ID	GL-132813
Guideline Name	Non-solid Dosage Forms
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Famotidine Suspension 40 mg/5mL, Generic Naproxen Suspension 125 mg/ 5mL, Generic Sevelamer packet, Brand Valsartan solution 4 mg, mL, Atorvaliq, Generic Baclofen solution 5 mg/5mL, Thyquidity, Flolipid, Zonisade, Norliqva, Katerzia, generic esomeprazole gran, Nexium gran, generic lansoprazole ODT, Prilosec Powder, Aspruzyo

Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN plans only	

Product Name	Generic Name	GPI	Brand/Generic
FAMOTIDINE	FAMOTIDINE FOR SUSP 40 MG/5ML	49200030001920	Generic
NAPROXEN	NAPROXEN SUSP 125 MG/5ML	66100060001805	Generic
SEVELAMER CARBONATE	SEVELAMER CARBONATE PACKET 0.8 GM	52800070053020	Generic

SEVELAMER CARBONATE	SEVELAMER CARBONATE PACKET 2.4 GM	52800070053040	Generic
VALSARTAN	VALSARTAN ORAL SOLN 4 MG/ML	36150080002025	Brand
ATORVALIQ	ATORVASTATIN CALCIUM SUSP 20 MG/5ML (4MG/ML) (BASE EQUIV)	39400010101810	Brand
BACLOFEN	BACLOFEN ORAL SOLN 5 MG/5ML	75100010002070	Generic
THYQUIDITY	LEVOTHYROXINE SODIUM ORAL SOLUTION 100 MCG/5ML	28100010102023	Brand
FLOLIPID	SIMVASTATIN SUSP 20 MG/5ML (4 MG/ML)	39400075001810	Brand
FLOLIPID	SIMVASTATIN SUSP 40 MG/5ML (8 MG/ML)	39400075001820	Brand
ZONISADE	ZONISAMIDE ORAL SUSP 100 MG/5ML (20 MG/ML)	72600090001820	Brand
NORLIQVA	AMLODIPINE BESYLATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	34000003102020	Brand
KATERZIA	AMLODIPINE BENZOATE ORAL SUSP 1 MG/ML (BASE EQUIVALENT)	34000003081820	Brand
ESOMEPRAZOLE MAGNESIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 10 MG	49270025103010	Generic
ESOMEPRAZOLE MAGNESIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 20 MG	49270025103020	Generic
ESOMEPRAZOLE MAGNESIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 40 MG	49270025103040	Generic
NEXIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACK 2.5 MG	49270025103004	Brand
NEXIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 5 MG	49270025103007	Brand
LANSOPRAZOLE ODT	LANSOPRAZOLE TAB DELAYED RELEASE ORALLY DISINTEGRATING 15 MG	4927004000H315	Generic
LANSOPRAZOLE ODT	LANSOPRAZOLE TAB DELAYED RELEASE ORALLY DISINTEGRATING 30 MG	4927004000H330	Generic
PRILOSEC	OMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 2.5 MG	49270060103020	Brand
PRILOSEC	OMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 10 MG	49270060103030	Brand
ASPRUZYO SPRINKLE	RANOLAZINE ER GRANULES PACKET 500 MG	32200040003020	Brand
ASPRUZYO SPRINKLE	RANOLAZINE ER GRANULES PACKET 1000 MG	32200040003040	Brand

1 - Unable to tolerate solid dose form

OR

2 - Age is less than 12 years old*

OR

3 - Minnesota Plans Only - Member has stage four metastatic cancer and the requested drug is being used as supportive care to treat symptoms directly related to their cancer or chemotherapy regimen

Notes	*Age edit does not apply to Zonisamide oral suspension because Zoni
	samide is only approved for age 16 and older.

Product Name: Generic Famotidine Suspension 40 mg/5mL, Generic Naproxen Suspension 125 mg/ 5mL, Generic Sevelamer packet, Brand Valsartan solution 4 mg, mL, Atorvaliq, Generic Baclofen solution 5 mg/5mL, Thyquidity, Flolipid, Zonisade, Norliqva, Katerzia, generic esomeprazole gran, Nexium gran, generic lansoprazole ODT, Prilosec Powder, Aspruzyo

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN plans only

Product Name	Generic Name	GPI	Brand/Generic
FAMOTIDINE	FAMOTIDINE FOR SUSP 40 MG/5ML	49200030001920	Generic
NAPROXEN	NAPROXEN SUSP 125 MG/5ML	66100060001805	Generic
SEVELAMER CARBONATE	SEVELAMER CARBONATE PACKET 0.8 GM	52800070053020	Generic
SEVELAMER CARBONATE	SEVELAMER CARBONATE PACKET 2.4 GM	52800070053040	Generic
VALSARTAN	VALSARTAN ORAL SOLN 4 MG/ML	36150080002025	Brand
ATORVALIQ	ATORVASTATIN CALCIUM SUSP 20 MG/5ML (4MG/ML) (BASE EQUIV)	39400010101810	Brand
BACLOFEN	BACLOFEN ORAL SOLN 5 MG/5ML	75100010002070	Generic
THYQUIDITY	LEVOTHYROXINE SODIUM ORAL SOLUTION 100 MCG/5ML	28100010102023	Brand
FLOLIPID	SIMVASTATIN SUSP 20 MG/5ML (4 MG/ML)	39400075001810	Brand
FLOLIPID	SIMVASTATIN SUSP 40 MG/5ML (8 MG/ML)	39400075001820	Brand

ZONISADE	ZONISAMIDE ORAL SUSP 100 MG/5ML (20 MG/ML)	72600090001820	Brand
NORLIQVA	AMLODIPINE BESYLATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	34000003102020	Brand
KATERZIA	AMLODIPINE BENZOATE ORAL SUSP 1 MG/ML (BASE EQUIVALENT)	34000003081820	Brand
ESOMEPRAZOLE MAGNESIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 10 MG	49270025103010	Generic
ESOMEPRAZOLE MAGNESIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 20 MG	49270025103020	Generic
ESOMEPRAZOLE MAGNESIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 40 MG	49270025103040	Generic
NEXIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACK 2.5 MG	49270025103004	Brand
NEXIUM	ESOMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 5 MG	49270025103007	Brand
LANSOPRAZOLE ODT	LANSOPRAZOLE TAB DELAYED RELEASE ORALLY DISINTEGRATING 15 MG	4927004000H315	Generic
LANSOPRAZOLE ODT	LANSOPRAZOLE TAB DELAYED RELEASE ORALLY DISINTEGRATING 30 MG	4927004000H330	Generic
PRILOSEC	OMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 2.5 MG	49270060103020	Brand
PRILOSEC	OMEPRAZOLE MAGNESIUM FOR DELAYED RELEASE SUSP PACKET 10 MG	49270060103030	Brand
ASPRUZYO SPRINKLE	RANOLAZINE ER GRANULES PACKET 500 MG	32200040003020	Brand
ASPRUZYO SPRINKLE	RANOLAZINE ER GRANULES PACKET 1000 MG	32200040003040	Brand

1 - Clinical documentation from the previous 12 months demonstrating a positive response to therapy

Notes	*Age edit does not apply to Zonisamide oral suspension because Zoni
	samide is only approved for age 16 and older.

Product Name: Generic Famotidine Suspension 40 mg/5mL, Generic Naproxen Suspension 125 mg/ 5mL, Generic Sevelamer packet, Brand Valsartan solution 4 mg, mL, Atorvaliq, Generic Baclofen solution 5 mg/5mL, Thyquidity, Flolipid, Zonisade, Norliqva, Katerzia, generic esomeprazole gran, Nexium gran, generic lansoprazole ODT, Prilosec Powder, Aspruzyo

Aspruzyo	
Approval Length	12/31/2039

Guideline Type Prior Authorization - All plans except IL and MN				
Product Name	Gener	ic Name	GPI	Brand/Generic
FAMOTIDINE	FAMOTIDINE FOR SUSP 40 MG/5ML		49200030001920	Generic
NAPROXEN	NAPRO	XEN SUSP 125 MG/5ML	66100060001805	Generic
SEVELAMER CARBONATE	SEVEL	AMER CARBONATE PACKET 0.8 GM	52800070053020	Generic
SEVELAMER CARBONATE	SEVEL	AMER CARBONATE PACKET 2.4 GM	52800070053040	Generic
VALSARTAN	VALSA	RTAN ORAL SOLN 4 MG/ML	36150080002025	Brand
ATORVALIQ		ASTATIN CALCIUM SUSP 20 MG/5ML IL) (BASE EQUIV)	39400010101810	Brand
BACLOFEN	BACLO	FEN ORAL SOLN 5 MG/5ML	75100010002070	Generic
THYQUIDITY	LEVOT 100 MC	HYROXINE SODIUM ORAL SOLUTION CG/5ML	28100010102023	Brand
FLOLIPID	SIMVAS	STATIN SUSP 20 MG/5ML (4 MG/ML)	39400075001810	Brand
FLOLIPID	SIMVAS	STATIN SUSP 40 MG/5ML (8 MG/ML)	39400075001820	Brand
ZONISADE	ZONISA MG/ML	AMIDE ORAL SUSP 100 MG/5ML (20)	72600090001820	Brand
NORLIQVA		DIPINE BESYLATE ORAL SOLN 1 MG/ML EQUIVALENT)	34000003102020	Brand
KATERZIA		DIPINE BENZOATE ORAL SUSP 1 MG/ML EQUIVALENT)	34000003081820	Brand
ESOMEPRAZOLE MAGNESIUM		PRAZOLE MAGNESIUM FOR DELAYED SE SUSP PACKET 10 MG	49270025103010	Generic
ESOMEPRAZOLE MAGNESIUM		PRAZOLE MAGNESIUM FOR DELAYED SE SUSP PACKET 20 MG	49270025103020	Generic
ESOMEPRAZOLE MAGNESIUM		PRAZOLE MAGNESIUM FOR DELAYED SE SUSP PACKET 40 MG	49270025103040	Generic
NEXIUM		EPRAZOLE MAGNESIUM FOR DELAYED SE SUSP PACK 2.5 MG	49270025103004	Brand
NEXIUM		PRAZOLE MAGNESIUM FOR DELAYED SE SUSP PACKET 5 MG	49270025103007	Brand
LANSOPRAZOLE ODT	_	PRAZOLE TAB DELAYED RELEASE Y DISINTEGRATING 15 MG	4927004000H315	Generic
LANSOPRAZOLE ODT	_	PRAZOLE TAB DELAYED RELEASE Y DISINTEGRATING 30 MG	4927004000H330	Generic
PRILOSEC		RAZOLE MAGNESIUM FOR DELAYED SE SUSP PACKET 2.5 MG	49270060103020	Brand
PRILOSEC		RAZOLE MAGNESIUM FOR DELAYED SE SUSP PACKET 10 MG	49270060103030	Brand
ASPRUZYO SPRINKLE	RANOL	AZINE ER GRANULES PACKET 500 MG	32200040003020	Brand

ASPRUZYO SPRINKLE	RANOLAZINE ER GRANULES PACKET 1000 MG	32200040003040	Brand
Approval Crite	ria		
1 - Unable to tolerate solid dose form			
OR			
2 - Age is less than 12 years old*			
Notes	*Age edit does not apply to Zonisar samide is only approved for age 16		on because Zoni

Date	Notes
11/28/2023	New Program

Nonprefe	rred Bowel Preparations	
The Stand Programmed Standings, The Review Standings and American	ome, som til Sakris jagski promiti aralin	

Guideline ID	GL-144876	
Guideline Name	onpreferred Bowel Preparations	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	7/20/2016
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Clenpiq, Plenvu, Sodium sulfate/Potassium sulfate/Magnesium sulfate, Osmoprep, Suflave				
Approval Length	12 month(s)	12 month(s)		
Therapy Stage	Initial Authorization	Initial Authorization		
Guideline Type	Prior Authorization-IL and MN Plans	Prior Authorization-IL and MN Plans Only		
Product Name	Generic Name	GPI	Brand/Generic	
CLENPIQ	SOD PICOSULFATE-MG OX-CITRIC AC SOL 10 MG-3.5 GM-12 GM/160ML	46992003452020	Brand	
CLENPIQ	OD PICOSULFATE-MG OX-CITRIC AC 46992003452030 Brand OL 10 MG-3.5 GM-12 GM/175ML		Brand	
SODIUM SULFATE/POTASSIUM	SOD SULFATE-POT SULF-MG SULF ORAL SOL 17.5-3.13-1.6 GM/177ML			

SULFATE/MAGNESIUM SULFATE			
PLENVU	PEG 3350-KCL-NACL-NA SULFATE-NA ASCORBATE-C FOR SOLN 140 GM	46992006302135	Brand
SUFLAVE	PEG 3350-KCL-NACL-NA SULFATE-MAG SULFATE FOR SOLN 178.7 GM	46992005382150	Brand
OSMOPREP	SOD PHOS MONO-SOD PHOS DI TABS 1.102-0.398 GM(1.5GM NA PHOS)	46109902120320	Brand

1 - Trial and failure, contraindication or intolerance to a preferred bowel preparation

Product Name: Clenpiq, Plenvu, Sodium sulfate/Potassium sulfate/Magnesium sulfate, Osmoprep, Suflave	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name GPI Bra		Brand/Generic
CLENPIQ	SOD PICOSULFATE-MG OX-CITRIC AC SOL 10 MG-3.5 GM-12 GM/160ML	46992003452020 Brand	
CLENPIQ	SOD PICOSULFATE-MG OX-CITRIC AC SOL 10 MG-3.5 GM-12 GM/175ML	46992003452030	Brand
SODIUM SULFATE/POTASSIUM SULFATE/MAGNESIUM SULFATE	SOD SULFATE-POT SULF-MG SULF ORAL SOL 17.5-3.13-1.6 GM/177ML	46992003602020	Generic
PLENVU	PEG 3350-KCL-NACL-NA SULFATE-NA ASCORBATE-C FOR SOLN 140 GM	46992006302135	Brand
SUFLAVE	PEG 3350-KCL-NACL-NA SULFATE-MAG SULFATE FOR SOLN 178.7 GM	46992005382150	Brand
OSMOPREP	SOD PHOS MONO-SOD PHOS DI TABS 1.102-0.398 GM(1.5GM NA PHOS)	46109902120320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Clenpiq, Plenvu, Sodium sulfate/Potassium sulfate/Magnesium sulfate, Osmoprep, Suflave	
Approval Length	12/31/2039
Guideline Type Prior Authorization-All plans except IL and MN	

Product Name	Generic Name GPI B		Brand/Generic
CLENPIQ	SOD PICOSULFATE-MG OX-CITRIC AC SOL 10 MG-3.5 GM-12 GM/160ML	46992003452020 Brand	
CLENPIQ	SOD PICOSULFATE-MG OX-CITRIC AC SOL 10 MG-3.5 GM-12 GM/175ML	46992003452030	Brand
SODIUM SULFATE/POTASSIUM SULFATE/MAGNESIUM SULFATE	SOD SULFATE-POT SULF-MG SULF ORAL SOL 17.5-3.13-1.6 GM/177ML	46992003602020	Generic
PLENVU	PEG 3350-KCL-NACL-NA SULFATE-NA ASCORBATE-C FOR SOLN 140 GM	46992006302135	Brand
SUFLAVE	PEG 3350-KCL-NACL-NA SULFATE-MAG SULFATE FOR SOLN 178.7 GM	46992005382150	Brand
OSMOPREP	SOD PHOS MONO-SOD PHOS DI TABS 1.102-0.398 GM(1.5GM NA PHOS)	46109902120320	Brand

1 - Trial and failure, contraindication or intolerance to a preferred bowel preparation

Date	Notes
3/26/2024	Added Osmoprep, and Suflave

Nonpreferred insulin			
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Guideline ID	GL-131426	
Guideline Name	lonpreferred insulin	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Apidra, Humalog Mix 50:50		
Approval Length 12 month(s)		
Therapy Stage	Therapy Stage Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
HUMALOG MIX 50/50	INSULIN LISPRO PROTAMINE & LISPRO INJ 100 UNIT/ML (50-50)	27104080001840	Brand
APIDRA SOLOSTAR	INSULIN GLULISINE SOLN PEN-INJECTOR INJ 100 UNIT/ML	2710400400D220	Brand
HUMALOG MIX 50/50 KWIKPEN	INSULIN LISPRO PROT & LISPRO SUS PEN-INJ 100 UNIT/ML (50-50)	2710408000D340	Brand
APIDRA	INSULIN GLULISINE INJ 100 UNIT/ML	27104004002022	Brand

1 - Diagnosis of diabetes mellitus

AND

2 - Trial and failure, contraindication or intolerance to use of insulin aspart (Novolog, Novolog Mix) including dose adjustments

Product Name: Apidra, Humalog Mix 50:50	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
HUMALOG MIX 50/50	INSULIN LISPRO PROTAMINE & LISPRO INJ 100 UNIT/ML (50-50)	27104080001840	Brand
APIDRA SOLOSTAR	INSULIN GLULISINE SOLN PEN-INJECTOR INJ 100 UNIT/ML	2710400400D220	Brand
HUMALOG MIX 50/50 KWIKPEN	INSULIN LISPRO PROT & LISPRO SUS PEN-INJ 100 UNIT/ML (50-50)	2710408000D340	Brand
APIDRA	INSULIN GLULISINE INJ 100 UNIT/ML	27104004002022	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Apidra, Humalog Mix 50:50				
Approval Length 12/31/2039				
Guideline Type		Prior Authorization-All plans except IL and MN		
Product Generic Name Name		ame	GPI	Brand/Generic

HUMALOG MIX 50/50	INSULIN LISPRO PROTAMINE & LISPRO INJ 100 UNIT/ML (50-50)	27104080001840	Brand
APIDRA SOLOSTAR	INSULIN GLULISINE SOLN PEN-INJECTOR INJ 100 UNIT/ML	2710400400D220	Brand
HUMALOG MIX 50/50 KWIKPEN	INSULIN LISPRO PROT & LISPRO SUS PEN-INJ 100 UNIT/ML (50-50)	2710408000D340	Brand
APIDRA	INSULIN GLULISINE INJ 100 UNIT/ML	27104004002022	Brand

1 - Diagnosis of diabetes mellitus

AND

2 - Trial and failure, contraindication or intolerance to use of insulin aspart (Novolog, Novolog Mix) including dose adjustments

Date	Notes
10/9/2023	New Program

Nonsteroidal Anti-inflammatory (NSAID) Combination		
(g) belongs an enderly belong to the section of the		

Guideline ID	GL-131404	
Guideline Name	Nonsteroidal Anti-inflammatory (NSAID) Combinations	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Ibuprofen/famotidine, Naproxen/esomeprazole	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
IBUPROFEN/FAMOTIDINE	IBUPROFEN-FAMOTIDINE TAB 800-26.6 MG	66109902320340	Generic
NAPROXEN/ESOMEPRAZOLE MAGNESIUM	NAPROXEN-ESOMEPRAZOLE MAGNESIUM TAB DR 375-20 MG	66109902440620	Generic
NAPROXEN/ESOMEPRAZOLE MAGNESIUM	NAPROXEN-ESOMEPRAZOLE MAGNESIUM TAB DR 500-20 MG	66109902440640	Generic

Approval Criteria

1 - Diagnosis of osteoarthritis, rheumatoid arthritis, or other pain-related condition requiring chronic NSAID use

AND

2 - Diagnosis of current or past gastric ulcer

AND

3 - Trial and failure after an adequate trial of the equivalent NSAID and histamine blocker or proton pump inhibitor as separate products

AND

4 - Prescriber supplies published literature to support the requested combination product will produce different clinical results than using the equivalent agents as separate products

OR

5 - For Minnesota Plans Only

MAGNESIUM

NAPROXEN/ESOMEPRAZOLE NAPROXEN-ESOMEPRAZOLE

5.1 Diagnosis of stage four metastatic cancer and the requested drug is being used to treat cancer-related pain

Product Name: Generic Ibuprofen/famotidine, Naproxen/esomeprazole				
Approval Length	12 r	12 month(s)		
Therapy Stage	Rea	Reauthorization		
Guideline Type	Pric	Prior Authorization-IL and MN Plans Only		
Product Name		Generic Name	GPI	Brand/Generic
IBUPROFEN/FAMOTIDINE		IBUPROFEN-FAMOTIDINE TAB 800-26.6 MG	66109902320340	Generic
NAPROXEN/ESOMEPRAZ MAGNESIUM	OLE	NAPROXEN-ESOMEPRAZOLE MAGNESIUM TAB DR 375-20 MG	66109902440620	Generic

MAGNESIUM TAB DR 500-20 MG

Generic

66109902440640

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

*Members new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) must meet the initial criteria for covera
ge

Product Name: Generic Ibuprofen/famotidine, Naproxen/esomeprazole	
Approval Length	12/31/2039
Guideline Type	Prior Authorization-All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
IBUPROFEN/FAMOTIDINE	IBUPROFEN-FAMOTIDINE TAB 800-26.6 MG	66109902320340	Generic
NAPROXEN/ESOMEPRAZOLE MAGNESIUM	NAPROXEN-ESOMEPRAZOLE MAGNESIUM TAB DR 375-20 MG	66109902440620	Generic
NAPROXEN/ESOMEPRAZOLE MAGNESIUM	NAPROXEN-ESOMEPRAZOLE MAGNESIUM TAB DR 500-20 MG	66109902440640	Generic

Approval Criteria

1 - Diagnosis of osteoarthritis, rheumatoid arthritis, or other pain-related condition requiring chronic NSAID use

AND

2 - Diagnosis of current or past gastric ulcer

AND

3 - Trial and failure after an adequate trial of the equivalent NSAID and histamine blocker or proton pump inhibitor as separate products

AND

4 - Prescriber supplies published literature to support the requested combination product will produce different clinical results than using the equivalent agents as separate products

Date	Notes
10/27/2023	New program

Ν	lorthera (dro	oxidopa)	
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Guideline ID	GL-129157
Guideline Name	Northera (droxidopa)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2023
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1. Criteria

Product Name: Generic Droxidopa*	
Approval Length	See note*
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
DROXIDOPA	DROXIDOPA CAP 100 MG	38700030000130	Generic
DROXIDOPA	DROXIDOPA CAP 200 MG	38700030000140	Generic
DROXIDOPA	DROXIDOPA CAP 300 MG	38700030000150	Generic

Approval Criteria

1 - Diagnosis of symptomatic neurogenic orthostatic hypotension caused by primary autonomic failure (Parkinson's disease, multiple system atrophy, and pure autonomic failure), dopamine beta-hydroxylase deficiency, or nondiabetic autonomic neuropathy

AND

2 - Prescribed by, or in consultation with, a Neurologist

AND

3 - Trial and failure, contraindication, or intolerance to both midodrine and fludrocortisone

Notes	* 2 months with partial fill
	(max 15 days/prescription)

Product Name: Generic Droxidopa		
Approval Length 12/31/2039		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
DROXIDOPA	DROXIDOPA CAP 100 MG	38700030000130	Generic
DROXIDOPA	DROXIDOPA CAP 200 MG	38700030000140	Generic
DROXIDOPA	DROXIDOPA CAP 300 MG	38700030000150	Generic

Approval Criteria

1 - Prescriber provides clinical documentation from the previous two months of demonstrated ongoing beneficial response to therapy.

Product Name: Generic Droxidopa			
Approval Length 12 month(s)			
Therapy Stage Initial Authorization			
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
DROXIDOPA	DROXIDOPA CAP 100 MG	38700030000130	Generic
DROXIDOPA	DROXIDOPA CAP 200 MG	38700030000140	Generic
DROXIDOPA	DROXIDOPA CAP 300 MG	38700030000150	Generic

1 - Diagnosis of symptomatic neurogenic orthostatic hypotension caused by primary autonomic failure (Parkinson's disease, multiple system atrophy, and pure autonomic failure), dopamine beta-hydroxylase deficiency, or nondiabetic autonomic neuropathy

AND

2 - Prescribed by, or in consultation with, a Neurologist

AND

3 - Trial and failure, contraindication, or intolerance to both midodrine and fludrocortisone

Product Name: Generic Droxidopa	
Approval Length 12/31/2039	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DROXIDOPA	DROXIDOPA CAP 100 MG	38700030000130	Generic
DROXIDOPA	DROXIDOPA CAP 200 MG	38700030000140	Generic
DROXIDOPA	DROXIDOPA CAP 300 MG	38700030000150	Generic

Approval Criteria

1 - Prescriber provides clinical documentation from the previous twelve months of demonstrated ongoing beneficial response to therapy.

Date	Notes
9/20/2023	New Program

Nucala (mepolizumab)						
	(g) the best department of the tense to the contract of the co					

Guideline ID GL-144880	
Guideline Name	Nucala (mepolizumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	4/19/2023
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Nucala SC				
Diagnosis		Eosinophilic Asthma		
Approval Length		12/31/2039		
Guideline Type		Prior Authorization - ALL Plans Except IL and MN Plans		
Product Name	Generic Na	me	GPI	Brand/Generic
NUCALA	MEPOLIZUMA INJECTOR 10	AB SUBCUTANEOUS SOLUTION AUTO- 00 MG/ML	4460405500D530	Brand
NUCALA	A MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML		4460405500E520	Brand
NUCALA	NUCALA MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML		4460405500E530	Brand

Approval Criteria
1 - Diagnosis of eosinophilic asthma
AND
2 - Prescribed by or in consultation with an asthma specialist (i.e., allergist, immunologist, pulmonologist)
AND
3 - Member is 6 years of age or older
AND
4 - One of the following:
4.1 All of the following:
4.1.1 Submission of medical records (e.g., chart notes) documenting that the blood eosinophil count is greater than or equal to 150 cells/mm3
AND
4.1.2 Other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease have been ruled out
AND
4.1.3 One of the following:
4.1.3.1 Symptoms are not well controlled or poorly controlled (Table 1) despite adherence* to a greater than or equal to a 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

- **4.1.3.2** Member has an intolerance to medium to high dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier. Exceptions based on adverse effects from medium to high dose ICS or comorbid conditions increasing long-term risks of adverse effects from high dose ICS or oral corticosteroids include one of the following:
 - Cataracts in members older than 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after consultation with an endocrinologist
 - Diagnosis of osteoporosis, whose treatment is resistant to FDA approved osteoporosis treatment

OR

4.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

MPR) greater than or equal to 70%, based on the previous 120 days of		•
**IL-5 inhibitor drugs in combination with omalizumab will be consider ed on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms . Tezepelumab, in combination with other	Notes	ed on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms . Tezepelumab, in combination with other biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both

Product Name: Nucala SC				
Diagnosis Eosinophilic Asthma				
Approval L	Approval Length 12 month(s)			
Guideline ⁻	Guideline Type Prior Authorization - IL and MN Plans			
Product Generic Name		GPI	Brand/Generic	
NUCALA MEPOLIZUMAB SUBCUTANEOUS SOLUTION AUTO- INJECTOR 100 MG/ML 4460405500D		4460405500D530	Brand	

NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML	4460405500E520	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML	4460405500E530	Brand

1 - Diagnosis of eosinophilic asthma

AND

2 - Prescribed by or in consultation with an asthma specialist (i.e., allergist, immunologist, pulmonologist)

AND

3 - Member is 6 years of age or older

AND

- 4 One of the following:
 - **4.1** All of the following:
- **4.1.1** Submission of medical records (e.g., chart notes) documenting that the blood eosinophil count is greater than or equal to 150 cells/mm3

AND

4.1.2 Other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease have been ruled out

AND

- **4.1.3** One of the following:
- 4.1.3.1 Symptoms are not well controlled or poorly controlled (Table 1) despite adherence*

to a greater than or equal to a 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

- **4.1.3.2** Member has an intolerance to medium to high dose inhaled corticosteroids in combination with a long-acting bronchodilator or leukotriene modifier. Exceptions based on adverse effects from medium to high dose ICS or comorbid conditions increasing long-term risks of adverse effects from high dose ICS or oral corticosteroids include one of the following:
 - Cataracts in members older than 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after consultation with an endocrinologist
 - Diagnosis of osteoporosis, whose treatment is resistant to FDA approved osteoporosis treatment

OR

4.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

,	
MPR) greater than of f prescription claims ** IL-5 inhibitor drug ed on a case-by-cas agent with combinat . Tezepelumab, in combined biologics, has not be extenuating circums	ment is defined as a medication possession ratio (or equal to 70%, based on the previous 120 days or some in combination with omalizumab will be consider see basis if each individual the ion high dose ICS/LABA did not control symptoms ombination with other seen studied and coverage is not allowed except in trances (applies to both eosinophilic asthma populations)

Product Name: Nucala SC				
Diagnosis	Eosinophilic Granulomatosis with Polyangitis			
Approval Length	12/31/2039	12/31/2039		
Guideline Type Prior Authorization - All Plans except IL and MN Plans		s		
Product Generic Name		GPI	Brand/Generic	

NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION AUTO- INJECTOR 100 MG/ML	4460405500D530	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML	4460405500E520	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML	4460405500E530	Brand

4	Diognosio	۰f	agginanhilia	aranulamatasia	with	nalyanaitia
1	- Diagnosis	OI	eosinophilic	granulomatosis	willi	polyandius

AND

2 - Prescribed by, or in consultation with, a provider experienced in the treatment EGPA (i.e. Allergist, Pulmonologist, or Rheumatologist)

AND

3 - Member is 18 years of age or older

AND

- 4 One of the following:
- **4.1** All of the following:
- **4.1.1** Disease is one of the following:
 - Relapsed
 - Refractory

AND

- **4.1.2** All of the following:
- **4.1.2.1** Blood eosinophil level greater than or equal to 10% or an absolute eosinophil count greater than 1000 cells/μL with other causes ruled out (i.e. hypereosinophilic syndromes, neoplastic disease, or parasitic disease)

А	N	D

- **4.1.2.2** At least TWO of the following organ systems or features of EGPA disease:
- **4.1.2.2.1** Histopathological evidence of one of the following:
- eosinophilic vasculitis (i.e. bleeding under skin, red rash, petechiae, fibrinoid degeneration, blood clots)
- perivascular eosinophilic infiltration (i.e. inflammatory cells around blood vessels, lichenoid infiltration)
- eosinophil-rich granulomatosis inflammation (i.e. nodules, thick aggregation of histiocytes)

OR

4.1.2.2.2 Neuropathy (i.e. mono or polyneuropathy, mononeuritis multiplex)

OR

4.1.2.2.3 Pulmonary infiltrates (i.e. asthma, chronic pneumonia, hemoptysis, cough)

OR

4.1.2.2.4 Sino-nasal abnormality (i.e. sinusitis, allergic rhinitis, polyposis)

OR

4.1.2.2.5 Cardiomyopathy (i.e. heart failure, myocarditis, pericarditis, subendocardial fibrosis)

OR

4.1.2.2.6 Glomerulonephritis (i.e. hematuria, red cell casts, proteinuria)

OR

4.1.2.2.7 Alveolar hemorrhage (by bronchoalveolar lavage)

OR

4.1.2.2.8 Palpable purpura (i.e. skin nodules, urticarial rash, digital ischemia)

OR

4.1.2.2.9 Positive antineutrophil cytoplasmic antibody [ANCA]

AND

- **4.1.3** Trial and failure, intolerance, or contraindication to an adequate 3-month trial of both of the following:
 - prednisone
 - At least ONE additional immunosuppressive agent (i.e. cyclophosphamide, azathioprine, or methotrexate)

AND

4.1.4 Submission of medical records (e.g., chart notes) documenting baseline disease severity assessment with an objective measure/tool (i.e. chronic oral corticosteroid dose, number of intermittent steroid bursts, Birmingham Vasculitis Activity Score BVAS, number urgent care, emergency room visits or hospitalizations etc.)

OR

4.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Nucala SC

Diagnosis	Eosinophilic Granulomatosis with Polyangitis
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION AUTO-INJECTOR 100 MG/ML	4460405500D530	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML	4460405500E520	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML	4460405500E530	Brand

1 - Diagnosis of eosinophilic granulomatosis with polyangitis

AND

2 - Prescribed by, or in consultation with, a provider experienced in the treatment EGPA (i.e. Allergist, Pulmonologist, or Rheumatologist)

AND

3 - Member is 18 years of age or older

AND

- 4 One of the following:
 - **4.1** All of the following:
 - **4.1.1** Disease is one of the following:
 - Relapsed
 - Refractory

AND

- **4.1.2** All of the following:
- **4.1.2.1** Blood eosinophil level greater than or equal to 10% or an absolute eosinophil count greater than 1000 cells/µL with other causes ruled out (i.e. hypereosinophilic syndromes, neoplastic disease, or parasitic disease)

AND

- **4.1.2.2** At least TWO of the following organ systems or features of EGPA disease:
- **4.1.2.2.1** Histopathological evidence of one of the following:
- eosinophilic vasculitis (i.e. bleeding under skin, red rash, petechiae, fibrinoid degeneration, blood clots)
- perivascular eosinophilic infiltration (i.e. inflammatory cells around blood vessels, lichenoid infiltration)
- eosinophil-rich granulomatosis inflammation (i.e. nodules, thick aggregation of histiocytes)

OR

4.1.2.2.2 Neuropathy (i.e. mono or polyneuropathy, mononeuritis multiplex)

OR

4.1.2.2.3 Pulmonary infiltrates (i.e. asthma, chronic pneumonia, hemoptysis, cough)

OR

4.1.2.2.4 Sino-nasal abnormality (i.e. sinusitis, allergic rhinitis, polyposis)

OR

4.1.2.2.5 Cardiomyopathy (i.e. heart failure, myocarditis, pericarditis, subendocardial fibrosis)

OR

4.1.2.2.6 Glomerulonephritis (i.e. hematuria, red cell casts, proteinuria)

OR

4.1.2.2.7 Alveolar hemorrhage (by bronchoalveolar lavage)

OR

4.1.2.2.8 Palpable purpura (i.e. skin nodules, urticarial rash, digital ischemia)

OR

4.1.2.2.9 Positive antineutrophil cytoplasmic antibody [ANCA]

AND

- **4.1.3** Trial and failure, intolerance, or contraindication to an adequate 3-month trial of both of the following:
 - prednisone
 - At least ONE additional immunosuppressive agent (i.e. cyclophosphamide, azathioprine, or methotrexate)

AND

4.1.4 Submission of medical records (e.g., chart notes) documenting baseline disease severity assessment with an objective measure/tool (i.e. chronic oral corticosteroid dose, number of intermittent steroid bursts, Birmingham Vasculitis Activity Score BVAS, number urgent care, emergency room visits or hospitalizations etc.)

OR

4.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Nucala SC		
Diagnosis	Hypereosinophilic Syndrome	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION AUTO- INJECTOR 100 MG/ML	4460405500D530	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML	4460405500E520	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML	4460405500E530	Brand

Approval Criteria

1 - Diagnosis of hypereosinophilic syndrome for greater than or equal to 6 months without an identifiable non-hematologic secondary cause (i.e. cancer, imatinib-sensitive conditions, etc.)

AND

- **2** Prescribed by or in consultation with one of the following:
 - hematologist
 - allergist
 - other specialist in the treatment of Hypereosinophilic Syndrome

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Blood eosinophil count of greater than or equal to 1,000 cells/mc on at least two occasions

AND

3.1.2 Trial and failure, contraindication, or intolerance to the use of at least one steroid-sparing preventive treatments for at least 4 weeks (e.g., hydroxyurea, interferon-alfa, cyclosporine, etc.)

OR

3.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Nucala SC		
Diagnosis	Hypereosinophilic Syndrome	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION AUTO- INJECTOR 100 MG/ML	4460405500D530	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML	4460405500E520	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML	4460405500E530	Brand

Approval Criteria

1 - Diagnosis of hypereosinophilic syndrome for greater than or equal to 6 months without an identifiable non-hematologic secondary cause (i.e. cancer, imatinib-sensitive conditions, etc.)

AND

- 2 Prescribed by or in consultation with one of the following:
 - hematologist
 - allergist

• other specialist in the treatment of Hypereosinophilic Syndrome

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Blood eosinophil count of greater than or equal to 1,000 cells/mc on at least two occasions

AND

3.1.2 Trial and failure, contraindication, or intolerance to the use of at least one steroid-sparing preventive treatments for at least 4 weeks (e.g., hydroxyurea, interferon-alfa, cyclosporine, etc.)

OR

3.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Nucala SC		
Diagnosis	Nasal Polyps	
Approval Length	12/31/2039	
Guideline Type Prior Authorization - All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION AUTO-INJECTOR 100 MG/ML	4460405500D530	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML	4460405500E520	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML	4460405500E530	Brand

Approval Criteria

- 1 Diagnosis of chronic rhinosinusitis with nasal polyposis including all of the following:
 - At least eight weeks of moderate to severe nasal congestion/blockage/obstruction OR diminished sense of smell or rhinorrhea
 - Submission of medical records (e.g., chart notes) documenting nasal polyps by direct exam, endoscopy, or sinus CT scan (i.e. nasal polyp score five out of eight)
 - No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation

AND

2 - Prescribed by, or in consultation with a specialist experienced in the treatment of nasal polyps (i.e., otolaryngologist, allergist)

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** One of the following:
 - Trial and failure, contraindication, or intolerance to oral corticosteroids for nasal polyps
 - Prior to surgery for nasal polyps greater than six months ago
 - Trial and failure, contraindication, or intolerance to 2 or more nasal steroid sprays (i.e. failed two nasal sprays)
 - Trial and failure, contraindication, or intolerance to IM injections for polyps with one previous nasal spray

AND

3.1.2 Will be used in combination with a nasal corticosteroid medication

OR

3.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Nucala SC

Diagnosis	Nasal Polyps
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION AUTO- INJECTOR 100 MG/ML	4460405500D530	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 40 MG/0.4ML	4460405500E520	Brand
NUCALA	MEPOLIZUMAB SUBCUTANEOUS SOLUTION PREF SYRINGE 100 MG/ML	4460405500E530	Brand

- **1** Diagnosis of chronic rhinosinusitis with nasal polyposis including all of the following:
 - At least eight weeks of moderate to severe nasal congestion/blockage/obstruction OR diminished sense of smell or rhinorrhea
 - Submission of medical records (e.g., chart notes) documenting nasal polyps by direct exam, endoscopy, or sinus CT scan (i.e. nasal polyp score five out of eight)
 - No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation

AND

2 - Prescribed by, or in consultation with a specialist experienced in the treatment of nasal polyps (i.e., otolaryngologist, allergist)

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** One of the following:
 - Trial and failure, contraindication, or intolerance to oral corticosteroids for nasal polyps
 - Prior to surgery for nasal polyps greater than six months ago
 - Trial and failure, contraindication, or intolerance to 2 or more nasal steroid sprays (i.e. failed two nasal sprays)

• Trial and failure, contraindication, or intolerance to IM injections for polyps with one previous nasal spray

AND

3.1.2 Will be used in combination with a nasal corticosteroid medication

OR

3.2 Continuation of prior therapy with mepolizumab, verified by paid claims or medical records (e.g. chart notes)

Notes	**Continuation of case-by case-approved IgE inhibitor and IL-5 inhibit
	or, or tezepelumab combination therapy will only be considered if ICS/
	LABA therapy was also continued AND there was reduction in oral ste
	roid dose, exacerbations, or hospitalizations

2. Background

Benefit/Coverage/Program Information			
TABLE 1 - Outcome Measure values for uncontrolled asthma			
Measure	Not Well Controlled	Very Poorly Controlled	
Baseline symptoms (outside of exacerbation)	Greater than 2 days/week	Throughout the day	
Nighttime awakening	1-3 times/week	Greater than or equal to 4 times/week	
Interference with normal activity	Some limitation	Extremely limited	
Short acting beta agonist use for symptom control	Greater than 2 days/week	Several times per day	
FEV1	60-80% predicted or personal best	Less than 60% predicted or personal best	
Asthma exacerbations requiring oral steroids	Yes	Yes	

greater than or equal to 2 times in the past year		
Asthma Control Test (ACT)	16 - 19	Less than or equal to 15

3. Definitions

Definition	Description
Relapsing EGPA	At least one confirmed EGPA relapse while the person was on prednisolone dose of greater than or equal to 7.5 mg (or equivalent) within the past 2 years that required an increase in oral corticosteroid dose, initiation/increased immunosuppressive therapy dose, or hospitalization.
Refractory EGPA	1) Failure to attain remission (BVAS = 0 and oral steroid dose less than or equal to 7.5 mg/day prednisolone or equivalent) within the last 6 months following induction treatment with a standard regimen (e.g., cyclophosphamide, methotrexate, azathioprine, mycophenolate, high dose steroids) administered for at least 3 months OR 2) within 6 months prior to initiation, recurrence of symptoms of EGPA while tapering oral steroids, occurring at any dose level greater than or equal to 7.5 mg/day prednisolone or equivalent.
Failure of an immunosuppressant	Defined as EGPA symptoms are not resolving or flare occurring with a prednisone dose change, hospitalization, OR contraindications/clinical inappropriateness to immunosuppressants (i.e., liver disease, fertility etc.).

4. Revision History

Date	Notes
4/9/2024	Guideline Update

Nuplazid	(Pimavanserin Tartrate)	
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Prior Authorization Guideline

Guideline ID	GL-131415	
Guideline Name	Nuplazid (Pimavanserin Tartrate)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan who are being treated for Parkinson's disease psychosis and are established on therapy will have coverage under their drug benefit with documentation of symptom improvement or disease stability.

1. Criteria

Product Na	Product Name: Nuplazid			
Approval L	ength	12 month(s)		
Therapy St	age	Initial Authorization		
Guideline 7	Guideline Type Prior Authorization-IL and MN Plans Only			
Product Name	Generic Name		GPI	Brand/Generic
NUPLAZID	PIMAVANSERIN TARTRATE CAP 34 MG (BASE EQUIVALENT)		59400028200120	Brand

NUPLAZID PIMAVANSERIN TARTRATE TAB 10 MG (BASE EQUIVALENT)	59400028200310	Brand
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1 - Diagnosis of Parkinson's disease psychosis with documented hallucinations or delusions

AND

2 - Drug is prescribed by, or in consultation with, a Neurologist

AND

3 - Dose reductions and alterations in scheduling of dopaminergic agents led to increased Parkinson's symptoms

Product Name: Nuplazid		
Approval Length 12 month(s)		
Therapy Stage	Stage Reauthorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
NUPLAZID	PIMAVANSERIN TARTRATE CAP 34 MG (BASE EQUIVALENT)	59400028200120	Brand
NUPLAZID	PIMAVANSERIN TARTRATE TAB 10 MG (BASE EQUIVALENT)	59400028200310	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Nuplazid			
Approval Length	Approval Length 12/31/2039		
Therapy Stage	Initial Authorization		

Guideline Type		Prior Authorization-All plans except IL and MN		
Product Name	Generic Name		GPI	Brand/Generic
NUPLAZID	PIMAVANSERIN TARTRATE CAP 34 MG (BASE EQUIVALENT)		59400028200120	Brand
NUPLAZID	PIMAVANSERIN TARTRATE TAB 10 MG (BASE EQUIVALENT)		59400028200310	Brand

1 - Diagnosis of Parkinson's disease psychosis with documented hallucinations or delusions

AND

2 - Drug is prescribed by, or in consultation with, a Neurologist

AND

3 - Dose reductions and alterations in scheduling of dopaminergic agents led to increased Parkinson's symptoms

2. Revision History

Date	Notes
10/9/2023	New program

Nι	Nuzyra (omadacycline)					
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Prior Authorization Guideline

Guideline ID	GL-144546	
Guideline Name	uzyra (omadacycline)	
Formulary	Quartz	

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Nuzyra Tab				
Approval L	Length 1 Time Approval			
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans		ns		
Product Name	Generic Na	me	GPI	Brand/Generic
NUZYRA	OMADACYCLINE TOSYLATE TAB 150 MG (BASE EQUIVALENT)		04200050200320	Brand

Approval Criteria

- 1 One of the following:
- **1.1** Member has been receiving drug during hospitalization and needs to complete the course of therapy as an outpatient

OR

- **1.2** ALL of the following:
- **1.2.1** Submission of medical records (e.g., chart notes) documenting BOTH of the following:
 - Outpatient treatment of bacterial resistant strains
 - Report of susceptibilities resistant to preferred alternatives

AND

1.2.2 Prescribed by or in consultation with an Infectious Disease Specialist

Product Name: Nuzyra Tab	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL Plans*

Product Name	Generic Name	GPI	Brand/Generic
NUZYRA	OMADACYCLINE TOSYLATE TAB 150 MG (BASE EQUIVALENT)	04200050200320	Brand

Approval Criteria

- **1** One of the following:
- **1.1** Member has been receiving drug during hospitalization and needs to complete the course of therapy as an outpatient

OR

- **1.2** ALL of the following:
- **1.2.1** Submission of medical records (e.g., chart notes) documenting BOTH of the following:
 - Outpatient treatment of bacterial resistant strains

• Report of susceptibilities resistant to preferred alternatives

AND

1.2.2 Prescribed by or in consultation with an Infectious Disease Specialist

OR

1.3 The requested FDA approved drug is being used for the long-term treatment of tickborne disease

Notes	*Members who are established on therapy will have coverage under their drug benefit for the remainder of the current treatment course *Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored
	overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through reauthorization criteria

Product Name: Nuzyra Tab	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL Plans*

Product Name	Generic Name	GPI	Brand/Generic
NUZYRA	OMADACYCLINE TOSYLATE TAB 150 MG (BASE EQUIVALENT)	04200050200320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

AND

2 - Drug is being used for the long-term treatment of tick borne disease

Notes	*Members who are established on therapy will have coverage under t heir drug benefit for the remainder of the current treatment course
	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through reauthorization criteria

Product Name: Nuzyra Tab				
Approval Length 12 month(s)				
Guideline Type Prior Authorization - MN Plans*				
Product Name	Generic Na	me	GPI	Brand/Generic
NUZYRA	OMADACYCLINE TOSYLATE TAB 150 MG (BASE EQUIVALENT)		04200050200320	Brand

- **1** One of the following:
- **1.1** Member has been receiving drug during hospitalization and needs to complete the course of therapy as an outpatient

OR

- **1.2** ALL of the following:
- **1.2.1** Submission of medical records (e.g., chart notes) documenting BOTH of the following:
 - Outpatient treatment of bacterial resistant strains
 - Report of susceptibilities resistant to preferred alternatives

AND

1.2.2 Prescribed by or in consultation with an Infectious Disease Specialist

*Members who are established on therapy will have coverage under their drug benefit for the remainder of the
current treatment course

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through reauthorization criteria.
reauthorization criteria

2. Revision History

Date	Notes
3/18/2024	Product name updated

Ocaliva (obeticholic acid)		
The Board Integration for the flags, etc. The Board, was been been an account, washed, and detect to the fine to be put	ik to a contributation.	

Prior Authorization Guideline

Guideline ID	GL-131406	
Guideline Name	Ocaliva (obeticholic acid)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2023
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1. Criteria

Product Name: Ocaliva	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
OCALIVA	OBETICHOLIC ACID TAB 5 MG	52750060000320	Brand
OCALIVA	OBETICHOLIC ACID TAB 10 MG	52750060000330	Brand

Approval Criteria

1 - Diagnosis of primary biliary cholangitis

AND

2 - Alkaline phosphatase level > 1.6 times the upper limit of normal (ULN) OR total bilirubin between 1-2 times the ULN

AND

3 - Obeticholic acid will be used in combination with ursodeoxycholic acid (UDCA or ursodiol) OR there is trial and failure, contraindication or intolerance to ursodeoxycholic acid

AND

4 - Person does not have compensated cirrhosis with portal hypertension or a history of decompensated cirrhosis

Product Name: Ocaliva	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
OCALIVA	OBETICHOLIC ACID TAB 5 MG	52750060000320	Brand
OCALIVA	OBETICHOLIC ACID TAB 10 MG	52750060000330	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Ocaliva	
Approval Length 12/31/2039	
Guideline Type Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
OCALIVA	OBETICHOLIC ACID TAB 5 MG	52750060000320	Brand
OCALIVA	OBETICHOLIC ACID TAB 10 MG	52750060000330	Brand

1 - Diagnosis of primary biliary cholangitis

AND

2 - Alkaline phosphatase level > 1.6 times the upper limit of normal (ULN) OR total bilirubin between 1-2 times the ULN

AND

3 - Obeticholic acid will be used in combination with ursodeoxycholic acid (UDCA or ursodiol) OR there is trial and failure, contraindication or intolerance to ursodeoxycholic acid

AND

4 - Person does not have compensated cirrhosis with portal hypertension or a history of decompensated cirrhosis

2. Revision History

Date	Notes
10/9/2023	New program

Off Label Administrative			
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Prior Authorization Guideline

Guideline ID	GL-135255	
Guideline Name	Off Label Administrative	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: A drug used for an off-label indication or non-FDA approved indication			
Diagnosis	Off-label indication		
Approval Length	12 month(s)		
Guideline Type	Administrative		
Duradicat Name Comm	de Nieme	ODI	D

Product Name Generic Name	GPI	Brand/Generic
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Approval Criteria

- **1** ONE of the following:
- **1.1** Diagnosis is supported as a use in American Hospital Formulary Service Drug Information (AHFS DI)

O	R

1.2 Diagnosis is supported in the FDA Uses/Non-FDA Uses section in DRUGDEX Evaluation with a Strength of Recommendation rating of IIb or better (see DRUGDEX Strength of Recommendation table in Background section)

OR

1.3 Provider submits two articles from major peer reviewed medical journals that present data supporting the proposed off-label use or uses as generally safe and effective unless there is clear and convincing contradictory evidence presented in a major peer-reviewed medical journal

AND

- 2 ONE of the following:
- **2.1** Trial and failure, contraindication or intolerance to an adequate trial of all formulary and/or over the counter (OTC) alternatives

OR

2.2 (Minnesota plans only) person with stage four metastatic cancer and the requested drug is being used as supportive care to treat symptoms directly related to their cancer or chemotherapy regimen

OR

2.3 (Illinois Plans only) The requested FDA approved drug is being used for the long-term treatment of tick-borne disease

2. Background

Clinical Practice Guidelines

DRUGDEX Strength of Recommendation

Class	Recommendation	Description
Class I	Recommended	The given test or treatment has been proven useful, and should be performed or administered.
Class IIa	Recommended, In Most Cases	The given test or treatment is generally considered to be useful, and is indicated in most cases.
Class IIb	Recommended, in Some Cases	The given test or treatment may be useful, and is indicated in some, but not most, cases.
Class III	Not Recommended	The given test or treatment is not useful, and should be avoided
Class Indeterminate	Evidence Inconclusive	

NCCN Categories of Evidence and Consensus [A]

Category	Level of Consensus
1	Based upon high-level evidence, there is uniform NCCN consensus that the intervention is appropriate.
2A	Based upon lower-level evidence, there is uniform NCCN consensus that the intervention is appropriate.
2B	Based upon lower-level evidence, there is NCCN consensus that the intervention is appropriate.
3	Based upon any level of evidence, there is major NCCN disagreement that the intervention is appropriate.

Lexi-Drugs: Strength of Recommendation for Inclusion in Lexi-Drugs for Oncology Off-Label Use and Level of Evidence Scale for Oncology Off-Label Use [5] Strength of Recommendation for Inclusion

Strong (for proposed	The evidence persuasively supports the off-label use (ie, Level
off-label use)	of Evidence A).

Equivocal (for proposed off-label use)	The evidence to support the off-label use is of uncertain clinical significance (ie, Level of Evidence B, C). Additional studies may be necessary to further define the role of this medication for the off-label use.
Against proposed off- label use	The evidence either advocates against the off-label use or suggests a lack of support for the off-label use (independent of Level of Evidence). Additional studies are necessary to define the role of this medication for the off-label use.

Level of Evidence Scale for Oncology Off-Label Use

_	
A	Consistent evidence from well-performed randomized, controlled trials or overwhelming evidence of some other form (eg, results of the introduction of penicillin treatment) to support off-label use. Further research is unlikely to change confidence in the estimate of benefit.
В	Evidence from randomized, controlled trials with important limitations (eg, inconsistent results, methodologic flaws, indirect, imprecise); or very strong evidence of some other research design. Further research (if performed) is likely to have an impact on confidence in the estimate of benefit and risk and may change the estimate.
С	Evidence from observational studies (eg, retrospective case series/reports providing significant impact on patient care); unsystematic clinical experience; or potentially flawed randomized, controlled trials (eg, when limited options exist for condition). Any estimate of effect is uncertain.
G	Use has been substantiated by inclusion in at least one evidence-based or consensus-based clinical practice guideline.

3. Revision History

Date	Notes
12/8/2023	2024 New Implementation

(Omnipod Insulin Delivery System					
•	and the state of t	te de la companya de				

Prior Authorization Guideline

Guideline ID GL-139181		
Guideline Name	Omnipod Insulin Delivery System	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/19/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Omnipod Dash, Omnipod 5				
Approval Length		12 month(s)		
Therapy St	tage Initial Authorization			
Guideline 7	Гуре	Prior Authorization		
Product Name	Generic Name		GPI	Brand/Generic
OMNIPOD DASH	*INSULIN INFUSION DISPOSABLE PUMP RESERVOIR***		97201030506300	Brand

PODS (GEN 4)			
OMNIPOD 5 G6 PODS (GEN 5)	*INSULIN INFUSION DISPOSABLE PUMP RESERVOIR***	97201030506300	Brand
OMNIPOD 5 G6 INTRO KIT (GEN 5)	*INSULIN INFUSION DISPOSABLE PUMP KIT***	97201030506400	Brand

1 - Prescribed by or in consultation with an Endocrinologist or other provider with expertise in the management of diabetes (e.g., Certified Diabetic Educator [CDE])

AND

- 2 One of the following:
 - 2.1 Diagnosis of type 1 diabetes mellitus or other type of insulin-deficient diabetes

OR

- **2.2** Both of the following:
- 2.2.1 Diagnosis of gestational diabetes

AND

2.2.2 Member is on an intensive insulin therapy regimen of at least 3 insulin injections per day with frequent self-adjustments of insulin dose

OR

- **2.3** All of the following:
- 2.3.1 Diagnosis of type 2 diabetes mellitus

AND

2.3.2 Evidence of adherence to an intensive insulin therapy regimen with at least three insulin injections per day, requiring frequent self-adjustments of insulin for at least 6 months

AND

- **2.3.3** At least ONE of the following criteria while on the intensive insulin therapy regimen:
 - Hemoglobin A1c greater than 7%
 - Recurrent hypoglycemia (less than 70mg/dL)
 - Dawn phenomenon (recurrent morning FBG greater than 200 mg/dL)
 - History of severe glycemic excursions
 - Fluctuations in blood sugar before mealtimes

INOTES QL = 10 Cartridges per 30 days		Notes	QL = 10 cartridges per 30 days
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Product Name: Omnipod Dash, Omnipod 5	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
OMNIPOD DASH PODS (GEN 4)	*INSULIN INFUSION DISPOSABLE PUMP RESERVOIR***	97201030506300	Brand
OMNIPOD 5 G6 PODS (GEN 5)	*INSULIN INFUSION DISPOSABLE PUMP RESERVOIR***	97201030506300	Brand
OMNIPOD 5 G6 INTRO KIT (GEN 5)	*INSULIN INFUSION DISPOSABLE PUMP KIT***	97201030506400	Brand

Approval Criteria

1 - Member has been evaluated within the past 12 months by an Endocrinologist or other diabetes specialist

AND

2 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the pump

Notes	QL = 10 cartridges per 30 days
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2. Revision History

Date	Notes
1/18/2024	Update Guideline

Opioid Risk Management Program 7	Day Opioid First Fill Exception
(3) The antique and in the first the second could could could select a part to a combination.	

Guideline ID	GL-134592	
Guideline Name	Opioid Risk Management Program 7 Day Opioid First Fill Exception	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: all opioids, including opioid containing cold products			
Approval Length	14 Day(s)		
Guideline Type DUR - Reject 88: Excd 7DS, review CDC guidelines, use lowest effective dose and shortest duration at start. Submit O/R code.			
Product Name Gene	ric Name	GPI	Brand/Generic

Approval Criteria

- 1 One of the following:
 - Long-term care resident

 - Receiving hospice, palliative, or other end-of-life care Treatment of cancer-related pain or sickle cell-related pain

 Prescriber attests that the current prescription is a continuation of a stable, on-going opioid treatment regimen

Date	Notes
11/27/2023	New Program

Opioid Risk Management Program: Opioid Concurrent Use Ed					
(3) behavior and solidar from the source and a dead of solid property conductions					

Guideline ID	GL-134593
Guideline Name	Opioid Risk Management Program: Opioid Concurrent Use Edit
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: all opioids, including opioid containing cold products				
Diagnosis		Opioid Dependency Stopped		
Approval Length		12 month(s)		
Guideline Type		DUR - Reject 88: Buprenorphine Hx:Call MD,E Naloxone for safety.	nter O/	R. Co-prescribe
Product Name Gene		ric Name	GPI	Brand/Generic

Approval Criteria

1 - Prescriber attests that the person has stopped opioid dependency treatment with a buprenorphine containing drug and is resuming other opioid treatment

Product Name: all opioids, including opioid containing cold products		
Diagnosis	Opioid Dependency Continued	
Approval Length	1 fill (14 days)	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
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1 - Prescriber attests that the person is continuing opioid dependency treatment with a buprenorphine containing drug but requires acute opioid treatment

Date	Notes
11/27/2023	New Program

Opioid Risk Management: Opioid dose greater than 120 Morphine Milligram Equivalents (MMI				
	St. Challenge with higher higher hand was a least the late to the			

Guideline ID	GL-134594
	Opioid Risk Management: Opioid dose greater than 120 Morphine Milligram Equivalents (MME)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: all opioids, including opioid containing cold products				
Approval Length		12/31/2039		
Guideline Type DUR - Reject 88: MED 120mg Exceeded; Ttl MME MED <total calculated="" cumulative="" med="">MG TO O/R, ENTER PSS CODE OR MALL HD</total>				
Product Name	Gene	ric Name	GPI	Brand/Generic

Approval Criteria

- 1 One of the following:
 - Long-term care resident
 - Receiving hospice, palliative, or other end of life care

- Treatment of cancer-related pain
- Treatment of sickle cell-related pain

Product Name: all opioids, including opioid containing cold products	
Approval Length	12 month(s)
Guideline Type	DUR - Reject 88: MED 120mg Exceeded; Ttl MME MED <total calculated="" cumulative="" med="">MG TO O/R, ENTER PSS CODE OR MALL HD</total>

Product Name	Generic Name	GPI	Brand/Generic
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- **1** All of the following:
 - Prescriber states the opioid dose requested is medically necessary
 - Documentation that the state prescription drug monitoring program (PDMP) site has been checked in the past month
 - Documentation of a current pain contract
 - Documentation that use of naloxone has been discussed
 - Documentation of urine compliance screen in the previous 12 months

Product Name: a	Product Name: all opioids, including opioid containing cold products			
Approval Length		14 Day(s)		
Guideline Type		DUR - Reject 88: MED 120mg Exceeded; Ttl M calculated cumulative MED >MG TO O/R, ENT MALL HD		
Product Name	Gene	ic Name	GPI	Brand/Generic

Approval Criteria

1 - Person is changing medications and the new medication regimen does not exceed 120 MME

Product Name: all opioids, including opioid containing cold products	
Approval Length	3 month(s)

Guideline Type		DUR - Reject 88: MED 120mg Exceeded; Ttl M calculated cumulative MED >MG TO O/R, ENT MALL HD		
Product Name	Gener	ic Name	GPI	Brand/Generic

Product Name	Generic Name	GPI	Brand/Generic
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- 1 Member discharged from an inpatient stay after a severe, acute trauma with ALL of the following:
 - Prescriber states the opioid dose requested is medically necessary
 - Documentation that the state PDMP site has been checked prior to discharge
 - Documentation that use of naloxone has been discussed

OR

- 2 Both of the following:
 - 2.1 Person has 2 or more fills of greater than 120 MME within the previous 6 months

AND

2.2 Provider attests that continuation of therapy greater than 120 MME is medically necessary

Date	Notes
11/27/2023	New Program

Opzelura (ruxolitinib)
The based region community of the first term to contact a state and factor to provide an executive and the contract and the c

Guideline ID	GL-136714
Guideline Name	Opzelura (ruxolitinib)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Opzelura	
Diagnosis	Mild to moderate atopic dermatitis
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
OPZELURA	RUXOLITINIB PHOSPHATE CREAM 1.5%	90272060503720	Brand

Approval Criteria

1 - Diagnosis of mild to moderate atopic dermatitis

2 - Trial and failure of or contraindication to topical corticosteroid.

AND

3 - Trial and failure of or contraindication to calcineurin inhibitor (e.g. pimecrolimus, tacrolimus)

AND

4 - Therapy will not be used in combination with other therapeutic biologics (e.g., dupilumab, omalizumab, Upadacitinib)

Product Name: Opzelura	
Diagnosis	Vitiligo
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
OPZELURA	RUXOLITINIB PHOSPHATE CREAM 1.5%	90272060503720	Brand

Approval Criteria

1 - Diagnosis of nonsegmental vitiligo

AND

2 - Prescribed by, or in consultation with, a Dermatologist

3 - Area being treated does not exceed 10% body surface area (BSA)

AND

- 4 Person meets one of the following:
 - 4.1 Trial and failure of or contraindication to a medium-to-high potency topical corticosteroid

OR

4.2 Person is treating vitiligo affecting one of the following areas: face, skin folds, and/or genitalia

OR

4.3 Person has steroid-induced atrophy

OR

4.4 Person has a history of long-term topical steroid use

AND

5 - Therapy will not be used in combination with other therapeutic biologics (e.g., dupilumab, omalizumab, Upadacitinib)

Product Name: Opzelura	
Diagnosis	All diagnoses
Approval Length	12 month(s)
Therapy Stage	Reauthorization

Guideline T	уре	Prior Authorization		
Product Name	Generic Na	ime	GPI	Brand/Generic
OPZELURA	RUXOLITINIE	3 PHOSPHATE CREAM 1.5%	90272060503720	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug and with improvement in symptoms (e.g. reduction in body surface area affected, reduced itching, repigmentation.

Date	Notes
12/1/2023	2024 New Implementation

Oral Calcitonin Gene-Related Peptide (CGRP) Inhibitor
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Guideline ID	GL-144570
Guideline Name	Oral Calcitonin Gene-Related Peptide (CGRP) Inhibitors
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Nurtec ODT, Ubrelvy	
Diagnosis Acute Migraine Treatment	
Approval Length	12/31/2039
Guideline Type	Prior Authorization- ALL Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand
UBRELVY	UBROGEPANT TAB 50 MG	67701080000320	Brand
UBRELVY	UBROGEPANT TAB 100 MG	67701080000340	Brand

Approval Criteria

- **1** ONE of the following:
- **1.1** Trial and failure or intolerance to at least two of the following:
 - sumatriptan
 - naratriptan
 - rizatriptan
 - eletriptan
 - zolmitriptan
 - almotriptan
 - frovatriptan

OR

- **1.2** Both of the following:
- **1.2.1** Contraindication to triptan use

AND

1.2.2 Trial and failure, contraindication or intolerance to two non-triptan, prescription strength analgesics that are effective for treatment of migraines according to the American Headache Society treatment guidelines (e.g., nonsteroidal anti-inflammatory drugs (NSAIDs), ergotamine derivatives)

Product Name: Nurtec ODT, Ubrelvy	
Diagnosis	Acute Migraine Treatment
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization- IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand
UBRELVY	UBROGEPANT TAB 50 MG	67701080000320	Brand
UBRELVY	UBROGEPANT TAB 100 MG	67701080000340	Brand

- 1 One of the following:
- **1.1** Trial and failure or intolerance to at least two of the following:
 - sumatriptan
 - naratriptan
 - rizatriptan
 - eletriptan
 - zolmitriptan
 - almotriptan
 - frovatriptan

OR

- 1.2 Both of the following
- 1.2.1 Contraindication to triptan use

AND

1.2.2 Trial and failure, contraindication or intolerance to two non-triptan, prescription strength analgesics that are effective for treatment of migraines according to the American Headache Society treatment guidelines (e.g., nonsteroidal anti-inflammatory drugs (NSAIDs), ergotamine derivatives)

Product Name: Nurtec ODT, Ubrelvy	
Diagnosis	Acute Migraine Treatment
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization- IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand
UBRELVY	UBROGEPANT TAB 50 MG	67701080000320	Brand
UBRELVY	UBROGEPANT TAB 100 MG	67701080000340	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Nurtec ODT, Qulipta	
Diagnosis	Prevention of Migraine
Approval Length	12/31/2039
Guideline Type	Prior Authorization- ALL Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
QULIPTA	ATOGEPANT TAB 10 MG	67701010000310	Brand
QULIPTA	ATOGEPANT TAB 30 MG	67701010000320	Brand
QULIPTA	ATOGEPANT TAB 60 MG	67701010000330	Brand
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand

Approval Criteria

1 - Member has greater than or equal to 4 migraine days per month with submission of medical records (e.g., chart notes) documenting that headaches are disabling (e.g., unable to work/attend school, unable to participate in activities of daily living [ADLs], moderate to severe MIDAS score)

AND

2 - Member is 18 years of age or older

AND

3 - Trial and failure, contraindication or intolerance to two generic preventive migraine medications (e.g., anti-hypertensives, antiepileptics, antidepressants, botulinum toxin)

- 4 Trial and failure, contraindication or intolerance to both of the following:
 - Aimovig
 - Emgality

AND

5 - Drug is not being used in combination with another CGRP inhibitor for the preventative treatment of migraines

Product Name: Nurtec ODT, Qulipta	
Diagnosis	Prevention of Migraine
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization- IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
QULIPTA	ATOGEPANT TAB 10 MG	67701010000310	Brand
QULIPTA	ATOGEPANT TAB 30 MG	67701010000320	Brand
QULIPTA	ATOGEPANT TAB 60 MG	67701010000330	Brand
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand

Approval Criteria

1 - Member has greater than or equal to 4 migraine days per month with submission of medical records (e.g., chart notes) documenting that headaches are disabling (e.g., unable to work/attend school, unable to participate in activities of daily living [ADLs], moderate to severe MIDAS score)

AND

2 - Member is 18 years of age or older

AND

3 - Trial and failure, contraindication or intolerance to two generic preventive migraine medications (e.g., anti-hypertensives, antiepileptics, antidepressants, botulinum toxin)

AND

- 4 Trial and failure, contraindication or intolerance to both of the following:
 - Aimovig
 - Emgality

AND

5 - Drug is not being used in combination with another CGRP inhibitor for the preventative treatment of migraines

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through
reauthorization criteria

Product Name: Nurtec ODT, Qulipta	
Diagnosis	Prevention of Migraine
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization- IL and MN Plans*

Product Name	Generic Name	GPI	Brand/Generic
QULIPTA	ATOGEPANT TAB 10 MG	67701010000310	Brand
QULIPTA	ATOGEPANT TAB 30 MG	67701010000320	Brand
QULIPTA	ATOGEPANT TAB 60 MG	67701010000330	Brand
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand

UBRELVY	UBROGEPANT TAB 50 MG	67701080000320	Brand
UBRELVY	UBROGEPANT TAB 100 MG	67701080000340	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member (as evidenced by coverage effective date of less than or equal to 90 days) is showing a response to therapy (e.g., symptom improvement such as decreased frequency or severity of headaches from baseline, reduced cluster headache frequency, improved ability to participate in therapies/ADLs, improved MIDAS score, less acute medication use, fewer ER/UC visits for migraine, ability to return to work/school)

AND

2 - Drug is not being used in combination with another CGRP inhibitor for the preventative treatment of migraines

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through
reauthorization criteria

Product Name: Nurtec ODT, Ubrelvy		
Diagnosis Acute treatment – Quantity Exception		
Approval Length 12/31/2039		
Guideline Type Quantity Exception - ALL Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand
UBRELVY	UBROGEPANT TAB 50 MG	67701080000320	Brand
UBRELVY	UBROGEPANT TAB 100 MG	67701080000340	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that member has 2 or more headaches per week

2 - Patient is on migraine headache prophylaxis treatment

Product Name: Nurtec ODT, Ubrelvy			
Diagnosis	Diagnosis Acute treatment – Quantity Exception		
Approval Length	12 month(s)		
Guideline Type Quantity Exception - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
NURTEC	RIMEGEPANT SULFATE TAB DISINT 75 MG	67701060707220	Brand
UBRELVY	UBROGEPANT TAB 50 MG	67701080000320	Brand
UBRELVY	UBROGEPANT TAB 100 MG	67701080000340	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that member has 2 or more headaches per week

AND

2 - Patient is on migraine headache prophylaxis treatment

Date	Notes
3/18/2024	Updated authorization type

Orencia	(abatacept	:)	
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Guideline ID	GL-144881
Guideline Name	Orencia (abatacept)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/18/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Orencia SC				
Diagnosis		Psoriatic Arthritis (PsA)		
Approval Length 12/31/2039				
Guideline T	Guideline Type Prior Authorization – All Plans except IL and MN Plans			S
Product Name	Generic Name GPI		GPI	Brand/Generic
ORENCIA CLICKJECT	ABATACEPT SUBCUTANEOUS SOLN AUTO- INJECTOR 125 MG/ML		6640001000D520	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.4ML		6640001000E510	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED 6640001000E515 Brand SYRINGE 87.5 MG/0.7ML			Brand

ORENCIA ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 125 MG/ML	6640001000E520	Brand
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1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - · actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

AND

- **3.1.2** Trial and failure, contraindication, or intolerance to TWO of the following:
 - Adalimumab
 - Etanercept
 - Certolizumab
 - Golimumab
 - Risankizumab
 - Upadacitinib
 - Guselkumab
 - Tofacitinib/Tofacitinib XR
 - Ustekinumab

OR

3.2 Continuation of prior therapy with abatacept, verified by paid claims or medical records (e.g. chart notes)

Product Name: Orencia SC	
Diagnosis Psoriatic Arthritis (PsA)	
Approval Length	12 month(s)
Guideline Type Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ORENCIA CLICKJECT	ABATACEPT SUBCUTANEOUS SOLN AUTO- INJECTOR 125 MG/ML	6640001000D520	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.4ML	6640001000E510	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 87.5 MG/0.7ML	6640001000E515	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 125 MG/ML	6640001000E520	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- **3** One of the following:
 - **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:

- · actively inflamed joints
- axial disease
- active skin, nail, or scalp psoriasis involvement
- dactylitis
- enthesitis

- **3.1.2** Trial and failure, contraindication, or intolerance to TWO of the following:
 - Adalimumab
 - Etanercept
 - Certolizumab
 - Golimumab
 - Risankizumab
 - Upadacitinib
 - Guselkumab
 - Tofacitinib/Tofacitinib XR
 - Ustekinumab

OR

3.2 Continuation of prior therapy with abatacept, verified by paid claims or medical records (e.g. chart notes)

Product Name: Orencia SC	
Diagnosis Moderate to Severely Active Rheumatoid Arthritis (RA)	
Approval Length	12/31/2039
Guideline Type Prior Authorization – All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ORENCIA CLICKJECT	ABATACEPT SUBCUTANEOUS SOLN AUTO- INJECTOR 125 MG/ML	6640001000D520	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.4ML	6640001000E510	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 87.5 MG/0.7ML	6640001000E515	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 125 MG/ML	6640001000E520	Brand

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - Methotrexate (MTX)*
 - Leflunomide
 - Hydroxychloroquine
 - Sulfasalazine

AND

- **3.1.2** Trial and failure, contraindication, or intolerance to TWO of the following:
 - Adalimumab
 - Certolizumab
 - Etanercept
 - Golimumab
 - Tofactinib (ER)
 - Upadacitinib

OR

3.2 Continuation of prior therapy with abatacept, verified by paid claims or medical records (e.g. chart notes)

Notes	* Absolute contraindications to methotrexate are pregnancy, nursing,
	alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm
	bocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Orencia SC	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis (RA)
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ORENCIA CLICKJECT	ABATACEPT SUBCUTANEOUS SOLN AUTO- INJECTOR 125 MG/ML	6640001000D520	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.4ML	6640001000E510	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 87.5 MG/0.7ML	6640001000E515	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 125 MG/ML	6640001000E520	Brand

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - Methotrexate (MTX)*
 - Leflunomide

- Hydroxychloroquine
- Sulfasalazine

- 3.1.2 Trial and failure, contraindication, or intolerance to TWO of the following
 - Adalimumab
 - Certolizumab
 - Etanercept
 - Golimumab
 - Tofactinib (ER)
 - Upadacitinib

OR

3.2 Continuation of prior therapy with abatacept, verified by paid claims or medical records (e.g. chart notes)

Notes	* Absolute contraindications to methotrexate are pregnancy, nursing,
	alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm
	bocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Orencia SC	
Diagnosis Juvenile Idiopathic Arthritis (JIA)	
Approval Length	12/31/2039
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ORENCIA CLICKJECT	ABATACEPT SUBCUTANEOUS SOLN AUTO- INJECTOR 125 MG/ML	6640001000D520	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.4ML	6640001000E510	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 87.5 MG/0.7ML	6640001000E515	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 125 MG/ML	6640001000E520	Brand

Approval Criteria		
1 - Diagnosis of juvenile idiopathic arthritis		
	AND	
2 - Prescribed by or in o	consultation with a rheumatologist	
	AND	
3 - One of the following	:	
3.1 All of the following	:	
3.1.1 Minimum 3-mor following:	nth trial and failure, contraindication, or intolerance to ONE of the	
 Methotrexate (MTX)* Leflunomide Hydroxychloroquine Sulfasalazine 		
	AND	
3.1.2 Trial and failure	3.1.2 Trial and failure, contraindication, or intolerance to TWO of the following:	
 Adalimumab Etanercept Tofacitinib/Tofacitinib XR 		
OR		
3.2 Continuation of prior therapy with abatacept, verified by paid claims or medical records (e.g. chart notes)		
Notes	* Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant anemia, or hypersensitivity to methotrexate.	

Product Name: Orencia SC		
Diagnosis Juvenile Idiopathic Arthritis (JIA)		
Approval Length	12 month(s)	
Guideline Type	Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ORENCIA CLICKJECT	ABATACEPT SUBCUTANEOUS SOLN AUTO- INJECTOR 125 MG/ML	6640001000D520	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.4ML	6640001000E510	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 87.5 MG/0.7ML	6640001000E515	Brand
ORENCIA	ABATACEPT SUBCUTANEOUS SOLN PREFILLED SYRINGE 125 MG/ML	6640001000E520	Brand

1 - Diagnosis of juvenile idiopathic arthritis

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Minimum 3-month trial and failure, contraindication, or intolerance to ONE of the following:
 - Methotrexate (MTX)*
 - Leflunomide
 - Hydroxychloroquine
 - Sulfasalazine

AND

- **3.1.2** Trial and failure, contraindication, or intolerance to TWO of the following:
 - Adalimumab
 - Etanercept
 - Tofacitinib/Tofacitinib XR

OR

3.2 Continuation of prior therapy with abatacept, verified by paid claims or medical records (e.g. chart notes)

Notes	* Absolute contraindications to methotrexate are pregnancy, nursing,
	alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm
	bocytopenia or significant anemia, or hypersensitivity to methotrexate.

Date	Notes
4/9/2024	Updated Product name

(g) behaviour and mining an article most a size of the first half parties are the article and a size of the article and a	ORFAI	OIN (Niti	sinone)	, Nityr (Nitisin	one
	The first integrounce to disphyse. The	file may face have record, numerical, or deleted. Welly that it is fo	a punto a fine acrossifie and loadies.			

Guideline ID	GL-129653	
Guideline Name	ORFADIN (Nitisinone), Nityr (Nitisinone)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Orfadin solution, Generic Nitisinone, Brand Nityr		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
NITISINONE	NITISINONE CAP 2 MG	30904045000110	Generic
NITISINONE	NITISINONE CAP 5 MG	30904045000120	Generic
NITISINONE	NITISINONE CAP 10 MG	30904045000130	Generic
NITISINONE	NITISINONE CAP 20 MG	30904045000140	Generic
ORFADIN	NITISINONE SUSP 4 MG/ML	30904045001820	Brand
NITYR	NITISINONE TAB 2 MG	30904045000310	Brand

NITYR	NITISINONE TAB 5 MG	30904045000320	Brand
NITYR	NITISINONE TAB 10 MG	30904045000330	Brand

1 - Diagnosis of hereditary tyrosinemia type I.

AND

2 - Detectable succinylacetone blood or urine levels.

Product Name: Orfadin solution, Generic Nitisinone, Brand Nityr		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
NITISINONE	NITISINONE CAP 2 MG	30904045000110	Generic
NITISINONE	NITISINONE CAP 5 MG	30904045000120	Generic
NITISINONE	NITISINONE CAP 10 MG	30904045000130	Generic
NITISINONE	NITISINONE CAP 20 MG	30904045000140	Generic
ORFADIN	NITISINONE SUSP 4 MG/ML	30904045001820	Brand
NITYR	NITISINONE TAB 2 MG	30904045000310	Brand
NITYR	NITISINONE TAB 5 MG	30904045000320	Brand
NITYR	NITISINONE TAB 10 MG	30904045000330	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Orfadin solution, Generic Nitisinone, Brand Nityr

Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
NITISINONE	NITISINONE CAP 2 MG	30904045000110	Generic
NITISINONE	NITISINONE CAP 5 MG	30904045000120	Generic
NITISINONE	NITISINONE CAP 10 MG	30904045000130	Generic
NITISINONE	NITISINONE CAP 20 MG	30904045000140	Generic
ORFADIN	NITISINONE SUSP 4 MG/ML	30904045001820	Brand
NITYR	NITISINONE TAB 2 MG	30904045000310	Brand
NITYR	NITISINONE TAB 5 MG	30904045000320	Brand
NITYR	NITISINONE TAB 10 MG	30904045000330	Brand

1 - Diagnosis of hereditary tyrosinemia type I.

AND

2 - Detectable succinylacetone blood or urine levels.

Date	Notes
10/25/2023	New Program

Otezla (apremilast)		
The hand the growth to display to the long hand to stand, consider a stand to the long passes to consolidate industrial.		

Guideline ID	GL-144891
Guideline Name	Otezla (apremilast)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/18/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Otezla				
Diagnosis		Plaque Psoriasis		
Approval Length		12/31/2039		
Guideline Type		Prior Authorization – All Plans except IL and MN Plans		
Product Name	Generic Name GPI Brand/Gene		Brand/Generic	
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG 667000150		6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG		66700015000330	Brand

1 - Diagnosis of mild to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

OR

3.2 Continuation of prior therapy with apremilast, verified by paid claims or medical records (e.g. chart notes)

Product Name: Otezla	
Diagnosis Plaque Psoriasis	
Approval Length	12 month(s)
Guideline Type Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG	6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG	66700015000330	Brand

Approval Criteria

1 - Diagnosis of mild to severe plaque psoriasis

2 - Prescribed by or in consultation with a dermatologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

OR

3.2 Continuation of prior therapy with apremilast, verified by paid claims or medical records (e.g. chart notes)

Product Name: Otezla	
Diagnosis	Psoriatic Arthritis (PsA)
Approval Length	12/31/2039
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG	6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG	66700015000330	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

3.2 Continuation of prior therapy with apremilast, verified by paid claims or medical records (e.g. chart notes)

Product Name: Otezla	
Diagnosis Psoriatic Arthritis (PsA)	
Approval Length	12 month(s)
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG	6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG	66700015000330	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

3.2 Continuation of prior therapy with apremilast, verified by paid claims or medical records (e.g. chart notes)

Product Name: Otezla	
Diagnosis Oral Ulcers Associated with Behçet's Disease	
Approval Length	12/31/2039
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG	6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG	66700015000330	Brand

Approval Criteria

1 - Diagnosis of Behçet's Disease with active oral ulcers

2 - Prescribed by or in consultation with a rheumatologist

Product Name: Otezla	
Diagnosis Oral Ulcers Associated with Behçet's Disease	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization – IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG	6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG	66700015000330	Brand

Approval Criteria

1 - Diagnosis of Behçet's Disease with active oral ulcers

AND

2 - Prescribed by or in consultation with a rheumatologist

Product Name: Otezla	
Diagnosis Oral Ulcers Associated with Behçet's Disease	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization – IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG	6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG	66700015000330	Brand

- 1 One of the following:
- **1.1** Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days)

OR

1.2 Member has a previous prior authorization with the plan or historical prior authorization for apremilast on file

AND

2 - Prescriber provides clinical documentation from the previous 12 months of the member's response to therapy including individual improvements in functional status related to therapeutic response

Product Name: Otezla	
Diagnosis Oral Ulcers Associated with Behçet's Disease	
Approval Length	12/31/2039
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization – All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
OTEZLA	APREMILAST TAB STARTER THERAPY PACK 10 MG & 20 MG & 30 MG	6670001500B720	Brand
OTEZLA	APREMILAST TAB 30 MG	66700015000330	Brand

Approval Criteria

1 - Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days)

2 - Prescriber provides clinical documentation from the previous 12 months of the member's response to therapy including individual improvements in functional status related to therapeutic response

Date	Notes
4/15/2024	2024 New Implementation

Oxazolidinone Antibiotic				
g baharaga antinagkan baha ja basan and anat a mila ing bahas apakaha anakaranta				

Guideline ID	GL-144550
Guideline Name	Oxazolidinone Antibiotic
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

1. Criteria

Product Name: Sivextro Tab				
Approval Length		14 Day (s)*		
Guideline Type		Prior Authorization - All Plans except IL and MN		
Product Name	Generic Name GPI Brand/G		Brand/Generic	
SIVEXTRO	TEDIZOLID PHOSPHATE TAB 200 MG		16230070200320	Brand

- **1** One of the following:
- **1.1** Submission of medical records (e.g., chart notes, paid claims) supporting use of the drug during the current hospitalization and needs to complete the course of therapy as an outpatient

OR

- **1.2** All of the following:
- 1.2.1 Used for outpatient treatment of resistant bacterial strains

AND

1.2.2 Report of susceptibilities documenting resistance to alternatives including linezolid

AND

1.2.3 Prescribed by, or in consultation with, an Infectious Disease specialist

OR

- **1.3** Both of the following:
- 1.3.1 Linezolid is the only viable alternative due to resistance

AND

1.3.2 Member is taking serotonergic agents (e.g. SSRI, tricyclic antidepressants, triptans, etc.)

Notes	*Approval duration: Approve for the duration of treatment (usual cours
	e 6-14 days, or 14 to 28 days for Vancomycin-resistant enterococcus)

Product Name: Sivextro	o Tab
Approval Length	12 month(s)

Guideline 1	Гуре	Prior Authorization - IL Plan and MN	l Plans	
Product Name	Generic Na	me	GPI	Brand/Generic
SIVEXTRO	TEDIZOLID P	HOSPHATE TAB 200 MG	16230070200320	Brand

- **1** One of the following:
- **1.1** Submission of medical records (e.g., chart notes, paid claims) supporting use of the drug during the current hospitalization and needs to complete the course of therapy as an outpatient

OR

- **1.2** All of the following:
- **1.2.1** Used for outpatient treatment of resistant bacterial strains

AND

1.2.2 Report of susceptibilities documenting resistance to alternatives including linezolid

AND

1.2.3 Prescribed by, or in consultation with, an Infectious Disease specialist

OR

- **1.3** Both of the following:
- **1.3.1** Linezolid is the only viable alternative due to resistance

1.3.2 Member is taking serotonergic agents (e.g. SSRI, tricyclic antidepressants, triptans, etc.)

OR

1.4 For IL Plans ONLY: The requested drug is being used for the long-term treatment of tick-borne disease

Date	Notes
3/18/2024	Updated product name

Oxbryta (voxelotor)	
(g) bibliothings and higher half up bear mad would state tell balled parts to mental articles.	

Guideline ID	GL-130600
Guideline Name	Oxbryta (voxelotor)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Oxbryta				
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization		
Product Name	Generic Name		GPI	Brand/Generic
OXBRYTA	VOXELOTOR TAB 300 MG		82805080000310	Brand
OXBRYTA	VOXELOTOR TAB 500 MG 82805080000320 Brand		Brand	

OXBRYTA	OXBRYTA VOXELOTOR TAB FOR ORAL SUSP 300 MG 82805080007320 Brand					
Approval (Criteria					
1 - Both of	the following:					
	gnosis of sickle cell disease mber has persistent anemia requiring transfusion	within the past 12	! months			
	AND					
2 - Prescrik	ped by or in consultation with one of the following	:				
	natologist ecialist with experience in the treatment of sickle o	cell disease				
	AND					
3 - One of t	the following:					
• Sub	mber is stable on hydroxyurea for at least 90 days omission of medical records (e.g., chart notes) do traindication, or intolerance to hydroxyurea		d failure,			
	AND					
4 - Membe	r's baseline hemoglobin (Hgb) is between 5.5 to ′	10.5 g/dL prior to ι	use of Oxybryta			
	AND					

5 - Requested medication will not be used in combination with Adakveo (crizanlizumab)

samples, and/or vouchers.		*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.
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Product Name: Oxbryta	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
OXBRYTA	VOXELOTOR TAB 300 MG	82805080000310	Brand
OXBRYTA	VOXELOTOR TAB 500 MG	82805080000320	Brand
OXBRYTA	VOXELOTOR TAB FOR ORAL SUSP 300 MG	82805080007320	Brand

- **1** Submission of medical records (e.g., chart notes) documenting from the previous 12 months positive clinical response to therapy as evidenced by one of the following:
 - Decreased frequency of sickle cell hospitalizations or urgent care visits
 - Decreased frequency of vaso-occlusive crisis
 - Reduction in use of pain medications
 - Improved quality of life (e.g. decreased pain, fewer missed day of work/school, increase in activities, etc.)
 - Reduced need for transfusions

Notes	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was
	initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Date	Notes
10/6/2023	2024 New Implementation

Oxerva	Oxervate (cenegermin)			
The black image current be displayed. The fit	rwy han haer recent, wraned, or dilatel. Verly that he his yell	to the amerificaci instinc		

Guideline ID	GL-137246
Guideline Name	Oxervate (cenegermin)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Na	ame: Oxervate			
Approval Le	approval Length 8 Week(s)^			
Guideline T	Type Prior Authorization			
Product Name	Generic Name		GPI	Brand/Generic
OXERVATE	CENEGERMIN-BKBJ OPHTH SOLN 0.002% (20 MCG/ML)		86770020202020	Brand

Approval Criteria

1 - Confirmed diagnosis of Stage 2* or Stage 3* Neurotrophic Keratitis

AND

2 - Prescribed by, or in consultation with, an ophthalmologist

AND

3 - Submission of medical records (e.g., chart notes) confirming decreased or loss of corneal sensitivity and corneal epithelium changes

AND

4 - Underlying conditions are being treated, if appropriate (e.g., herpetic eye disease, diabetes, dry eye, multiple sclerosis, etc.)

AND

- **5** Failure to improve with conservative management after an adequate trial of one of the following for at least two weeks:
 - Ocular lubricants
 - Artificial tears

AND

6 - Discontinuation of ophthalmic steroids or avoidance of ophthalmic preservatives

Notes	*Stage 2 (Moderate) = NK exhibits nonhealing persistent epithelial def ect (PED); Stage 3 (Severe) = NK exhibits corneal ulceration involving subepithel ial (stromal) tissue which may progress to corneal
	perforation. ^ Maximum coverage is limited to 56 days per lifetime approval. Oxerv ate is hard-coded with a quantity limit of 56 days of therapy per lifetim e. Subsequent request will be reviewed using the off-label guideline

Date	Notes
12/6/2023	New program

Oxymorphone Hydrochloride		
The bill delicting source budging on The first transport from the county constant, or about the first first to price the countrils and states.		

Guideline ID	GL-129859
Guideline Name	Oxymorphone Hydrochloride
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Oxymorphone Immediate Release (IR), Oxymorphone Extended Release (ER)				
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization - IL Plan		
Product Name	Ge	neric Name GPI Brand/Generi		Brand/Generic
OXYMORPHONE HYDROCHLORIDE	ОХ	YMORPHONE HCL TAB 5 MG	65100080100305	Generic
OXYMORPHONE HYDROCHLORIDE	ОХ	YMORPHONE HCL TAB 10 MG	65100080100310	Generic
OXYMORPHONE HYDROCHLORIDE ER	ОХ	YMORPHONE HCL TAB ER 12HR 5 MG	65100080107405	Generic

OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 7.5 MG	65100080107407	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 10 MG	65100080107410	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 15 MG	65100080107415	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 20 MG	65100080107420	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 30 MG	65100080107430	Generic
OXYMORPHONE HYDROCHLORIDEER	OXYMORPHONE HCL TAB ER 12HR 40 MG	65100080107440	Generic

1 - For Oxymorphone IR requests ONLY: Trial and failure, contraindication, or intolerance to two generic immediate-release narcotics

OR

- **2** For Oxymorphone ER requests ONLY: Trial and failure, contraindication, or intolerance to BOTH of the following:
 - generic extended release morphine
 - extended release oxycodone

less rer-s will g	ember new to the plan (as evidenced by coverage effective date of than or equal to 90 days) who initiated therapy using a manufacture sponsored free drug program, provider samples, and/or vouchers go through initial criteria, otherwise for continuation of therapy for to plan, reauthorization criteria applies
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Product Name: Oxymorphone Immediate Release (IR), Oxymorphone Extended Release (ER)		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - MN Plan	

Product Name	Generic Name	GPI	Brand/Generic
OXYMORPHONE HYDROCHLORIDE	OXYMORPHONE HCL TAB 5 MG	65100080100305	Generic
OXYMORPHONE HYDROCHLORIDE	OXYMORPHONE HCL TAB 10 MG	65100080100310	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 5 MG	65100080107405	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 7.5 MG	65100080107407	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 10 MG	65100080107410	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 15 MG	65100080107415	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 20 MG	65100080107420	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 30 MG	65100080107430	Generic
OXYMORPHONE HYDROCHLORIDEER	OXYMORPHONE HCL TAB ER 12HR 40 MG	65100080107440	Generic

- 1 For Oxymorphone IR requests ONLY, One of the following:
- 1.1 Trial and failure, contraindication, or intolerance to two generic immediate-release narcotics

OR

- 1.2 Both of the following

 - Member has stage four metastatic cancer The requested drug is being used to treat cancer-related pain

OR

- **2** For Oxymorphone ER requests ONLY, one of the following:
- **2.1** Trial and failure, contraindication, or intolerance to BOTH of the following:
 - generic extended release morphine extended release oxycodone

OR

2.2 Both of the following:

- Member has stage four metastatic cancer
- The requested drug is being used to treat cancer-related pain

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for
	new to plan, reauthorization criteria applies

Product Name: Oxymorphone Immediate Release (IR), Oxymorphone Extended Release (ER)		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
OXYMORPHONE HYDROCHLORIDE	OXYMORPHONE HCL TAB 5 MG	65100080100305	Generic
OXYMORPHONE HYDROCHLORIDE	OXYMORPHONE HCL TAB 10 MG	65100080100310	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 5 MG	65100080107405	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 7.5 MG	65100080107407	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 10 MG	65100080107410	Generic

OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 15 MG	65100080107415	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 20 MG	65100080107420	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 30 MG	65100080107430	Generic
OXYMORPHONE HYDROCHLORIDEER	OXYMORPHONE HCL TAB ER 12HR 40 MG	65100080107440	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, regulatorization criteria applies
	new to plan, reauthorization criteria applies

Product Name: Oxymorphone Immediate Release (IR), Oxymorphone Extended Release (ER)		
Approval Length	12/31/2039	
Guideline Type Prior Authorization - All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
OXYMORPHONE HYDROCHLORIDE	OXYMORPHONE HCL TAB 5 MG	65100080100305	Generic
OXYMORPHONE HYDROCHLORIDE	OXYMORPHONE HCL TAB 10 MG	65100080100310	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 5 MG	65100080107405	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 7.5 MG	65100080107407	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 10 MG	65100080107410	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 15 MG	65100080107415	Generic

OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 20 MG	65100080107420	Generic
OXYMORPHONE HYDROCHLORIDE ER	OXYMORPHONE HCL TAB ER 12HR 30 MG	65100080107430	Generic
OXYMORPHONE HYDROCHLORIDEER	OXYMORPHONE HCL TAB ER 12HR 40 MG	65100080107440	Generic

1 - For Oxymorphone IR requests ONLY: Trial and failure, contraindication, or intolerance to two generic immediate-release narcotics

OR

- **2** For Oxymorphone ER requests ONLY: Trial and failure, contraindication, or intolerance to BOTH of the following:
 - generic extended release morphine
 - extended release oxycodone

Date	Notes
8/14/2023	2024 New Implementation

Palforzia (peanut powder)			
(2) Section frequency strategy for the section sense and account of their last parties to contribute states.			

Guideline ID	GL-129373
Guideline Name	Palforzia (peanut powder)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Palforzia		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization - ALL Plans*		

Product Name	Generic Name	GPI	Brand/Generic
PALFORZIA INITIAL DOSE ESCALATION	PEANUT POWDER-DNFP STARTER PACK 0.5 & 1 & 1.5 & 3 & 6 MG	2010004020H510	Brand
PALFORZIA LEVEL 1	PEANUT POWDER-DNFP CAP SPRINKLE PACK 3 X 1 MG (3 MG DOSE)	2010004020H525	Brand
PALFORZIA LEVEL 2	PEANUT POWDER-DNFP CAP SPRINKLE PACK 6 X 1 MG (6 MG DOSE)	2010004020H530	Brand
PALFORZIA LEVEL 3	PEANUT POWDER-DNFP PACK 2 X 1 MG & 10 MG (12 MG DOSE)	2010004020H535	Brand

PALFORZIA LEVEL 4	PEANUT POWDER-DNFP CAP SPRINKLE PACK 20 MG (20 MG DOSE)	2010004020H540	Brand
PALFORZIA LEVEL 5	PEANUT POWDER-DNFP CAP SPRINKLE PACK 2 X 20 MG (40 MG DOSE)	2010004020H545	Brand
PALFORZIA LEVEL 6	PEANUT POWDER-DNFP CAP SPRINKLE PACK 4 X 20 MG (80 MG DOSE)	2010004020H550	Brand
PALFORZIA LEVEL 7	PEANUT POWDER-DNFP PACK 20 MG & 100 MG (120 MG DOSE)	2010004020H555	Brand
PALFORZIA LEVEL 8	PEANUT POWDER-DNFP PACK 3 X 20 MG & 100 MG (160 MG DOSE)	2010004020H560	Brand
PALFORZIA LEVEL 9	PEANUT POWDER-DNFP PACK 2 X 100 MG (200 MG DOSE)	2010004020H565	Brand
PALFORZIA LEVEL 10	PEANUT POWDER-DNFP PACK 2 X 20 MG & 2 X 100 MG (240 MG DOSE)	2010004020H570	Brand
PALFORZIA LEVEL 11 (TITRATION)	PEANUT ALLERGEN POWDER-DNFP TITRATION PACKET 300 MG	20100040203030	Brand
PALFORZIA LEVEL 11 (MAINTENANCE)	PEANUT ALLERGEN POWDER-DNFP MAINTENANCE PACKET 300 MG	20100040203050	Brand

1 - Submission of medical records (e.g., chart notes) documenting systemic allergic reaction to peanuts (e.g., anaphylaxis, tongue/throat swelling, shortness of breath/wheezing the requires treatment, urticaria, angioedema, hypotension, and/or vomiting that occurs within 1-2 hours after ingestion of peanut)

AND

2 - Submission of medical records (e.g., chart notes) documenting a positive skin prick test (wheal diameter greater than or equal to 3 mm) OR peanut specific IgE (greater than or equal to 0.35 kUA/L) within the past 12 months

AND

3 - Used in conjunction with a peanut-avoidance diet

4 - Patient is 4 years of age or older, to less than or equal to 17 years of age

AND

5 - Prescribed by or in consultation with an allergist/immunologist

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through reauthorization criteria
reauthorization criteria

Product Name: Palforzia		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - ALL Plans*	

Product Name	Generic Name	GPI	Brand/Generic
PALFORZIA INITIAL DOSE ESCALATION	PEANUT POWDER-DNFP STARTER PACK 0.5 & 1 & 1.5 & 3 & 6 MG	2010004020H510	Brand
PALFORZIA LEVEL 1	PEANUT POWDER-DNFP CAP SPRINKLE PACK 3 X 1 MG (3 MG DOSE)	2010004020H525	Brand
PALFORZIA LEVEL 2	PEANUT POWDER-DNFP CAP SPRINKLE PACK 6 X 1 MG (6 MG DOSE)	2010004020H530	Brand
PALFORZIA LEVEL 3	PEANUT POWDER-DNFP PACK 2 X 1 MG & 10 MG (12 MG DOSE)	2010004020H535	Brand
PALFORZIA LEVEL 4	PEANUT POWDER-DNFP CAP SPRINKLE PACK 20 MG (20 MG DOSE)	2010004020H540	Brand
PALFORZIA LEVEL 5	PEANUT POWDER-DNFP CAP SPRINKLE PACK 2 X 20 MG (40 MG DOSE)	2010004020H545	Brand
PALFORZIA LEVEL 6	PEANUT POWDER-DNFP CAP SPRINKLE PACK 4 X 20 MG (80 MG DOSE)	2010004020H550	Brand
PALFORZIA LEVEL 7	PEANUT POWDER-DNFP PACK 20 MG & 100 MG (120 MG DOSE)	2010004020H555	Brand
PALFORZIA LEVEL 8	PEANUT POWDER-DNFP PACK 3 X 20 MG & 100 MG (160 MG DOSE)	2010004020H560	Brand
PALFORZIA LEVEL 9	PEANUT POWDER-DNFP PACK 2 X 100 MG (200 MG DOSE)	2010004020H565	Brand
PALFORZIA LEVEL 10	PEANUT POWDER-DNFP PACK 2 X 20 MG & 2 X 100 MG (240 MG DOSE)	2010004020H570	Brand

PALFORZIA LEVEL 11 (TITRATION)	PEANUT ALLERGEN POWDER-DNFP TITRATION PACKET 300 MG	20100040203030	Brand
PALFORZIA LEVEL 11 (MAINTENANCE)	PEANUT ALLERGEN POWDER-DNFP MAINTENANCE PACKET 300 MG	20100040203050	Brand

- 1 Submission of medical records (e.g., chart notes) documenting one of the following:
 - Member has a persistent peanut allergy as documented in an allergy/immunology clinic visit within the past 12 months
 - Member has a documented positive skin prick test (wheal diameter greater than or equal to 3 mm) or peanut specific IgE (greater than or equal to 0.35 kUA/L) within the past 12 months

AND

2 - Used in conjunction with a peanut-avoidance diet

AND

3 - Prescribed by or in consultation with an allergist/immunologist

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through
reauthorization criteria

Date	Notes
8/4/2023	2024 New Implementation

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Guideline ID	GL-138053
Guideline Name Palynziq	
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Palynziq (10 and 20 mg dose)				
Approval Length		4 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization - All plans except IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
PALYNZIQ PEGVALIASE-PQPZ SUBCUTANEOUS SOLN SYRINGE 2.5 MG/0.5ML			3090855040E510	Brand

PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 10 MG/0.5ML	3090855040E520	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 20 MG/ML	3090855040E530	Brand

1 - Diagnosis of Phenylketonuria (PKU)

AND

2 - Member is 18 years of age or older

AND

- **3** Blood phenylalanine (Phe) concentration greater than 600 micromol/L (10 mg/dL) despite both of the following:
 - Six months of adherent use of a Phe restricted diet
 - Two-month trial and failure, contraindication, or intolerance of sapropterin

AND

4 - Sapropterin must be discontinued prior to start of Palynziq

Product Name: Palynziq (10 and 20 mg dose)		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 2.5 MG/0.5ML	3090855040E510	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 10 MG/0.5ML	3090855040E520	Brand

· ·	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 20 MG/ML	3090855040E530	Brand
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1 - Diagnosis of Phenylketonuria (PKU)

AND

2 - Member is 18 years of age or older

AND

- **3** Blood phenylalanine (Phe) concentration greater than 600 micromol/L (10 mg/dL) despite both of the following:
 - Six months of adherent use of a Phe restricted diet
 - Two-month trial and failure, contraindication, or intolerance of sapropterin

AND

4 - Sapropterin must be discontinued prior to start of Palynziq

Product Name: Palynziq (40 mg dose)		
Approval Length 4 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Quantity Limit - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 2.5 MG/0.5ML	3090855040E510	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 10 MG/0.5ML	3090855040E520	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 20 MG/ML	3090855040E530	Brand

1 - Diagnosis of Phenylketonuria (PKU)

AND

2 - Member is 18 years of age or older

AND

- **3** Blood phenylalanine (Phe) concentration greater than 600 micromol/L (10 mg/dL) despite both of the following:
 - Six months of adherent use of a Phe restricted diet
 - Two-month trial and failure, contraindication, or intolerance of sapropterin

AND

4 - Sapropterin must be discontinued prior to start of Palynziq

- **5** One of the following:
 - 24-week trial of 20 mg/day dosing and failure to achieve a 20% reduction from baseline in Phe levels
 - Phe levels remain greater than 600 micromol/L

Product Name: Palynziq (40 mg dose)		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Quantity Limit- IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 2.5 MG/0.5ML	3090855040E510	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 10 MG/0.5ML	3090855040E520	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 20 MG/ML	3090855040E530	Brand

1 - Diagnosis of Phenylketonuria (PKU)

AND

2 - Member is 18 years of age or older

AND

- **3** Blood phenylalanine (Phe) concentration greater than 600 micromol/L (10 mg/dL) despite both of the following:
 - Six months of adherent use of a Phe restricted diet
 - Two-month trial and failure, contraindication, or intolerance of sapropterin

AND

4 - Sapropterin must be discontinued prior to start of Palynziq

- 5 One of the following:
 - 24-week trial of 20 mg/day dosing and failure to achieve a 20% reduction from baseline in Phe levels
 - Phe levels remain greater than 600 micromol/L

Product Name: Palynziq	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 2.5 MG/0.5ML	3090855040E510	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 10 MG/0.5ML	3090855040E520	Brand
PALYNZIQ	PEGVALIASE-PQPZ SUBCUTANEOUS SOLN PREF SYRINGE 20 MG/ML	3090855040E530	Brand

1 - Used in conjunction with a Phe restricted diet

AND

- **2** Submission of medical records (e.g., chart notes) documenting ONE of the following:
 - 20% reduction in Phe levels from baseline
 - Phe levels remain greater than 600 micromol/L

AND

3 - Not on concurrent sapropterin

Date	Notes
12/20/2023	Update

Parathyroid Hormone Analogues for Osteoporosis		

Guideline ID	GL-144692	
Guideline Name	Parathyroid Hormone Analogues for Osteoporosis	
Formulary	Quartz	

Guideline Note:

Effective Date:	4/21/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

1. Criteria

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos				
Diagnosis		Osteoporosis in Postmenopausal Women		
Approval Leng	th	24 month(s)		
Therapy Stage)	Initial Authorization		
Guideline Type	9	Prior Authorization - All Plans except IL and MN Plans		
Product Name	Generic	: Name	GPI	Brand/Generic

TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

3 - Diagnosis of osteoporosis with a T-score of less than or equal to -2.5 at the femoral neck, total hip, lumbar spine, or 33% (one-third) radius

AND

- 4 Very high risk of fracture defined by AT LEAST ONE of the following:
 - Recent fracture (e.g. within past 12 months)
 - Fracture while on approved osteoporosis therapy
 - Multiple fractures
 - Fractures while on drugs causing skeletal harm (e.g. long-term glucocorticoid use)
 - Very low T-score (less than -3.0)
 - High risk for falls
 - History of injurious falls

AND

5 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos		
Diagnosis	Osteoporosis in Postmenopausal Women	
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

3 - Diagnosis of osteoporosis with a T-score of less than or equal to -2.5 at the femoral neck, total hip, lumbar spine, or 33% (one-third) radius

- 4 Very high risk of fracture defined by AT LEAST ONE of the following:
 - Recent fracture (e.g. within past 12 months)
 - Fracture while on approved osteoporosis therapy
 - Multiple fractures
 - Fractures while on drugs causing skeletal harm (e.g. long-term glucocorticoid use)
 - Very low T-score (less than -3.0)
 - High risk for falls

• History of injurious falls

AND

5 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos				
Diagnosis	Osteopenia in Postmenopausal Women			
Approval Length	24 month(s)			
Therapy Stage	Initial Authorization			
Guideline Type	Prior Authorization - All Plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

Approval Criteria

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

3 - Diagnosis of osteopenia with a T-score between -1.0 and -2.5 at the femoral neck, total hip, lumbar spine, or 33% (one-third) radius

AND

4 - 10-year probability of a hip fracture of at least 3% or major osteoporosis-related fracture of at least 20%

AND

- **5** Very high risk of fracture defined by AT LEAST ONE of the following:
 - Recent fracture (e.g. within past 12 months)
 - Fracture while on approved osteoporosis therapy
 - Multiple fractures
 - Fractures while on drugs causing skeletal harm (e.g. long-term glucocorticoid use)
 - Very high FRAX (major osteoporotic fracture > 30%, hip fracture > 4.5%)
 - High risk for falls
 - History of injurious falls

AND

6 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos				
Diagnosis	Osteopenia in Postmenopausal Women			
Approval Length	12 month(s)			
Therapy Stage	Initial Authorization			
Guideline Type	Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

3 - Diagnosis of osteopenia with a T-score between -1.0 and -2.5 at the femoral neck, total hip, lumbar spine, or 33% (one-third) radius

AND

4 - 10-year probability of a hip fracture of at least 3% or major osteoporosis-related fracture of at least 20%

AND

- 5 Very high risk of fracture defined by AT LEAST ONE of the following:
 - Recent fracture (e.g. within past 12 months)
 - Fracture while on approved osteoporosis therapy
 - Multiple fractures
 - Fractures while on drugs causing skeletal harm (e.g. long-term glucocorticoid use)
 - Very high FRAX (major osteoporotic fracture > 30%, hip fracture > 4.5%)
 - High risk for falls
 - History of injurious falls

AND

6 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos			
Diagnosis	Osteoporosis Due to Prolonged Steroid Use		
Approval Length	24 month(s)		
Therapy Stage	erapy Stage Initial Authorization		
Guideline Type Prior Authorization - All Plans except IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

3 - Trial and failure (e.g. low-trauma fractures or decrease in BMD while on alendronate 70 mg per week), contraindication, or intolerance to therapy with at least one bisphosphonate

AND

4 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos		
Diagnosis Osteoporosis Due to Prolonged Steroid Use		
Approval Length 12 month(s)		

Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

3 - Trial and failure (e.g. low-trauma fractures or decrease in BMD while on alendronate 70 mg per week), contraindication, or intolerance to therapy with at least one bisphosphonate

AND

4 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos		
Diagnosis	Primary or Hypogonadal Osteoporosis in Men	
Approval Length	24 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

- 3 One of the following:
- **3.1** Trial and failure (e.g. low-trauma fractures or decrease in BMD while on alendronate 70 mg per week), contraindication, or intolerance to therapy with at least one bisphosphonate

OR

3.2 T-score of less than -2.5 and at least one fragility fracture

AND

4 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos			
Diagnosis Primary or Hypogonadal Osteoporosis in Men			
Approval Length 12 month(s)			

Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Medication will be self-administered or administered by a family member or friend

AND

2 - Will not be used in combination with anti-resorptive therapy or after denosumab therapy

AND

- 3 One of the following:
- **3.1** Trial and failure (e.g. low-trauma fractures or decrease in BMD while on alendronate 70 mg per week), contraindication, or intolerance to therapy with at least one bisphosphonate

OR

3.2 T-score of less than -2.5 and at least one fragility fracture

AND

4 - Applies to Teriparatide only: Trial and failure, contraindication, or intolerance to abaloparatide

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos

Diagnosis	All Indications
Approval Length	24 Month(s)*
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) or who are established on therapy, will have coverage under their drug benefit for the remainder of the current treatment course (up to 24 months total). Restrictions to specific network pharmacies and participation in medication management programs may apply.

Notes	*Maximum coverage is limited to a 24 months per lifetime approval. T
	eriparatide and Tymlos are hard-coded with a quantity limit of 24 mont
	hs of therapy per lifetime. Subsequent request will be reviewed using t
	he off-label guideline.

Product Name: Brand Teriparatide Generic Teriparatide, Tymlos	
Diagnosis	All Indications
Approval Length	12 Month(s)*
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TYMLOS	ABALOPARATIDE SUBCUTANEOUS SOLN PEN- INJECTOR 3120 MCG/1.56ML	3004400500D230	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 620 MCG/2.48ML	3004407000D221	Brand
TERIPARATIDE	TERIPARATIDE (RECOMBINANT) SOLN PEN-INJ 600 MCG/2.4ML	3004407000D220	Generic

1 - Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) or who are established on therapy, will have coverage under their drug benefit for the remainder of the current treatment course (up to 24 months total). Restrictions to specific network pharmacies and participation in medication management programs may apply.

*Maximum coverage is limited to 24 months per lifetime approval. Teri
paratide and Tymlos are hard-coded with a quantity limit of 24 months
of therapy per lifetime. Subsequent request will be reviewed using the
off-label guideline.

Date	Notes
3/20/2024	Updated guideline

Peghigrastim
The Mandel Anger count to displayers. Therefore, record, research, and defined in the Bankla kit profits the accounting and trades.

Guideline ID	GL-129860
Guideline Name	Pegfilgrastim
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Fulphila, Fylnetra, Nyvepria, Stimufend, Udenyca, Ziextenzo				
Approval Length		12 month(s)		
Guideline Type		Prior Authorization		
Product Name	Generic Name		GPI	Brand/Generic
FULPHILA	PEGFILGRASTIM-JMDB SOLN PREFILLED SYRINGE 6 MG/0.6ML		8240157020E520	Brand
FYLNETRA	PEGFILGRASTIM-PBBK SOLN PREFILLED SYRINGE 6 MG/0.6ML		8240157060E520	Brand
NYVEPRIA	PEGFILGRASTIM-APGF SOLN PREFILLED SYRINGE 6 MG/0.6ML		8240157002E520	Brand
STIMUFEND	PEGFILGRASTIM-FPGK SOLN PREFILLED SYRINGE 6 MG/0.6ML		8240157015E520	Brand
UDENYCA	A PEGFILGRASTIM-CBQV SOLN AUTO-INJECTOR 6 8240157010D520 Brand MG/0.6ML		Brand	

UDENYCA	PEGFILGRASTIM-CBQV SOLN PREFILLED SYRINGE 6 MG/0.6ML	8240157010E520	Brand
ZIEXTENZO	PEGFILGRASTIM-BMEZ SOLN PREFILLED SYRINGE 6 MG/0.6ML	8240157005E520	Brand

- 1 One of the following:
- **1.1** Both of the following:
- **1.1.1** Trial and failure (e.g., febrile neutropenia, delay in chemotherapy), contraindication, or intolerance to a filgrastim drug product

AND

1.1.2 Trial and failure, contraindication, or intolerance to use of Ziextenzo in the clinic as a clinic administered drug

OR

- 1.2 Both of the following (Applies to Minnesota Plans ONLY):
 - Member has stage four metastatic cancer
 - The requested drug is being used as supportive care for their cancer treatment

Notes	*Pharmacy benefit coverage information (preferred/nonpreferred statu
	s, restriction, etc) only applies to plans with
	Quartz pharmacy coverage

Date	Notes
10/12/2023	2024 New Implementation

Pe	Pegylated Interferons					
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Guideline ID	GL-129861
Guideline Name	Pegylated Interferons
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Pegasys				
Approval Length Approval Durations: Hepatitis = 48 weeks. Other indications = 12 months				
Therapy Stage	Initial Authorization			
Guideline Type	Prior Authorization			
Draduat Canari	- Name			

Product Name	Generic Name	GPI	Brand/Generic
PEGASYS	PEGINTERFERON ALFA-2A SOLN PREFILLED SYR 180 MCG/0.5ML	1235306005E540	Brand
PEGASYS	PEGINTERFERON ALFA-2A INJ 180 MCG/ML	12353060052020	Brand

Approval Criteria

of the following:
chronic hepatitis B chronic hepatitis B
AND
npensated liver disease
AND
of the following:
on
OR
g is being used alone or in a combination regimen that has a class 1 ruse from the National Comprehensive Cancer Network (NCCN) in the member*
AND
e self-administered by member e administered by a family member
*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was nitiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Name: Pegasys			
Approval Length Approval Durations: Hepatitis = 48 weeks. Other indications months			
Therapy Stage	Reauthorization		
Guideline Type	Prior Authorization		

Product Name	Generic Name	GPI	Brand/Generic
PEGASYS	PEGINTERFERON ALFA-2A SOLN PREFILLED SYR 180 MCG/0.5ML	1235306005E540	Brand
PEGASYS	PEGINTERFERON ALFA-2A INJ 180 MCG/ML	12353060052020	Brand

1	- (Or	ıe	of	the	fol	low	ing	:

- **1.1** All of the following:
- **1.1.1** Diagnosis of one of the following:
 - HBeAg positive chronic hepatitis B
 - HBeAg negative chronic hepatitis B

AND

1.1.2 Member has compensated liver disease

AND

- **1.1.3** Evidence of both of the following:
 - Viral replication
 - Liver inflammation

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

AND

- 2 One of the following;
 - Medication will be self-administered by member
 - Medication will be administered by a family member

AND

3 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months, response to therapy is stable or improvement seen on therapy with evidence-based clinical rationale to support continuing therapy

AND

4 - Restrictions to specific network pharmacies and participation in medication management programs may apply

Notes	*Continuation of therapy/coverage criteria will not be applied to person
	s who were not previously approved for coverage whose therapy was
	initiated using a manufacturer-sponsored free drug program, provider
	samples, and/or vouchers.

Date	Notes
8/14/2023	2024 New Implementation

ŀ	Pradaxa Oral Pellets						
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Guideline ID	GL-129132	
Guideline Name	Pradaxa Oral Pellets	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Pradaxa Oral Pellets	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 20 MG	83337030203020	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 30 MG	83337030203025	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 40 MG	83337030203030	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 50 MG	83337030203035	Brand

PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 110 MG	83337030203040	Brand
PRADAXA	PRADAXA DABIGATRAN ETEXILATE MESYLATE PELLET PACK 150 MG		Brand

1 - Trial and failure, contraindication, or intolerance to rivaroxaban suspension

Product Name: Pradaxa Oral Pellets		
Approval Length	Approval Length 12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Step Therapy - IL or MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 20 MG	83337030203020	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 30 MG	83337030203025	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 40 MG	83337030203030	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 50 MG	83337030203035	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 110 MG	83337030203040	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 150 MG	83337030203045	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Pradaxa Oral Pellets	
Approval Length 12/31/2039	
Guideline Type Step Therapy - All Plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 20 MG	83337030203020	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 30 MG	83337030203025	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 40 MG	83337030203030	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 50 MG	83337030203035	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 110 MG	83337030203040	Brand
PRADAXA	DABIGATRAN ETEXILATE MESYLATE PELLET PACK 150 MG	83337030203045	Brand

1 - Trial and failure, contraindication, or intolerance to rivaroxaban suspension

Date	Notes
8/25/2023	New Program

Preferred and Unrestricted Insulin Quantity Limit Excepti		
(g) beliefung mensebagik belieg natura kent onto a dan ak kebuhayan kentakan kentakan		

Guideline ID	GL-139113	
Guideline Name	Preferred and Unrestricted Insulin Quantity Limit Exception	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/17/2024
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1. Criteria

Product Name: Novolin N, Novolin R, Novolin 70/30, Novolog, Novolog 70/30, Humulin R U500, Semglee-yfgn				
Approval Length		12/31/2039		
Guideline Type		Quantity Limit - All plans except IL and MN Plans		
Product Name	Gen	Generic Name GPI Brand/Generic		
NOVOLIN N	INSU UNIT	LIN NPH (HUMAN) (ISOPHANE) INJ 100 /ML	27104020001805	Brand
NOVOLIN R	INSU	LIN REGULAR (HUMAN) INJ 100 UNIT/ML	27104010002005	Brand
NOVOLIN N FLEXPEN RELION		LIN NPH (HUMAN) (ISOPHANE) SUSP INJECTOR 100 UNIT/ML	2710402000D320	Brand
NOVOLIN N FLEXPEN		LIN NPH (HUMAN) (ISOPHANE) SUSP INJECTOR 100 UNIT/ML	2710402000D320	Brand
NOVOLIN N RELION	INSU UNIT	LIN NPH (HUMAN) (ISOPHANE) INJ 100 ML	27104020001805	Brand

1			
NOVOLIN 70/30 FLEXPEN RELION	INSULIN NPH & REGULAR SUSP PEN-INJ 100 UNIT/ML (70-30)	2710409000D320	Brand
NOVOLIN 70/30 FLEXPEN	INSULIN NPH & REGULAR SUSP PEN-INJ 100 UNIT/ML (70-30)	2710409000D320	Brand
NOVOLIN 70/30 RELION	INSULIN NPH ISOPHANE & REGULAR HUMAN INJ 100 UNIT/ML (70-30)	27104090001810	Brand
NOVOLIN 70/30	INSULIN NPH ISOPHANE & REGULAR HUMAN INJ 100 UNIT/ML (70-30)	27104090001810	Brand
NOVOLOG FLEXPEN	INSULIN ASPART SOLN PEN-INJECTOR 100 UNIT/ML	2710400200D220	Brand
NOVOLOG FLEXPEN RELION	INSULIN ASPART SOLN PEN-INJECTOR 100 UNIT/ML	2710400200D220	Brand
NOVOLOG PENFILL	INSULIN ASPART SOLN CARTRIDGE 100 UNIT/ML	2710400200E220	Brand
NOVOLOG RELION	INSULIN ASPART INJ SOLN 100 UNIT/ML	27104002002022	Brand
NOVOLOG	INSULIN ASPART INJ SOLN 100 UNIT/ML	27104002002022	Brand
NOVOLOG MIX 70/30 PREFILLED FLEXPEN	INSULIN ASPART PROT & ASPART SUS PEN- INJ 100 UNIT/ML (70-30)	2710407000D320	Brand
NOVOLOG MIX 70/30 PREFILLED FLEXPEN RELION	INSULIN ASPART PROT & ASPART SUS PENINJ 100 UNIT/ML (70-30)	2710407000D320	Brand
NOVOLOG MIX 70/30 RELION	INSULIN ASPART PROT & ASPART (HUMAN) INJ 100 UNIT/ML (70-30)	27104070001820	Brand
NOVOLOG MIX 70/30	INSULIN ASPART PROT & ASPART (HUMAN) INJ 100 UNIT/ML (70-30)	27104070001820	Brand
HUMULIN R U-500 KWIKPEN	INSULIN REGULAR (HUMAN) SOLN PEN- INJECTOR 500 UNIT/ML	2710401000D250	Brand
NOVOLIN R RELION	INSULIN REGULAR (HUMAN) INJ 100 UNIT/ML	27104010002005	Brand
HUMULIN R U-500 (CONCENTRATED)	INSULIN REGULAR (HUMAN) INJ 500 UNIT/ML	27104010002015	Brand
SEMGLEE	INSULIN GLARGINE-YFGN SOLN PEN- INJECTOR 100 UNIT/ML	2710400390D220	Brand
SEMGLEE	INSULIN GLARGINE-YFGN INJ 100 UNIT/ML	27104003902020	Brand

 $\boldsymbol{1}$ - Submission of medical records (e.g., chart notes) documenting insulin dosing and directions

AND

2 - Member requires more than 150 units per 30 days or, for U500 pens, more than 100 units per 30 days, or U500 vial, more than 333 units per 30 days based on daily prescribed dosing

The approval edits for quantity limit exceptions should be rounded up to allow the full trade package size, when necessary (e.g., the request ed quantity is 45ml per 30 days and the product is available in 30ml and 100ml, approve to a quantity of 60ml per 30 days, if the guideline criteria has been met).
riteria nas been met).

Product Name: Novolin N, Novolin R, Novolin 70/30, Novolog, Novolog 70/30, Humulin R U500, Semglee-yfgn		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Quantity Limit - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
NOVOLIN N	INSULIN NPH (HUMAN) (ISOPHANE) INJ 100 UNIT/ML	27104020001805	Brand
NOVOLIN R	INSULIN REGULAR (HUMAN) INJ 100 UNIT/ML	27104010002005	Brand
NOVOLIN R FLEXPEN RELION	INSULIN REGULAR (HUMAN) SOLN PEN- INJECTOR 100 UNIT/ML	2710401000D220	Brand
NOVOLIN R FLEXPEN	INSULIN REGULAR (HUMAN) SOLN PEN- INJECTOR 100 UNIT/ML	2710401000D220	Brand
NOVOLIN R RELION	INSULIN REGULAR (HUMAN) INJ 100 UNIT/ML	27104010002005	Brand
NOVOLIN N FLEXPEN RELION	INSULIN NPH (HUMAN) (ISOPHANE) SUSP PEN-INJECTOR 100 UNIT/ML	2710402000D320	Brand
NOVOLIN N FLEXPEN	INSULIN NPH (HUMAN) (ISOPHANE) SUSP PEN-INJECTOR 100 UNIT/ML	2710402000D320	Brand
NOVOLIN N RELION	INSULIN NPH (HUMAN) (ISOPHANE) INJ 100 UNIT/ML	27104020001805	Brand
NOVOLIN 70/30 FLEXPEN RELION	INSULIN NPH & REGULAR SUSP PEN-INJ 100 UNIT/ML (70-30)	2710409000D320	Brand
NOVOLIN 70/30 FLEXPEN	INSULIN NPH & REGULAR SUSP PEN-INJ 100 UNIT/ML (70-30)	2710409000D320	Brand
NOVOLIN 70/30 RELION	INSULIN NPH ISOPHANE & REGULAR HUMAN INJ 100 UNIT/ML (70-30)	27104090001810	Brand
NOVOLIN 70/30	INSULIN NPH ISOPHANE & REGULAR HUMAN INJ 100 UNIT/ML (70-30)	27104090001810	Brand

NOVOLOG FLEXPEN	INSULIN ASPART SOLN PEN-INJECTOR 100 UNIT/ML	2710400200D220	Brand
NOVOLOG FLEXPEN RELION	INSULIN ASPART SOLN PEN-INJECTOR 100 UNIT/ML	2710400200D220	Brand
NOVOLOG PENFILL	INSULIN ASPART SOLN CARTRIDGE 100 UNIT/ML	2710400200E220	Brand
NOVOLOG RELION	INSULIN ASPART INJ SOLN 100 UNIT/ML	27104002002022	Brand
NOVOLOG	INSULIN ASPART INJ SOLN 100 UNIT/ML	27104002002022	Brand
NOVOLOG MIX 70/30 PREFILLED FLEXPEN	INSULIN ASPART PROT & ASPART SUS PENINJ 100 UNIT/ML (70-30)	2710407000D320	Brand
NOVOLOG MIX 70/30 PREFILLED FLEXPEN RELION	INSULIN ASPART PROT & ASPART SUS PENINJ 100 UNIT/ML (70-30)	2710407000D320	Brand
NOVOLOG MIX 70/30 RELION	INSULIN ASPART PROT & ASPART (HUMAN) INJ 100 UNIT/ML (70-30)	27104070001820	Brand
NOVOLOG MIX 70/30	INSULIN ASPART PROT & ASPART (HUMAN) INJ 100 UNIT/ML (70-30)	27104070001820	Brand
HUMULIN R U-500 KWIKPEN	INSULIN REGULAR (HUMAN) SOLN PEN- INJECTOR 500 UNIT/ML	2710401000D250	Brand
HUMULIN R U-500 (CONCENTRATED)	INSULIN REGULAR (HUMAN) INJ 500 UNIT/ML	27104010002015	Brand
SEMGLEE	INSULIN GLARGINE-YFGN SOLN PEN- INJECTOR 100 UNIT/ML	2710400390D220	Brand
SEMGLEE	INSULIN GLARGINE-YFGN INJ 100 UNIT/ML	27104003902020	Brand

1 - Submission of medical records (e.g., chart notes) documenting insulin dosing and directions

AND

2 - Member requires more than 150 units per 30 days or, for U500 pens, more than 100 units per 30 days or, for U500 vial, more than 333 units per 30 days based on daily prescribed dosing

Notes	The approval edits for quantity limit exceptions should be rounded up to allow the full trade package size, when necessary (e.g., the request ed quantity is 45ml per 30 days and the product is available in 30ml and 100ml, approve to a quantity of 60ml per 30 days, if the guideline contests have been mot.)
Notes	o allow the full trade package size, when necessary (e.g., the requeed quantity is 45ml per 30 days and the product is available in 30ml

Product Name: Novolin N, Novolin R, Novolin 70/30, Novolog, Novolog 70/30, Humulin R U500, Semglee-yfgn		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Quantity Limit - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
NOVOLIN N	INSULIN NPH (HUMAN) (ISOPHANE) INJ 100 UNIT/ML	27104020001805	Brand
NOVOLIN R	INSULIN REGULAR (HUMAN) INJ 100 UNIT/ML	27104010002005	Brand
NOVOLIN R FLEXPEN RELION	INSULIN REGULAR (HUMAN) SOLN PENINJECTOR 100 UNIT/ML	2710401000D220	Brand
NOVOLIN R FLEXPEN	INSULIN REGULAR (HUMAN) SOLN PENINJECTOR 100 UNIT/ML	2710401000D220	Brand
NOVOLIN R RELION	INSULIN REGULAR (HUMAN) INJ 100 UNIT/ML	27104010002005	Brand
NOVOLIN N FLEXPEN RELION	INSULIN NPH (HUMAN) (ISOPHANE) SUSP PEN-INJECTOR 100 UNIT/ML	2710402000D320	Brand
NOVOLIN N FLEXPEN	INSULIN NPH (HUMAN) (ISOPHANE) SUSP PEN-INJECTOR 100 UNIT/ML	2710402000D320	Brand
NOVOLIN N RELION	INSULIN NPH (HUMAN) (ISOPHANE) INJ 100 UNIT/ML	27104020001805	Brand
HUMULIN 70/30 KWIKPEN	INSULIN NPH & REGULAR SUSP PEN-INJ 100 UNIT/ML (70-30)	2710409000D320	Brand
NOVOLIN 70/30 FLEXPEN RELION	INSULIN NPH & REGULAR SUSP PEN-INJ 100 UNIT/ML (70-30)	2710409000D320	Brand
NOVOLIN 70/30 FLEXPEN	INSULIN NPH & REGULAR SUSP PEN-INJ 100 UNIT/ML (70-30)	2710409000D320	Brand
NOVOLIN 70/30 RELION	INSULIN NPH ISOPHANE & REGULAR HUMAN INJ 100 UNIT/ML (70-30)	27104090001810	Brand
NOVOLIN 70/30	INSULIN NPH ISOPHANE & REGULAR HUMAN INJ 100 UNIT/ML (70-30)	27104090001810	Brand
NOVOLOG FLEXPEN	INSULIN ASPART SOLN PEN-INJECTOR 100 UNIT/ML	2710400200D220	Brand
NOVOLOG FLEXPEN RELION	INSULIN ASPART SOLN PEN-INJECTOR 100 UNIT/ML	2710400200D220	Brand
NOVOLOG PENFILL	INSULIN ASPART SOLN CARTRIDGE 100 UNIT/ML	2710400200E220	Brand
NOVOLOG RELION	INSULIN ASPART INJ SOLN 100 UNIT/ML	27104002002022	Brand
NOVOLOG	INSULIN ASPART INJ SOLN 100 UNIT/ML	27104002002022	Brand

NOVOLOG MIX 70/30 PREFILLED FLEXPEN	INSULIN ASPART PROT & ASPART SUS PENINJ 100 UNIT/ML (70-30)	2710407000D320	Brand
NOVOLOG MIX 70/30 PREFILLED FLEXPEN RELION	INSULIN ASPART PROT & ASPART SUS PENINJ 100 UNIT/ML (70-30)	2710407000D320	Brand
NOVOLOG MIX 70/30 RELION	INSULIN ASPART PROT & ASPART (HUMAN) INJ 100 UNIT/ML (70-30)	27104070001820	Brand
NOVOLOG MIX 70/30	INSULIN ASPART PROT & ASPART (HUMAN) INJ 100 UNIT/ML (70-30)	27104070001820	Brand
HUMULIN R U-500 KWIKPEN	INSULIN REGULAR (HUMAN) SOLN PEN- INJECTOR 500 UNIT/ML	2710401000D250	Brand
HUMULIN R U-500 (CONCENTRATED)	INSULIN REGULAR (HUMAN) INJ 500 UNIT/ML	27104010002015	Brand
SEMGLEE	INSULIN GLARGINE-YFGN SOLN PEN- INJECTOR 100 UNIT/ML	2710400390D220	Brand
SEMGLEE	INSULIN GLARGINE-YFGN INJ 100 UNIT/ML	27104003902020	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Notes	The approval edits for quantity limit exceptions should be rounded up to allow the full trade package size, when necessary (e.g., the request ed quantity is 45ml per 30 days and the product is available in 30ml and 100ml, approve to a quantity of 60ml per 30 days, if the guideline c
	riteria has been met).

Date	Notes
1/17/2024	Update program

[3] The Mindellings and Analysis, To Andrew Standard Control and Analysis and Analy	ptio

Guideline ID GL-131588	
Guideline Name	Preferred Blood Glucose Test Strips Quantity Limit Exception
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Na	Product Name: Onetouch Verio, Onetouch Ultra			
Approval Length Guideline Type		12/31/2039		
		Quantity Limit - All plans except IL and MN Plans		
Product Name	Generic Na	ame	GPI	Brand/Generic
ONETOUCH VERIO TEST STRIPS	GLUCOSE BLOOD TEST STRIP		94100030006100	Brand
ONETOUCH ULTRA	GLUCOSE E	BLOOD TEST STRIP	94100030006100	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting directions and frequency of blood sugar checks indicating member requires more than 200 strips per 30 days

Product Name: Onetouch Verio, Onetouch Ultra			
Approval Length 12 month(s)			
Guideline Type	Quantity Limit - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
ONETOUCH VERIO TEST STRIPS	GLUCOSE BLOOD TEST STRIP	94100030006100	Brand
ONETOUCH ULTRA	GLUCOSE BLOOD TEST STRIP	94100030006100	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting directions and frequency of blood sugar checks indicating member requires more than 200 strips per 30 days

Date	Notes
10/24/2023	2024 New Implementation

Prevymis (letermovir)
Sign financing many tradegapes from the species conserved, a common and final financing from the second burst

Guideline ID	GL-144555
Guideline Name	Prevymis (letermovir)
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Na	me: Prevym	nis Tab		
Approval L	ength	1 Course up to 200 Days		
Guideline Type Product Generic Na Name PREVYMIS LETERMOVIE		Prior Authorization - ALL Plans Exce	ept IL and MN Pla	ns
		ime	GPI	Brand/Generic
		R TAB 240 MG	12200045000320	Brand
PREVYMIS	LETERMOVI	R TAB 480 MG	12200045000340	Brand

Approval Criteria

1 - Requested drug is being used for cytomegalovirus (CMV) prophylaxis with one of the following:

- Post allogenic hematopoietic stem cell transplant
- Post kidney transplant

AND

2 - Submission of medical records (e.g., chart notes) documenting cytomegalovirus (CMV)-seropositive recipient (R+) or a CMV positive donor (D+)

AND

- **3** One of the following:
 - Drug is initiated within the first allogenic hematopoietic stem cell transplant: 28 days post-transplant
 - Drug is initiated within the first kidney transplant: 7 days post-transplant

AND

4 - Submission of medical records (e.g., chart notes) documenting that member does not have active CMV infection (CMV PCR level over 250 IU/mL) and is not receiving preemptive treatment (e.g., foscarnet)

AND

- **5** Prescribed by or in consultation with one of the following:
 - hematologist
 - oncologist
 - · infectious disease specialist
 - transplant specialist

*Member new to the plan (as evidenced by coverage effective less than or equal to 90 days) who are established on therapy ve coverage under their drug benefit for the remainder of the eatment course (to a maximum of day 200 post-transplant) ***Member new to the plan (as evidenced by coverage effection of less than or equal to 90 days) who initiated therapy using a cturer-sponsored free drug program, provider samples, and/o	y, will ha current tr ive date a manufa
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rs will go through initial criteria, otherwise for continuation of therapy f
or new to plan, reauthorization criteria applies

Product Name: Prevymis Tab	
Approval Length	12 months with 7 fills
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
PREVYMIS	LETERMOVIR TAB 240 MG	12200045000320	Brand
PREVYMIS	LETERMOVIR TAB 480 MG	12200045000340	Brand

- **1** Requested drug is being used for cytomegalovirus (CMV) prophylaxis with one of the following:
 - Post allogenic hematopoietic stem cell transplant
 - Post kidney transplant

AND

2 - Submission of medical records (e.g., chart notes) documenting cytomegalovirus (CMV)-seropositive recipient (R+) or a CMV positive donor (D+)

AND

- 3 One of the following:
 - Drug is initiated within the first allogenic hematopoietic stem cell transplant: 28 days post-transplant
 - Drug is initiated within the first kidney transplant: 7 days post-transplant

AND

4 - Submission of medical records (e.g., chart notes) documenting that member does not

have active CMV infection (CMV PCR level over 250 IU/mL) and is not receiving preemptive treatment (e.g., foscarnet)

AND

- **5** Prescribed by or in consultation with one of the following:
 - hematologist
 - oncologist
 - infectious disease specialist
 - transplant specialist

Notes *Member new to the plan (as evidenced by coverage effective date		
ve coverage under their drug benefit for the remainder of the curre eatment course (to a maximum of day 200 post-transplant) ***Member new to the plan (as evidenced by coverage effective day of less than or equal to 90 days) who initiated therapy using a man cturer-sponsored free drug program, provider samples, and/or vou	Notes	***Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufa cturer-sponsored free drug program, provider samples, and/or vouche rs will go through initial criteria, otherwise for continuation of therapy f

Product Name: Prevymis Tab	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
PREVYMIS	LETERMOVIR TAB 240 MG	12200045000320	Brand
PREVYMIS	LETERMOVIR TAB 480 MG	12200045000340	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting evidence-based clinical rationale for using a duration beyond 200 days post-transplant

*Member new to the plan (as evidenced by coverage effective date of
less than or equal to 90 days) who are established on therapy, will ha
ve coverage under their drug benefit for the remainder of the current tr
eatment course (to a maximum of day 200 post-transplant)

***Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufa cturer-sponsored free drug program, provider samples, and/or vouche rs will go through initial criteria, otherwise for continuation of therapy f
or new to plan, reauthorization criteria applies

Date	Notes
3/18/2024	Updated product name

Pulmonary Arterial Hypertension (PAH)) Agents
(3) below the second of the se	

Guideline ID	GL-129862
Guideline Name	Pulmonary Arterial Hypertension (PAH) Agents
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Preferred Drugs: Adempas, generic ambrisentan, generic bosentan, Opsumit, generic sildenafil, generic tadalafil, Uptravi				
Approval Leng	th	12 month(s)		
Therapy Stage)	Initial Authorization	Initial Authorization	
Guideline Type	Э	Prior Authorization - IL and MN Plans		
Product Name	Generio	Name GPI Brand/Generic		
AMBRISENTAN	AMBRISENTAN TAB 5 MG		40160007000310	Generic
AMBRISENTAN	AMBRISENTAN TAB 10 MG		40160007000320	Generic
BOSENTAN	BOSENTAN TAB 62.5 MG		40160015000320	Generic
BOSENTAN	BOSENTAN TAB 125 MG		40160015000330	Generic
OPSUMIT	MACITEN	NTAN TAB 10 MG	40160050000320	Brand

			
ADEMPAS	RIOCIGUAT TAB 0.5 MG	40134050000310	Brand
ADEMPAS	RIOCIGUAT TAB 1 MG	40134050000320	Brand
ADEMPAS	RIOCIGUAT TAB 1.5 MG	40134050000330	Brand
ADEMPAS	RIOCIGUAT TAB 2 MG	40134050000340	Brand
ADEMPAS	RIOCIGUAT TAB 2.5 MG	40134050000350	Brand
UPTRAVI TITRATION PACK	SELEXIPAG TAB THERAPY PACK 200 MCG (140) & 800 MCG (60)	4012007000B720	Brand
UPTRAVI	SELEXIPAG TAB 200 MCG	40120070000310	Brand
UPTRAVI	SELEXIPAG TAB 400 MCG	40120070000315	Brand
UPTRAVI	SELEXIPAG TAB 600 MCG	40120070000320	Brand
UPTRAVI	SELEXIPAG TAB 800 MCG	40120070000325	Brand
UPTRAVI	SELEXIPAG TAB 1000 MCG	40120070000330	Brand
UPTRAVI	SELEXIPAG TAB 1200 MCG	40120070000335	Brand
UPTRAVI	SELEXIPAG TAB 1400 MCG	40120070000340	Brand
UPTRAVI	SELEXIPAG TAB 1600 MCG	40120070000345	Brand
UPTRAVI	SELEXIPAG FOR IV SOLN 1800 MCG	40120070002120	Brand
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 20 MG	40143060100320	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE FOR SUSPENSION 10 MG/ML	40143060101920	Generic
SILDENAFIL	SILDENAFIL CITRATE FOR SUSPENSION 10 MG/ML	40143060101920	Generic
TADALAFIL	TADALAFIL TAB 20 MG (PAH)	40143080000320	Generic
			,

1 - Diagnosis of pulmonary arterial hypertension

AND

- **2** Prescribed by or in consultation with one of the following:

 - Cardiologist Pulmonologist

rer-sponsored free drug program, provider samples, and/or vouchers	Notes	Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
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Product Name: Non-Preferred Drugs: Orenitram, Ventavis		
Approval Length 12 month(s)		
Therapy Stage	herapy Stage Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
VENTAVIS	ILOPROST INHALATION SOLUTION 10 MCG/ML	40170060002020	Brand
VENTAVIS	ILOPROST INHALATION SOLUTION 20 MCG/ML	40170060002040	Brand
ORENITRAM TITRATION KIT MONTH 1	TREPROSTINIL TAB ER TITR PK (MO1) 126 X0.125MG & 42 X0.25MG	4017008005C110	Brand
ORENITRAM TITRATION KIT MONTH 2	TREPROSTINIL TAB ER TITR PK (MO2) 126 X0.125MG & 210 X0.25MG	4017008005C120	Brand
ORENITRAM TITRATION KIT MONTH 3	TREPROSTINIL TAB ER TITR PK(MO3)126X0.125MG&42X0.25MG&84X1MG	4017008005C130	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 0.125 MG (BASE EQUIV)	40170080050410	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 0.25 MG (BASE EQUIV)	40170080050415	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 1 MG (BASE EQUIV)	40170080050420	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 2.5 MG (BASE EQUIV)	40170080050425	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 5 MG (BASE EQUIV)	40170080050435	Brand

1 - Diagnosis of pulmonary arterial hypertension

AND

- **2** Prescribed by or in consultation with one of the following:
 - Cardiologist
 - Pulmonologist

AND

3 - Trial and failure, contraindication, or intolerance to inhaled treprostinil (Tyvaso)

Notes	Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for
	new to plan, reauthorization criteria applies

Product Name: Preferred Drugs: Adempas, generic ambrisentan, generic bosentan, Opsumit, generic sildenafil, generic tadalafil, Uptravi; and Non-Preferred Drugs: Orenitram, Ventavis

Approval Length 12 month(s)

Therapy Stage Reauthorization

Guideline Type Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
AMBRISENTAN	AMBRISENTAN TAB 5 MG	40160007000310	Generic
AMBRISENTAN	AMBRISENTAN TAB 10 MG	40160007000320	Generic
BOSENTAN	BOSENTAN TAB 62.5 MG	40160015000320	Generic
BOSENTAN	BOSENTAN TAB 125 MG	40160015000330	Generic
OPSUMIT	MACITENTAN TAB 10 MG	40160050000320	Brand
ADEMPAS	RIOCIGUAT TAB 0.5 MG	40134050000310	Brand
ADEMPAS	RIOCIGUAT TAB 1 MG	40134050000320	Brand
ADEMPAS	RIOCIGUAT TAB 1.5 MG	40134050000330	Brand
ADEMPAS	RIOCIGUAT TAB 2 MG	40134050000340	Brand
ADEMPAS	RIOCIGUAT TAB 2.5 MG	40134050000350	Brand

UPTRAVI TITRATION PACK	SELEXIPAG TAB THERAPY PACK 200 MCG (140) & 800 MCG (60)	4012007000B720	Brand
UPTRAVI	SELEXIPAG TAB 200 MCG	40120070000310	Brand
UPTRAVI	SELEXIPAG TAB 400 MCG	40120070000315	Brand
UPTRAVI	SELEXIPAG TAB 600 MCG	40120070000320	Brand
UPTRAVI	SELEXIPAG TAB 800 MCG	40120070000325	Brand
UPTRAVI	SELEXIPAG TAB 1000 MCG	40120070000330	Brand
UPTRAVI	SELEXIPAG TAB 1200 MCG	40120070000335	Brand
UPTRAVI	SELEXIPAG TAB 1400 MCG	40120070000340	Brand
UPTRAVI	SELEXIPAG TAB 1600 MCG	40120070000345	Brand
UPTRAVI	SELEXIPAG FOR IV SOLN 1800 MCG	40120070002120	Brand
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 20 MG	40143060100320	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE FOR SUSPENSION 10 MG/ML	40143060101920	Generic
SILDENAFIL	SILDENAFIL CITRATE FOR SUSPENSION 10 MG/ML	40143060101920	Generic
TADALAFIL	TADALAFIL TAB 20 MG (PAH)	40143080000320	Generic
VENTAVIS	ILOPROST INHALATION SOLUTION 10 MCG/ML	40170060002020	Brand
VENTAVIS	ILOPROST INHALATION SOLUTION 20 MCG/ML	40170060002040	Brand
ORENITRAM TITRATION KIT MONTH 1	TREPROSTINIL TAB ER TITR PK (MO1) 126 X0.125MG & 42 X0.25MG	4017008005C110	Brand
ORENITRAM TITRATION KIT MONTH 2	TREPROSTINIL TAB ER TITR PK (MO2) 126 X0.125MG & 210 X0.25MG	4017008005C120	Brand
ORENITRAM TITRATION KIT MONTH 3	TREPROSTINIL TAB ER TITR PK(MO3)126X0.125MG&42X0.25MG&84X1MG	4017008005C130	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 0.125 MG (BASE EQUIV)	40170080050410	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 0.25 MG (BASE EQUIV)	40170080050415	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 1 MG (BASE EQUIV)	40170080050420	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 2.5 MG (BASE EQUIV)	40170080050425	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 5 MG (BASE EQUIV)	40170080050435	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for
new to plan, reauthorization criteria applies

Product Name: Preferred Drugs: Adempas, generic ambrisentan, generic bosentan, Opsumit, generic sildenafil, generic tadalafil, Uptravi

Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
AMBRISENTAN	AMBRISENTAN TAB 5 MG	40160007000310	Generic
AMBRISENTAN	AMBRISENTAN TAB 10 MG	40160007000320	Generic
BOSENTAN	BOSENTAN TAB 62.5 MG	40160015000320	Generic
BOSENTAN	BOSENTAN TAB 125 MG	40160015000330	Generic
OPSUMIT	MACITENTAN TAB 10 MG	40160050000320	Brand
ADEMPAS	RIOCIGUAT TAB 0.5 MG	40134050000310	Brand
ADEMPAS	RIOCIGUAT TAB 1 MG	40134050000320	Brand
ADEMPAS	RIOCIGUAT TAB 1.5 MG	40134050000330	Brand
ADEMPAS	RIOCIGUAT TAB 2 MG	40134050000340	Brand
ADEMPAS	RIOCIGUAT TAB 2.5 MG	40134050000350	Brand
UPTRAVI TITRATION PACK	SELEXIPAG TAB THERAPY PACK 200 MCG (140) & 800 MCG (60)	4012007000B720	Brand
UPTRAVI	SELEXIPAG TAB 200 MCG	40120070000310	Brand
UPTRAVI	SELEXIPAG TAB 400 MCG	40120070000315	Brand
UPTRAVI	SELEXIPAG TAB 600 MCG	40120070000320	Brand
UPTRAVI	SELEXIPAG TAB 800 MCG	40120070000325	Brand
UPTRAVI	SELEXIPAG TAB 1000 MCG	40120070000330	Brand

UPTRAVI	SELEXIPAG TAB 1200 MCG	40120070000335	Brand
UPTRAVI	SELEXIPAG TAB 1400 MCG	40120070000340	Brand
UPTRAVI	SELEXIPAG TAB 1600 MCG	40120070000345	Brand
UPTRAVI	SELEXIPAG FOR IV SOLN 1800 MCG	40120070002120	Brand
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 20 MG	40143060100320	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE FOR SUSPENSION 10 MG/ML	40143060101920	Generic
SILDENAFIL	SILDENAFIL CITRATE FOR SUSPENSION 10 MG/ML	40143060101920	Generic
TADALAFIL	TADALAFIL TAB 20 MG (PAH)	40143080000320	Generic

1 - Diagnosis of pulmonary arterial hypertension

AND

- **2** Prescribed by or in consultation with one of the following:

 - Cardiologist Pulmonologist

Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for
new to plan, reauthorization criteria applies

Product Name: Non-Preferred Drugs: Orenitram, Ventavis		
Approval Length	12/31/2039	
Guideline Type Prior Authorization - All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
VENTAVIS	ILOPROST INHALATION SOLUTION 10 MCG/ML	40170060002020	Brand
VENTAVIS	ILOPROST INHALATION SOLUTION 20 MCG/ML	40170060002040	Brand

ORENITRAM TITRATION KIT MONTH 1	TREPROSTINIL TAB ER TITR PK (MO1) 126 X0.125MG & 42 X0.25MG	4017008005C110	Brand
ORENITRAM TITRATION KIT MONTH 2	TREPROSTINIL TAB ER TITR PK (MO2) 126 X0.125MG & 210 X0.25MG	4017008005C120	Brand
ORENITRAM TITRATION KIT MONTH 3	TREPROSTINIL TAB ER TITR PK(MO3)126X0.125MG&42X0.25MG&84X1MG	4017008005C130	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 0.125 MG (BASE EQUIV)	40170080050410	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 0.25 MG (BASE EQUIV)	40170080050415	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 1 MG (BASE EQUIV)	40170080050420	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 2.5 MG (BASE EQUIV)	40170080050425	Brand
ORENITRAM	TREPROSTINIL DIOLAMINE TAB ER 5 MG (BASE EQUIV)	40170080050435	Brand

1 - Diagnosis of pulmonary arterial hypertension

AND

- **2** Prescribed by or in consultation with one of the following:

 - Cardiologist Pulmonologist

AND

3 - Trial and failure, contraindication, or intolerance to inhaled treprostinil (Tyvaso)

Notes	Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu
	rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

Date	Notes
10/12/2023	2024 New Implementation

Pyrukynd
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Guideline ID	GL-130133	
Guideline Name	Pyrukynd	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Pyrukynd	
Approval Length	6 month(s)
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 5 MG	8587005070B710	Brand
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 7 X 20 MG & 7 X 5 MG	8587005070B720	Brand
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 7 X 50 MG & 7 X 20 MG	8587005070B735	Brand

PYRUKYND	MITAPIVAT SULFATE TAB 5 MG	85870050700310	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 20 MG	85870050700325	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 50 MG	85870050700340	Brand

1 - Diagnosis of pyruvate kinase deficiency

AND

2 - Prescribed by, or in consultation with, a Hematologist or other expert in treating hemolytic anemia

AND

3 - Hemoglobin less than or equal to 10 mg/dL

AND

4 - Greater than or equal to 1 red blood cell (RBC) transfusion in the past 12 months

AND

5 - Member is 18 years of age or older

Notes	Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for
	new to plan, reauthorization criteria applies

Product Name: Pyrukynd	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 5 MG	8587005070B710	Brand
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 7 X 20 MG & 7 X 5 MG	8587005070B720	Brand
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 7 X 50 MG & 7 X 20 MG	8587005070B735	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 5 MG	85870050700310	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 20 MG	85870050700325	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 50 MG	85870050700340	Brand

1 - Diagnosis of pyruvate kinase deficiency

AND

2 - Prescribed by, or in consultation with, a Hematologist or other expert in treating hemolytic anemia

AND

3 - Hemoglobin less than or equal to 10 mg/dL

AND

4 - Greater than or equal to 1 red blood cell (RBC) transfusion in the past 12 months

AND

5 - Member is 18 years of age or older

		Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
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Product Name: Pyrukynd	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 5 MG	8587005070B710	Brand
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 7 X 20 MG & 7 X 5 MG	8587005070B720	Brand
PYRUKYND TAPER PACK	MITAPIVAT SULFATE TAB THERAPY PACK 7 X 50 MG & 7 X 20 MG	8587005070B735	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 5 MG	85870050700310	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 20 MG	85870050700325	Brand
PYRUKYND	MITAPIVAT SULFATE TAB 50 MG	85870050700340	Brand

1 - Diagnosis of pyruvate kinase deficiency

AND

2 - Submission of medical records (e.g., chart notes) documenting that within the past 6 months (for initial starts) or past 12 months the member demonstrates positive clinical response to therapy

Notes	Member new to the plan (as evidenced by coverage effective date of I ess than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
	new to plan, readmonzation entena applies

Date	Notes
10/6/2023	2024 New Implementation

Qbrexza (Glycopyrronium topical)				
The interfuence of the figure is the first term and, would a little and doubt him to comit arithms.				

Guideline ID	GL-129624
Guideline Name	Qbrexza (Glycopyrronium topical)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Qbrexza		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
QBREXZA	GLYCOPYRRONIUM TOSYLATE PAD 2.4% (BASE EQUIVALENT)	90970030204320	Brand

Approval Criteria

1 - Diagnosis of axillary hyperhidrosis with clinical documentation of a persistent or chronic

cutaneous condition due to excessive sweating (e.g. skin maceration, dermatitis, fungal infections)

AND

2 - Failure of an adequate trial or intolerance to BOTH prescription strength topical aluminum antiperspirants and an oral anticholinergic drug such as glycopyrrolate or oxybutynin

Product Name: Qbrexza		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
QBREXZA	GLYCOPYRRONIUM TOSYLATE PAD 2.4% (BASE EQUIVALENT)	90970030204320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Qbrexza		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
QBREXZA	GLYCOPYRRONIUM TOSYLATE PAD 2.4% (BASE EQUIVALENT)	90970030204320	Brand

Approval Criteria

1 - Diagnosis of axillary hyperhidrosis with clinical documentation of a persistent or chronic cutaneous condition due to excessive sweating (e.g. skin maceration, dermatitis, fungal infections)

AND

2 - Failure of an adequate trial or intolerance to BOTH prescription strength topical aluminum antiperspirants and an oral anticholinergic drug such as glycopyrrolate or oxybutynin

Date	Notes
10/6/2023	New Program

Quantity Limit Exceptions
Control of the second s

Guideline ID	GL-144892
Guideline Name	Quantity Limit Exceptions
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	1/18/2017
P&T Revision Date:	7/18/2023

1. Criteria

Diagnosis		CRITERIA FOR COVERAGE OF QUANTITY EXCEPTIONS (DOSE ≤ MAXIMUM DOSE IN PRESCRIBING INFORMATION) - Titration or loading dose		
Approval Length		One Time Fill		
Guideline Type		Administrative		
Product Name Gene		ric Name	GPI	Brand/Generic
Approval Criteria				
1 - Request is for a titration or loading dose				

Diagnosis	CRITERIA FOR COVERAGE OF QUANTITY EXCEPTIONS (DOSE ≤ MAXIMUM DOSE IN PRESCRIBING INFORMATION)
Approval Length	12 month(s)
Guideline Type	Administrative
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Product Name	Generic Name	GPI	Brand/Generic
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1 - Person is on a dose alternating schedule

OR

2 - For topical applications: person requires a larger quantity to cover a larger surface area

OR

3 - Requested strength/dose is commercially unavailable

OR

4 - Requested strength/dose is listed in the prescribing information as the standard maintenance dosing regimen for the submitted diagnosis

OR

5 - If the request is for reauthorization, prescriber submits medical records (e.g., chart notes) of documentation from the previous 12 months that the member is continuing therapy with the requested drug and dosing regimen

Diagnosis	CRITERIA FOR COVERAGE OF QUANTITY EXCEPTIONS (DOSE > MAXIMUM DOSE IN PRESCRIBING INFORMATION)
Approval Length	12 month(s)

Guideline Type	Administrative		
Product Name	Generic Name	GPI	Brand/Generic

- 1 Both of the following:
- **1.1** One of the following:
- **1.1.1** Higher dose or quantity is supported by clinical research in two articles from major peer reviewed medical journals that present data supporting the proposed higher than maximum doses for the diagnosis provided as generally safe and effective

OR

1.1.2 Higher dose or quantity is supported by American Hospital Formulary Service Drug Information or Micromedex DRUGDEX System

AND

- **1.2** One of the following
- **1.2.1** Maximum doses specified under the quantity restriction have been tried for an adequate period of time and been deemed ineffective in the treatment of the member's disease or medical condition

OR

1.2.2 If lower doses have not been tried, there is clinical support (i.e., clinical literature, patient attributes, or characteristics of the drug) that the number of doses available under the quantity restriction will be ineffective in the treatment of the member's disease or medical condition

OR

2 - If the request is for reauthorization, prescriber submits medical records (e.g., chart notes) of documentation from the previous 12 months that the member is continuing therapy with the requested drug and dosing regimen

Diagnosis		All Indications		
Approval Length		12 month(s)		
Therapy Stage		Reauthorization		
Guideline Type		Administrative		
Product Name	Gene	ric Name	GPI	Brand/Generic

1 - Prescriber submits medical records (e.g., chart notes) of documentation from the previous 12 months that the member is continuing therapy with the requested drug and dosing regimen

Date	Notes
3/26/2024	Updated Guideline

Radicava (Edaravone)
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Guideline ID	GL-144894
Guideline Name	Radicava (Edaravone)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	8/16/2017
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Radicava ORS				
Approval L	Length 12 month(s)			
Guideline 7	Type Prior Authorization			
Product Name	Generic Name		GPI	Brand/Generic
RADICAVA ORS STARTER KIT	EDARAVONE ORAL SUSP 105 MG/5ML		74509030001820	Brand
RADICAVA ORS	EDARAVONE	ORAL SUSP 105 MG/5ML	74509030001820	Brand

Approval Criteria
1 - Diagnosis of definite or probable ALS based on El Escorial revised Airlie House diagnostic criteria
AND
2 - Prescribed by, or in consultation with, a Neurologist or other specialist in treating amyotrophic lateral sclerosis (ALS)
AND
3 - Age 20-75
AND
4 - One of the following:
4.1 All of the following:
4.1.1 Independent living status (i.e., Japan ALS Severity Classification Grade 1 or 2)
AND
4.1.2 Score of ≥ 2 on all 12 items of the ALS Functional Rating Scale (ALSFRS-R) (assessed and documented within the last 3 months)
AND
4.1.3 FVC % predicted ≥ 80% (assessed and documented within the last 3 months)
AND
4.1.4 Duration of disease from the first symptom of 2 years or less

AND

4.1.5 Current use of riluzole or documented contraindication/intolerance/ lack of therapeutic effect of therapy

OR

4.2 Continuation of prior therapy with edaravone, verified by paid claims or medical records (e.g. chart notes)

Date	Notes
3/26/2024	Updated Guideline

Rayos (prednisone DR)
(g) behaviorage and behave. We have to see that and a state and a state of the behaviorage and the behavio

Guideline ID	GL-136613
Guideline Name	Rayos (prednisone DR)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
P&T Approval Date:	
P&T Revision Date:	

1. Criteria

RAYOS

RAYOS

Approval Length 12 month(s) Therapy Stage Initial Authorization	Product Name: Rayos			
Therapy Stage Initial Authorization	Approval Length	12 month(s)		
	Therapy Stage	Initial Authorization		
Guideline Type Prior Authorization-IL and MN Plans Only	Guideline Type	Prior Authorization-IL and MN Plan	s Only	
Product Generic Name GPI Brand/Gen		lame	GPI	Brand/Generic
RAYOS PREDNISONE TAB DELAYED RELEASE 1 MG 22100045000610 Brand	RAYOS PREDNIS	NE TAB DELAYED RELEASE 1 MG	22100045000610	Brand

PREDNISONE TAB DELAYED RELEASE 2 MG

PREDNISONE TAB DELAYED RELEASE 5 MG

Brand

Brand

22100045000620

22100045000630

- 1 One of the following:
- **1.1** Both of the following:
- **1.1.1** Therapy-limiting intolerance of immediate-release prednisone despite dose adjustment and/or timing modification

OR

1.1.2 The prescriber provides an evidence-based clinical rationale for why the side effects are not likely to occur with the extended-release formulation

OR

1.2 Minnesota plans only: Member with stage four metastatic cancer and the requested drug is being used as supportive care to treat fatigue related to their cancer diagnosis or chemotherapy regimen

Product Name: Rayos	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
RAYOS	PREDNISONE TAB DELAYED RELEASE 1 MG	22100045000610	Brand
RAYOS	PREDNISONE TAB DELAYED RELEASE 2 MG	22100045000620	Brand
RAYOS	PREDNISONE TAB DELAYED RELEASE 5 MG	22100045000630	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Rayos	
Approval Length	12 month(s)
Guideline Type	Prior Authorization-All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
RAYOS	PREDNISONE TAB DELAYED RELEASE 1 MG	22100045000610	Brand
RAYOS	PREDNISONE TAB DELAYED RELEASE 2 MG	22100045000620	Brand
RAYOS	PREDNISONE TAB DELAYED RELEASE 5 MG	22100045000630	Brand

1 - Therapy-limiting intolerance of immediate-release prednisone despite dose adjustment and/or timing modification

AND

2 - The prescriber provides an evidence-based clinical rationale for why the side effects are not likely to occur with the extended-release formulation

Date	Notes
11/21/2023	Update program

Relyvrio (sodium phenylbutyrate and taurursodi		

Guideline ID	GL-131273	
Guideline Name	Relyvrio (sodium phenylbutyrate and taurursodiol)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Relyvrio	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - ALL Plans

Product Name	Generic Name	GPI	Brand/Generic
RELYVRIO	SODIUM PHENYLBUTYRATE-TAURURSODIOL POWD PACK 3-1 GM	74509902703020	Brand

Approval Criteria

1 - Diagnosis of definite or probable amyotrophic lateral sclerosis (ALS)

	AND
2 - Member is 18 years	of age or older
	AND
3 - Submission of medi greater than 60%, withi	cal records (e.g., chart notes) documenting Slow vital capacity (SVC) in the past 3 months
	AND
4 - Member has not cui	rrently had a tracheostomy or on permanent assisted ventilation
	AND
5 - Duration of disease	from the first symptom, is of 18 months or less
	AND
6 - Member is currently using riluzole or has a documented contraindication/intolerance/or lack of therapeutic effect of therapy	
AND	
7 - Prescribed by or in	consultation with one of the following:
 neurologist other specialist in the treatment of ALS 	
Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

Product Name: Relyvrio	
Approval Length 12 month(s)	
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - ALL Plans	

Product Name	Generic Name	GPI	Brand/Generic
RELYVRIO	SODIUM PHENYLBUTYRATE-TAURURSODIOL POWD PACK 3-1 GM	74509902703020	Brand

1 - Submission of medical records (e.g., chart notes) documenting that the use of the drug has slowed the progression of ALS and function is improved relative to the expected natural course of the disease

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for
new to plan, reauthorization criteria applies

Date	Notes
10/6/2023	2024 New Implementation

Repatha (evolocumab)		
(2) The interpretation of the first the second country and a first to be presented as control.		

Guideline ID	GL-144897
Guideline Name	Repatha (evolocumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	2/20/2019
P&T Revision Date:	7/18/2023

Note:

1. Criteria

Product Name	Product Name: Repatha			
Approval Leng	jth	12 month(s)		
Therapy Stage)	Initial Authorization		
Guideline Type		Prior Authorization		
Product Name	Generic	Name	GPI	Brand/Generic

^{*}Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

REPATHA SURECLICK	EVOLOCUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 140 MG/ML	3935002000D520	Brand
REPATHA PUSHTRONEX SYSTEM	EVOLOCUMAB SUBCUTANEOUS SOLN CARTRIDGE/INFUSOR 420 MG/3.5ML	3935002000E230	Brand
REPATHA	EVOLOCUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 140 MG/ML	3935002000E520	Brand

- 1 Diagnosis of one of the following:
 - Heterozygous Familial Hypercholesteremia
 - Homozygous Familial Hypercholesterolemia
 - Established arteriosclerotic cardiovascular disease (ASCVD)

AND

2 - Submission of medical records (e.g., chart notes) documenting that medication is prescribed by or in consultation with a cardiologist, endocrinologist, or lipidologist

AND

- **3** All of the following:
- **3.1** Member has LDL-C greater than or equal to 70 mg/dL while on maximally tolerated statin doses

AND

- **3.2** One of the following:
- **3.2.1** All of the following:
- **3.2.1.1** Member is statin tolerant and will continue statin treatment in combination with PCSK9

AND

3.2.1.2 One of the following:

- Adherent treatment with a high potency statin (ex. atorvastatin 40-80 mg daily, rosuvastatin 20-40 mg daily) for a minimum of 8 weeks duration
- Member cannot tolerate high potency statin and adherent treatment with a maximally tolerated dose of any statin for a minimum of 8 weeks duration

OR

- **3.2.2** Member is statin intolerant as defined by all of the following:
 - Member was unable to tolerate at least two statins with one started at the lowest starting dose
 - Statin dose reduction was attempted to resolve symptoms or lab abnormalities (not discontinuation)
 - Symptoms or lab abnormalities reversed with statin discontinuation but returned with re-challenge of statins
 - Symptoms or lab abnormalities are not due to established predispositions such as drug interactions, significant changes in physical activity, or underlying muscle disease

OR

3.2.3 Member has a contraindication to statin use such as active liver disease or persistently elevated serum transaminases

Product Name: Repatha	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
REPATHA SURECLICK	EVOLOCUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 140 MG/ML	3935002000D520	Brand
REPATHA PUSHTRONEX SYSTEM	EVOLOCUMAB SUBCUTANEOUS SOLN CARTRIDGE/INFUSOR 420 MG/3.5ML	3935002000E230	Brand
REPATHA	EVOLOCUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 140 MG/ML	3935002000E520	Brand

- 1 One of the following:
- **1.1** Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days)

OR

1.2 Member has a previous prior authorization with the plan or historical prior authorization for evolocumab on file

AND

2 - Submission of medical records (e.g., chart notes) within the past 12 month demonstrating a reduction in LDL-C from baseline

AND

3 - Member continues treatment with baseline lipid-lowering therapies

Product Name: Repatha	
Approval Length	12/31/2039
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
REPATHA SURECLICK	EVOLOCUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 140 MG/ML	3935002000D520	Brand
REPATHA PUSHTRONEX SYSTEM	EVOLOCUMAB SUBCUTANEOUS SOLN CARTRIDGE/INFUSOR 420 MG/3.5ML	3935002000E230	Brand
REPATHA	EVOLOCUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 140 MG/ML	3935002000E520	Brand

1 - Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days)

AND

2 - Submission of medical records (e.g., chart notes) within the past 12 month demonstrating a reduction in LDL-C from baseline

AND

3 - Member continues treatment with baseline lipid-lowering therapies

Product Name: Repatha	
Approval Length	12 month(s)
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
REPATHA SURECLICK	EVOLOCUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 140 MG/ML	3935002000D520	Brand
REPATHA PUSHTRONEX SYSTEM	EVOLOCUMAB SUBCUTANEOUS SOLN CARTRIDGE/INFUSOR 420 MG/3.5ML	3935002000E230	Brand
REPATHA	EVOLOCUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 140 MG/ML	3935002000E520	Brand

Approval Criteria

1 - Diagnosis of primary hyperlipidemia

AND

2 - Submission of medical records (e.g., chart notes) documenting that medication is prescribed by or in consultation with a cardiologist, endocrinologist, or lipidologist

AND

- 3 One of the following:
- **3.1** Paid claims or submission of medical records (e.g. chart notes) show there have been adequate trials of ALL appropriate formulary therapeutic alternatives and there was (a) inadequate clinical response, (b) inappropriate clinical response, (c) intolerance or (d) allergy to those medications

OR

3.2 An exception to the formulary may be considered when ALL appropriate formulary therapeutic alternatives have not been tried and there is submission of medical record documentation (e.g. chart notes) demonstrating that ALL appropriate formulary therapeutic alternatives will be (a) ineffective, (b) less effective or (c) will result in adverse effects

Date	Notes
3/26/2024	Updated Guideline

Restricted Diclofe	nac
(3) This belongs are budget. While, the best was, same, a size only while pure to worther	

Guideline ID	GL-131458
Guideline Name	Restricted Diclofenac
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Zipsor, Generic Cambia	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
	DICLOFENAC POTASSIUM (MIGRAINE) PACKET 50 MG	67600040103020	Generic
DICLOFENAC POTASSIUM	DICLOFENAC POTASSIUM CAP 25 MG	66100007100120	Generic

Approval Criteria

1 - Trial and failure (with maximized doses) or intolerance of a preferred oral diclofenac

AND

2 - Trial and failure (with maximized doses) or intolerance of another preferred oral nonsteroidal anti-inflammatory drug (NSAID)

Product Name: Diclofenac 2% solution (generic Pennsaid), Diclofenac 1.5% solution,
Diclofenac 3% gel

Approval Length 12/31/2039

Guideline Type Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
DICLOFENAC SODIUM	DICLOFENAC SODIUM SOLN 2%	90210030302030	Generic
DICLOFENAC SODIUM	DICLOFENAC SODIUM SOLN 1.5%	90210030302025	Generic
DICLOFENAC SODIUM	DICLOFENAC SODIUM (ACTINIC KERATOSES) GEL 3%	90374035304020	Generic

Approval Criteria

- **1** Both of the following:
- **1.1** Trial and failure (with maximized doses), contraindication or intolerance to two preferred oral NSAIDs

AND

1.2 Trial and failure of maximized dosing of generic diclofenac 1% gel

OR

2 - (Diclofenac 3% gel only) Used for short-term treatment of actinic keratosis

Product Name: Generic Zipsor, Generic Cambia

Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
	DICLOFENAC POTASSIUM (MIGRAINE) PACKET 50 MG	67600040103020	Generic
DICLOFENAC POTASSIUM	DICLOFENAC POTASSIUM CAP 25 MG	66100007100120	Generic

- 1 Both of the following:
- 1.1 Trial and failure (with maximized doses) or intolerance of a preferred oral diclofenac

AND

1.2 Trial and failure (with maximized doses) or intolerance of another preferred oral nonsteroidal anti-inflammatory drug (NSAID)

OR

2 - (Minnesota plans only) – Member has stage four metastatic cancer and the requested drug is being used to treat cancer-related pain

Product Name: Diclofenac 2% solution (generic Pennsaid), Diclofenac 1.5% solution, Diclofenac 3% gel	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
DICLOFENAC SODIUM	DICLOFENAC SODIUM SOLN 2%	90210030302030	Generic
DICLOFENAC SODIUM	DICLOFENAC SODIUM SOLN 1.5%	90210030302025	Generic

DICLOFENAC DIC SODIUM 3%	CLOFENAC SODIUM (ACTINIC KERATOSES) GEL	90374035304020	Generic
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- **1** Both of the following:
- **1.1** Trial and failure (with maximized doses), contraindication or intolerance to two preferred oral NSAIDs

AND

1.2 Trial and failure of maximized dosing of generic diclofenac 1% gel

OR

2 - (Diclofenac 3% gel only) Used for short-term treatment of actinic keratosis

OR

3 - (Minnesota plans only) – Member has stage four metastatic cancer and the requested drug is being used to treat cancer-related pain

Product Name: Generic Zipsor, Generic Cambia, Diclofenac 2% solution (generic Pennsaid), Diclofenac 1.5% solution, Diclofenac 3% gel	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DICLOFENAC POTASSIUM	DICLOFENAC POTASSIUM (MIGRAINE) PACKET 50 MG	67600040103020	Generic
DICLOFENAC SODIUM	DICLOFENAC SODIUM SOLN 2%	90210030302030	Generic
DICLOFENAC SODIUM	DICLOFENAC SODIUM SOLN 1.5%	90210030302025	Generic

DICLOFENAC SODIUM	DICLOFENAC SODIUM (ACTINIC KERATOSES) GEL 3%	90374035304020	Generic
DICLOFENAC POTASSIUM	DICLOFENAC POTASSIUM CAP 25 MG	66100007100120	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
10/24/2023	2024 New Implementation

Restricted Inhaled Corticosteroid				
[2] The beat image context to deploye. The firms of their terms context, and	use a dear the first the provide a confidence of the confidence of			

Guideline ID	GL-143612
Guideline Name	Restricted Inhaled Corticosteroid
Formulary	Quartz

Guideline Note:

Effective Date:	4/1/2024
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1. Criteria

Product Name: Pulmicort Flexhaler, Alvesco			
Approval Length 12 month(s)			
Therapy Stage	herapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
	BUDESONIDE INHAL AERO POWD 90 MCG/ACT (BREATH ACTIVATED)	44400015008009	Brand
	BUDESONIDE INHAL AERO POWD 180 MCG/ACT (BREATH ACTIVATED)	44400015008018	Brand
ALVESCO	CICLESONIDE INHAL AEROSOL 80 MCG/ACT	44400017003420	Brand
ALVESCO	CICLESONIDE INHAL AEROSOL 160 MCG/ACT	44400017003440	Brand

1 - Submission of medical records (e.g., chart notes) documenting trial and failure (e.g., objective change in symptom control such as Asthma Control Test score, nocturnal awakenings, increased rescue inhaler use, etc.) with an equipotent dose as outlined in the Global Initiative for Asthma (GINA) guidelines, intolerance, or contraindication to a preferred fluticasone product (e.g., Arnuity Ellipta or Flovent)

OR

2 - The requested drug is being used as an add-on inhaler to a preferred fluticasone or mometasone maintenance inhaler for patients who need to "step-up" their asthma treatment plan (i.e. go from Green Zone to Yellow Zone, etc.) due to an increase in symptoms

OR

3 - (For Pulmicort only): Submission of medical records (e.g., chart notes) documenting a current pregnancy

Product Name: Pulmicort Flexhaler, Alvesco			
Approval Length	Approval Length 12 month(s)		
Therapy Stage	Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
	BUDESONIDE INHAL AERO POWD 90 MCG/ACT (BREATH ACTIVATED)	44400015008009	Brand
PULMICORT FLEXHALER	BUDESONIDE INHAL AERO POWD 180 MCG/ACT (BREATH ACTIVATED)	44400015008018	Brand
ALVESCO	CICLESONIDE INHAL AEROSOL 80 MCG/ACT	44400017003420	Brand
ALVESCO	CICLESONIDE INHAL AEROSOL 160 MCG/ACT	44400017003440	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Pulmicort Flexhaler, Alvesco		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
PULMICORT FLEXHALER	BUDESONIDE INHAL AERO POWD 90 MCG/ACT (BREATH ACTIVATED)	44400015008009	Brand
PULMICORT FLEXHALER	BUDESONIDE INHAL AERO POWD 180 MCG/ACT (BREATH ACTIVATED)	44400015008018	Brand
ALVESCO	CICLESONIDE INHAL AEROSOL 80 MCG/ACT	44400017003420	Brand
ALVESCO	CICLESONIDE INHAL AEROSOL 160 MCG/ACT	44400017003440	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting trial and failure (e.g., objective change in symptom control such as Asthma Control Test score, nocturnal awakenings, increased rescue inhaler use, etc.) with an equipotent dose as outlined in the Global Initiative for Asthma (GINA) guidelines, intolerance, or contraindication to a preferred fluticasone product (e.g., Arnuity Ellipta or Flovent)

OR

2 - The requested drug is being used as an add-on inhaler to a preferred fluticasone or mometasone maintenance inhaler for patients who need to "step-up" their asthma treatment plan (i.e. go from Green Zone to Yellow Zone, etc.) due to an increase in symptoms

OR

3 - (For Pulmicort only): Submission of medical records (e.g., chart notes) documenting a current pregnancy

Date	Notes
2/28/2024	Removed Asmanex

Restricted Long-acting Morphine Sulfat	te
The best requirement of the first transfer o	

Prior Authorization Guideline

Guideline ID	GL-131573	
Guideline Name	Restricted Long-acting Morphine Sulfate	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Morphine ER capsules (Kadian and Evinza equivalent)				
Approval Length		12/31/2039		
Guideline T	ype	Prior Authorization - All plans except IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 10 MG		65100055107010	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 20 MG		65100055107020	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 30 MG 65100055107030 Generic		Generic	

MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 50 MG	65100055107040	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 60 MG	65100055107045	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 80 MG	65100055107050	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 100 MG	65100055107060	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 30 MG	65100055207020	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 45 MG	65100055207025	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 60 MG	65100055207030	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 75 MG	65100055207035	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 90 MG	65100055207040	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 120 MG	65100055207050	Generic

1 - Trial and failure, contraindication or intolerance to an equivalent dose of generic extended release morphine tablets (MS Contin equivalent)

Product Name: Morphine ER capsules (Kadian and Evinza equivalent)				
Approval Le	ength	12 month(s)		
Therapy St	age	Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans				
Product Name	Generic Name		GPI	Brand/Generic

MORPHINE SULFATE CAP ER 24HR 10 MG	65100055107010	Generic
MORPHINE SULFATE CAP ER 24HR 20 MG	65100055107020	Generic
MORPHINE SULFATE CAP ER 24HR 30 MG	65100055107030	Generic
MORPHINE SULFATE CAP ER 24HR 50 MG	65100055107040	Generic
MORPHINE SULFATE CAP ER 24HR 60 MG	65100055107045	Generic
MORPHINE SULFATE CAP ER 24HR 80 MG	65100055107050	Generic
MORPHINE SULFATE CAP ER 24HR 100 MG	65100055107060	Generic
MORPHINE SULFATE BEADS CAP ER 24HR 30 MG	65100055207020	Generic
MORPHINE SULFATE BEADS CAP ER 24HR 45 MG	65100055207025	Generic
MORPHINE SULFATE BEADS CAP ER 24HR 60 MG	65100055207030	Generic
MORPHINE SULFATE BEADS CAP ER 24HR 75 MG	65100055207035	Generic
MORPHINE SULFATE BEADS CAP ER 24HR 90 MG	65100055207040	Generic
MORPHINE SULFATE BEADS CAP ER 24HR 120 MG	65100055207050	Generic
	MORPHINE SULFATE CAP ER 24HR 20 MG MORPHINE SULFATE CAP ER 24HR 30 MG MORPHINE SULFATE CAP ER 24HR 50 MG MORPHINE SULFATE CAP ER 24HR 60 MG MORPHINE SULFATE CAP ER 24HR 80 MG MORPHINE SULFATE CAP ER 24HR 100 MG MORPHINE SULFATE BEADS CAP ER 24HR 30 MG MORPHINE SULFATE BEADS CAP ER 24HR 45 MG MORPHINE SULFATE BEADS CAP ER 24HR 60 MG MORPHINE SULFATE BEADS CAP ER 24HR 60 MG MORPHINE SULFATE BEADS CAP ER 24HR 75 MG MORPHINE SULFATE BEADS CAP ER 24HR 90 MG	MORPHINE SULFATE CAP ER 24HR 20 MG MORPHINE SULFATE CAP ER 24HR 30 MG MORPHINE SULFATE CAP ER 24HR 50 MG MORPHINE SULFATE CAP ER 24HR 60 MG MORPHINE SULFATE CAP ER 24HR 60 MG MORPHINE SULFATE CAP ER 24HR 80 MG MORPHINE SULFATE CAP ER 24HR 100 MG MORPHINE SULFATE CAP ER 24HR 100 MG MORPHINE SULFATE CAP ER 24HR 30 MG MORPHINE SULFATE BEADS CAP ER 24HR 30 MG MORPHINE SULFATE BEADS CAP ER 24HR 45 MG MORPHINE SULFATE BEADS CAP ER 24HR 45 MG MORPHINE SULFATE BEADS CAP ER 24HR 60 MG MORPHINE SULFATE BEADS CAP ER 24HR 60 MG MORPHINE SULFATE BEADS CAP ER 24HR 75 MG MORPHINE SULFATE BEADS CAP ER 24HR 75 MG MORPHINE SULFATE BEADS CAP ER 24HR 90 MG MORPHINE SULFATE BEADS CAP ER 24HR 90 MG 65100055207040

1 - Trial and failure, contraindication or intolerance to an equivalent dose of generic extended-release morphine tablets (MS Contin equivalent)

OR

2 - (Minnesota plans only) – Member has stage four metastatic cancer and the requested drug is being used to treat cancer-related pain

Product Name: Morphine ER capsules (Kadian and Evinza equivalent)		
Approval Length	Approval Length 12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 10 MG	65100055107010	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 20 MG	65100055107020	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 30 MG	65100055107030	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 50 MG	65100055107040	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 60 MG	65100055107045	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 80 MG	65100055107050	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE CAP ER 24HR 100 MG	65100055107060	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 30 MG	65100055207020	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 45 MG	65100055207025	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 60 MG	65100055207030	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 75 MG	65100055207035	Generic
MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 90 MG	65100055207040	Generic

MORPHINE SULFATE ER	MORPHINE SULFATE BEADS CAP ER 24HR 120 MG	65100055207050	Generic	
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1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

2. Revision History

Date	Notes
10/13/2023	2024 New Implementation

Restricted Methotrexate Injection	
(2) The contract of the contra	

Prior Authorization Guideline

Guideline ID	GL-131419
Guideline Name	Restricted Methotrexate Injection
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Rasuvo, Otrexup, Reditrex		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 7.5 MG/0.15ML	6625005000D510	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 10 MG/0.2ML	6625005000D512	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 12.5 MG/0.25ML	6625005000D517	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 15 MG/0.3ML	6625005000D519	Brand

RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 17.5 MG/0.35ML	6625005000D522	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 20 MG/0.4ML	6625005000D525	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 20 MG/0.4ML	6625005000D525	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 22.5 MG/0.45ML	6625005000D527	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 25 MG/0.5ML	6625005000D535	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 30 MG/0.6ML	6625005000D545	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 10 MG/0.4ML	6625005000D515	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 12.5 MG/0.4ML	6625005000D518	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 15 MG/0.4ML	6625005000D520	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 17.5 MG/0.4ML	6625005000D523	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 22.5 MG/0.4ML	6625005000D528	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 25 MG/0.4ML	6625005000D530	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 7.5 MG/0.3ML	6625005000E508	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 10 MG/0.4ML	6625005000E510	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 12.5 MG/0.5ML	6625005000E512	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 15 MG/0.6ML	6625005000E515	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 17.5 MG/0.7ML	6625005000E522	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 20 MG/0.8ML	6625005000E526	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 22.5 MG/0.9ML	6625005000E532	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 25 MG/ML	6625005000E536	Brand

1 - Documented disability that does not allow administration of methotrexate from conventional vials utilizing conventional syringes

AND

2 - The person or a family member/caregiver are self-administering the medication

Product Name: Rasuvo, Otrexup, Reditrex		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name GPI Brand/G		Brand/Generic
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 7.5 MG/0.15ML	6625005000D510	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 10 MG/0.2ML	6625005000D512	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 12.5 MG/0.25ML	6625005000D517	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 15 MG/0.3ML	6625005000D519	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 17.5 MG/0.35ML	6625005000D522	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 20 MG/0.4ML	6625005000D525	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 20 MG/0.4ML	6625005000D525	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 22.5 MG/0.45ML	6625005000D527	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 25 MG/0.5ML	6625005000D535	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 30 MG/0.6ML	6625005000D545	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 10 MG/0.4ML	6625005000D515	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 12.5 MG/0.4ML	6625005000D518	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 15 MG/0.4ML	6625005000D520	Brand

OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 17.5 MG/0.4ML	6625005000D523	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 22.5 MG/0.4ML	6625005000D528	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 25 MG/0.4ML	6625005000D530	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 7.5 MG/0.3ML	6625005000E508	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 10 MG/0.4ML	6625005000E510	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 12.5 MG/0.5ML	6625005000E512	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 15 MG/0.6ML	6625005000E515	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 17.5 MG/0.7ML	6625005000E522	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 20 MG/0.8ML	6625005000E526	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 22.5 MG/0.9ML	6625005000E532	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 25 MG/ML	6625005000E536	Brand

1 - Documentation from the past 12 months that the person is continuing therapy with the requested drug

Product Name: Rasuvo, Otrexup, Reditrex				
Approval Length 12/31/2039				
Guideline ⁻	Туре	Prior Authorization-All plans except	IL and MN	
Product Name	Generic Name GPI Brand/Gener		Brand/Generic	
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 7.5 MG/0.15ML		6625005000D510	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 10 MG/0.2ML		6625005000D512	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 12.5 MG/0.25ML 66.		6625005000D517	Brand
RASUVO	SUVO METHOTREXATE SOLN PF AUTO-INJECTOR 15 6625005000D519 Brand		Brand	

RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 17.5 MG/0.35ML	6625005000D522	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 20 MG/0.4ML	6625005000D525	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 20 MG/0.4ML	6625005000D525	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 22.5 MG/0.45ML	6625005000D527	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 25 MG/0.5ML	6625005000D535	Brand
RASUVO	METHOTREXATE SOLN PF AUTO-INJECTOR 30 MG/0.6ML	6625005000D545	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 10 MG/0.4ML	6625005000D515	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 12.5 MG/0.4ML	6625005000D518	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 15 MG/0.4ML	6625005000D520	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 17.5 MG/0.4ML	6625005000D523	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 22.5 MG/0.4ML	6625005000D528	Brand
OTREXUP	METHOTREXATE SOLN PF AUTO-INJECTOR 25 MG/0.4ML	6625005000D530	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 7.5 MG/0.3ML	6625005000E508	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 10 MG/0.4ML	6625005000E510	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 12.5 MG/0.5ML	6625005000E512	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 15 MG/0.6ML	6625005000E515	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 17.5 MG/0.7ML	6625005000E522	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 20 MG/0.8ML	6625005000E526	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 22.5 MG/0.9ML	6625005000E532	Brand
REDITREX	METHOTREXATE SOLN PREFILLED SYRINGE 25 MG/ML	6625005000E536	Brand

1 - Documented disability that does not allow administration of methotrexate from conventional vials utilizing conventional syringes

AND

2 - The person or a family member/caregiver are self-administering the medication

2. Revision History

Date	Notes
10/24/2023	New Program

Restricted Minocycline ER			
g - handrage considered hand and some some some and basic problem confluenced.			

Prior Authorization Guideline

Guideline ID	GL-137244
Guideline Name	Restricted Minocycline ER
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Brand Coremino, Generic Solodyn, generic minocycline extended release		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 55 MG	04000040107522	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 65 MG	04000040107525	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 80 MG	04000040107528	Generic

MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 105 MG	04000040107533	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 115 MG	04000040107535	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 45 MG	04000040107520	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 45 MG	04000040107520	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 90 MG	04000040107530	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 90 MG	04000040107530	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 135 MG	04000040107540	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 135 MG	04000040107540	Generic

1 - Trial and intolerance (clinically significant side effects that limit use) from a preferred minocycline product at equivalent doses

AND

2 - The prescriber provides an evidence-based clinical rationale why a different result would be expected with use of minocycline ER

Product Name: Brand Coremino, Generic Solodyn, generic minocycline extended release			
Approval Length 12 month(s)			
Therapy Stage Reauthorization			
Guideline Type	Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 55 MG	04000040107522	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 65 MG	04000040107525	Generic

MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 80 MG	04000040107528	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 105 MG	04000040107533	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 115 MG	04000040107535	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 45 MG	04000040107520	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 45 MG	04000040107520	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 90 MG	04000040107530	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 90 MG	04000040107530	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 135 MG	04000040107540	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 135 MG	04000040107540	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Brand Coremino, Generic Solodyn, generic minocycline extended release		
Approval Length One fill		
Guideline Type Prior Authorization-All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 55 MG	04000040107522	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 65 MG	04000040107525	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 80 MG	04000040107528	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 105 MG	04000040107533	Generic

MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 115 MG	04000040107535	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 45 MG	04000040107520	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 45 MG	04000040107520	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 90 MG	04000040107530	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 90 MG	04000040107530	Generic
COREMINO	MINOCYCLINE HCL TAB ER 24HR 135 MG	04000040107540	Generic
MINOCYCLINE HYDROCHLORIDE ER	MINOCYCLINE HCL TAB ER 24HR 135 MG	04000040107540	Generic

1 - Trial and intolerance (clinically significant side effects that limit use) from a preferred minocycline product at equivalent doses

AND

2 - The prescriber provides an evidence-based clinical rationale why a different result would be expected with use of minocycline ER

2. Revision History

Date	Notes
12/6/2023	New program

Restricted Non-preferred Medications
(S) behavioral mental balance behavior to the color of th

Prior Authorization Guideline

Guideline ID	GL-144916	
Guideline Name	Restricted Non-preferred Medications	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	5/18/2016
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Restricted Non-preferred Drugs Greater than or equal to 5 therapeutic alternatives					
Diagnosis	Diagnosis Illinois Plan ONLY				
Approval Length	Approval Length 12 month(s)				
Guideline Type Administrative					
Product Name Generic Name GPI Brand/Gen			Brand/Generic		
Approval Criteria					
1 - Both of the following:					

- **1.1** The requested medication has a diagnosis that is one of the following:
 - Food Drug Administration (FDA)-approved indication
 - A medically accepted indication that is supported by nationally recognized compendia (see table of Compendia Requirements within the Background Section)

AND

- **1.2** For drugs with greater than or equal to 5 therapeutic alternatives (in the same drug class or different drug class that treats the same disease as the requested drug) one of the following:
- **1.2.1** Trial and failure or intolerance, or contraindication to at least 2 preferred alternatives in the same drug class

OR

1.2.2 Trial and failure to at least 2 preferred alternatives from other drug classes that treat the same disease if there are not 2 formulary alternatives in the same drug class and provider attests that it is clinically appropriate treatment of the diagnosis

OR

1.2.3 Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on nonpreferred therapy, will have coverage under their drug benefit with submission of medical records (e.g., chart notes) documenting symptom improvement or disease stability

OR

2 - The requested drug is FDA approved for the treatment of tick-borne disease

Product Name: Restricted Non-preferred Drugs less than 5 therapeutic alternatives		
Diagnosis	Illinois Plan ONLY	
Approval Length	12 month(s)	
Guideline Type	Administrative	

Product Name	Generic Name	GPI	Brand/Generic
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- **1** Both of the following:
- **1.1** The requested medication has a diagnosis that is one of the following:
 - Food Drug Administration (FDA)-approved indication
 - A medically accepted indication that is supported by nationally recognized compendia (see table of Compendia Requirements within the Background Section)

AND

- **1.2** For drugs with less than 5 therapeutic alternatives (in the same drug class or different drug class that treats the same disease as the requested drug) one of the following:
- **1.2.1** Trial and failure or intolerance, or contraindication to at least 1 preferred alternative in the same drug class

OR

1.2.2 Trial and failure to at least 1 preferred alternative from other drug classes that treat the same disease if there are not 1 formulary alternatives in the same drug class and provider attests that it is clinically appropriate treatment of the diagnosis

OR

1.2.3 Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on nonpreferred therapy, will have coverage under their drug benefit with submission of medical records (e.g., chart notes) documenting symptom improvement or disease stability

OR

2 - The requested drug is for the treatment of tick-borne disease

Product Name: Restricted Non-preferred Drugs Greater than or equal to 5 therapeutic alternatives			
Diagnosis	Minnesota Plans ONLY		
Approval Length	12 month(s)		
Therapy Stage	Initial Authorization		
Guideline Type	Administrative		

Product Name	Generic Name	GPI	Brand/Generic
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- 1 Both of the following:
- **1.1** The requested medication has a diagnosis that is one of the following:
 - Food Drug Administration (FDA)-approved indication
 - A medically accepted indication that is supported by nationally recognized compendia (see table of Compendia Requirements within the Background Section)

AND

- **1.2** For drugs with greater than or equal to 5 therapeutic alternatives (in the same drug class or different drug class that treats the same disease as the requested drug) one of the following:
- **1.2.1** Trial and failure or intolerance, or contraindication to at least 2 preferred alternatives in the same drug class

OR

1.2.2 Trial and failure to at least 2 preferred alternatives from other drug classes that treat the same disease if there are not 2 formulary alternatives in the same drug class and provider attests that it is clinically appropriate treatment of the diagnosis

OR

1.2.3 Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on nonpreferred therapy, will have coverage under

their drug benefit with submission of medical recor	ds (e.g., chart notes) documenting
symptom improvement or disease stability	

OR

- 2 Both of the following:
- **2.1** Provider attests the patient has emotional disturbance or mental illness

AND

2.2 Prescriber submits medical records (e.g., chart notes) that all equivalent drugs in the formulary were considered and it has been determined that the drug prescribed will best treat the person's condition

OR

3 - For continuation of care: the person has been treated for 90 days prior to the change, the medication is working, and the prescriber attests the drug prescribed will best treat the person's condition

OR

4 - The requested drug is for stage four metastatic cancer and prescribed drug is used for cancer related treatment including but not limited to: pain, constipation, nausea, or prevention/treatment of infection

Product Name: Restricted Non-preferred Drugs less than 5 therapeutic alternatives					
Diagnosis Minnesota Plans ONLY					
Approval Length 12 month(s)					
Therapy Stage Initial Authorization					
Guideline Type		Administrative			
Product Name Generic Name		ric Name	GPI	Brand/Generic	
Approval Criteria					

- **1** Both of the following:
- **1.1** The requested medication has a diagnosis that is one of the following:
 - Food Drug Administration (FDA)-approved indication
 - A medically accepted indication that is supported by nationally recognized compendia (see table of Compendia Requirements within the Background Section)

AND

- **1.2** For drugs with less than 5 therapeutic alternatives (in the same drug class or different drug class that treats the same disease as the requested drug) one of the following:
- **1.2.1** Trial and failure or intolerance, or contraindication to at least 1 preferred alternative in the same drug class

OR

1.2.2 Trial and failure to at least 1 preferred alternative from other drug classes that treat the same disease if there are not 1 formulary alternatives in the same drug class and provider attests that it is clinically appropriate treatment of the diagnosis

OR

- 2 Both of the following:
- **2.1** Provider attests the patient has emotional disturbance or mental illness

AND

2.2 Submission of medical records (e.g., chart notes) that all equivalent drugs in the formulary were considered and it has been determined that the drug prescribed will best treat the person's condition

OR

3 - For continuation of care: the person has been treated for 90 days prior to the change, the

medication is working, and the prescriber attests the drug prescribed will best treat the person's condition

OR

4 - The requested drug is for stage four metastatic cancer and prescribed drug is used for cancer related treatment including but not limited to: pain, constipation, nausea, or prevention/treatment of infection

Product Name: Restricted Non-preferred Drugs greater than or equal to 5 therapeutic alternatives			
Approval Length 12 month(s)			
Therapy Stage	Initial Authorization		
Guideline Type Administrative - All plans except IL and MN			
Product Name Generic Name GPI Brand/Generic			

Approval Criteria

- **1** The requested medication has a diagnosis that is one of the following:
 - Food Drug Administration (FDA)-approved indication
 - A medically accepted indication that is supported by nationally recognized compendia (see table of Compendia Requirements within the Background Section)

AND

- **2** For drugs with greater than or equal to 5 therapeutic alternatives (in the same drug class or different drug class that treats the same disease as the requested drug) one of the following:
- **2.1** Trial and failure or intolerance, or contraindication to at least 2 preferred alternatives in the same drug class

OR

2.2 Trial and failure to at least 2 preferred alternatives from other drug classes that treat the

same disease if there are not 2 formulary alternatives in the same drug class and provider attests that it is clinically appropriate treatment of the diagnosis

OR

2.3 Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on nonpreferred therapy, will have coverage under their drug benefit with submission of medical records (e.g., chart notes) documenting symptom improvement or disease stability

Product Name: Restricted Non-preferred Drugs less than 5 therapeutic alternatives				
Approval Length 12 month(s)				
Therapy Stage		Initial Authorization		
Guideline Type Administrative - All other plans except IL and MN				
Product Name Generic Name GPI Brand/Generic				

Approval Criteria

- **1** The requested medication has a diagnosis that is one of the following:
 - Food Drug Administration (FDA)-approved indication
 - A medically accepted indication that is supported by nationally recognized compendia (see table of Compendia Requirements within the Background Section)

AND

- **2** For drugs with less than 5 therapeutic alternatives (in the same drug class or different drug class that treats the same disease as the requested drug) one of the following:
- **2.1** Trial and failure or intolerance, or contraindication to at least 1 preferred alternative in the same drug class

OR

2.2 Trial and failure to at least 1 preferred alternative from other drug classes that treat the same disease if there are not 1 formulary alternatives in the same drug class and provider attests that it is clinically appropriate treatment of the diagnosis

OR

2.3 Member is new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who are established on nonpreferred therapy, will have coverage under their drug benefit with submission of medical records (e.g., chart notes) documenting symptom improvement or disease stability

Product Name: All Indications above			
Diagnosis	All Plans		
Approval Length	12 month(s)		
Therapy Stage	Reauthorization		
Guideline Type Administrative			
Product Name Generic Name GPI Brand/Generi			

Approval Criteria

1 - Paid claims or submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

2. Revision History

Date	Notes
3/27/2024	Updated approval duration to "months"

Restricted Nonpreferred Proton Pump Inhibitor (F							
(g) beliefung-manadaya balan sa kata mana anna a dan ka kata ka anna ka mana ka ata ka							

Prior Authorization Guideline

Guideline ID	GL-131574	
Guideline Name	Restricted Nonpreferred Proton Pump Inhibitor (PPI)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic dexlansoprazole					
Approval Length	12/31/2039	12/31/2039			
Guideline Type	Prior Authorization - All Plans excep	Prior Authorization - All Plans except IL and MN Plans			
Product Name	Generic Name	GPI	Brand/Generic		
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 30 MG	49270020006520	Generic		
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 60 MG	49270020006530	Generic		

Approval Criteria

1 - Trial and failure, contraindication, or intolerance to at least THREE of the following preferred PPI alternatives:

- omeprazole pantoprazole
- lansoprazole
- rabeprazole tablets
- esomeprazole capsules

Product Name: Generic dexlansoprazole		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL Plan	

Product Name	Generic Name	GPI	Brand/Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 30 MG	49270020006520	Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 60 MG	49270020006530	Generic

- 1 Trial and failure, contraindication, or intolerance to at least THREE of the following preferred PPI alternatives:
 - omeprazole
 - pantoprazole
 - lansoprazole
 - rabeprazole tablets
 - esomeprazole capsules

Product Name: Generic dexlansoprazole					
Approval Length		12 month(s)			
Therapy Stage		Initial Authorization			
Guideline Type		Prior Authorization - MN Plan			
Product Name	Ge	eneric Name	GPI	Brand/Generic	
DEXLANSOPRAZOLE		XLANSOPRAZOLE CAP DELAYED LEASE 30 MG	49270020006520	Generic	
DEXLANSOPRAZOLE		XLANSOPRAZOLE CAP DELAYED LEASE 60 MG	49270020006530	Generic	

- **1** Trial and failure, contraindication, or intolerance to at least THREE of the following preferred PPI alternatives:
 - omeprazole
 - pantoprazole
 - lansoprazole
 - rabeprazole tablets
 - esomeprazole capsules

OR

2 - Diagnosis of stage four metastatic cancer and the requested drug is being used as supportive care to treat symptoms directly related to their cancer or chemotherapy regimen

Product Name: Generic dexlansoprazole		
Diagnosis	Quantity Exception	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 30 MG	49270020006520	Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 60 MG	49270020006530	Generic

Approval Criteria

1 - Member has extraesophageal symptoms

OR

2 - The requested dosing schedule cannot be met using commercially available dose forms within the quantity limit and the prescriber provides an evidence-based rationale for using a dose outside of the quantity limit

OR

3 - For use in compounded prescriptions where the quantity limit would prevent the pharmacy from being able to make the compounded formulation

Product Name: Generic dexlansoprazole	
Diagnosis	Quantity Exception
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 30 MG	49270020006520	Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 60 MG	49270020006530	Generic

Approval Criteria

1 - Member has extraesophageal symptoms

OR

2 - The requested dosing schedule cannot be met using commercially available dose forms within the quantity limit and the prescriber provides an evidence-based rationale for using a dose outside of the quantity limit

OR

3 - For use in compounded prescriptions where the quantity limit would prevent the pharmacy from being able to make the compounded formulation

Product Name: Generic dexlansoprazole	
Approval Length	12 month(s)

Therapy Stage	Reauthorization		
Guideline Type	Prior Authorization - IL and MN Plans		
Product Name	Generic Name	GPI	Brand/Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 30 MG	49270020006520	Generic
DEXLANSOPRAZOLE	DEXLANSOPRAZOLE CAP DELAYED RELEASE 60 MG	49270020006530	Generic

1 - Prescriber provides clinical documentation from the past 12 months that the person is continuing therapy with the requested drug

2. Revision History

Date	Notes
10/6/2023	2024 New Implementation

Restricted Oral Antipsychotics	Step
The Mandang and Antiques Track to the last area and areas, a seek with district project for smooth and common	

Prior Authorization Guideline

Guideline ID	GL-127882
Guideline Name	Restricted Oral Antipsychotics Step
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Asenapine, Vraylar, Fanapt, Caplyta, Generic Lurasidone, Rexulti		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 2.5 MG (BASE EQUIV)	59155015100710	Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 5 MG (BASE EQUIV)	59155015100720	Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 10 MG (BASE EQUIV)	59155015100730	Generic
VRAYLAR	CARIPRAZINE HCL CAP THERAPY PACK 1.5 MG (1) & 3 MG (6)	5940001810B220	Brand
VRAYLAR	CARIPRAZINE HCL CAP 1.5 MG (BASE EQUIVALENT)	59400018100120	Brand

VRAYLAR	CARIPRAZINE HCL CAP 3 MG (BASE EQUIVALENT)	59400018100130	Brand
VRAYLAR	CARIPRAZINE HCL CAP 4.5 MG (BASE EQUIVALENT)	59400018100140	Brand
VRAYLAR	CARIPRAZINE HCL CAP 6 MG (BASE EQUIVALENT)	59400018100150	Brand
FANAPT	ILOPERIDONE TAB 1 MG	59070035000310	Brand
FANAPT	ILOPERIDONE TAB 2 MG	59070035000320	Brand
FANAPT	ILOPERIDONE TAB 4 MG	59070035000340	Brand
FANAPT	ILOPERIDONE TAB 6 MG	59070035000360	Brand
FANAPT	ILOPERIDONE TAB 8 MG	59070035000380	Brand
FANAPT	ILOPERIDONE TAB 10 MG	59070035000385	Brand
FANAPT	ILOPERIDONE TAB 12 MG	59070035000390	Brand
FANAPT TITRATION PACK	ILOPERIDONE TAB 1 MG & 2 MG & 4 MG & 6 MG TITRATION PAK	59070035006320	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 10.5 MG	59400022400110	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 21 MG	59400022400115	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 42 MG	59400022400120	Brand
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 20 MG	59400023100310	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 40 MG	59400023100320	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 60 MG	59400023100330	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 80 MG	59400023100340	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 120 MG	59400023100350	Generic
REXULTI	BREXPIPRAZOLE TAB 0.25 MG	59250020000310	Brand
REXULTI	BREXPIPRAZOLE TAB 0.5 MG	59250020000320	Brand
REXULTI	BREXPIPRAZOLE TAB 1 MG	59250020000330	Brand
REXULTI	BREXPIPRAZOLE TAB 2 MG	59250020000340	Brand
REXULTI	BREXPIPRAZOLE TAB 3 MG	59250020000350	Brand
REXULTI	BREXPIPRAZOLE TAB 4 MG	59250020000360	Brand

1 - Trial and failure, contraindication, or intolerance of a preferred second-generation antipsychotic (i.e., aripiprazole, olanzapine, risperidone, quetiapine, or ziprasidone).

OR

2 - For Minnesota Plans Only: If requested drug is prescribed for emotional disturbance or mental illness, approve if submission of medical records (e.g., chart notes) documenting that all equivalent drugs in the formulary were considered and it has been determined that the drug prescribed will best treat the member's condition

Product Name: Generic Asenapine, Vraylar, Fanapt, Caplyta, Generic Lurasidone, Rexulti		
Approval Length	Approval Length 12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 2.5 MG (BASE EQUIV)	59155015100710	Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 5 MG (BASE EQUIV)	59155015100720	Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 10 MG (BASE EQUIV)	59155015100730	Generic
VRAYLAR	CARIPRAZINE HCL CAP THERAPY PACK 1.5 MG (1) & 3 MG (6)	5940001810B220	Brand
VRAYLAR	CARIPRAZINE HCL CAP 1.5 MG (BASE EQUIVALENT)	59400018100120	Brand
VRAYLAR	CARIPRAZINE HCL CAP 3 MG (BASE EQUIVALENT)	59400018100130	Brand
VRAYLAR	CARIPRAZINE HCL CAP 4.5 MG (BASE EQUIVALENT)	59400018100140	Brand
VRAYLAR	CARIPRAZINE HCL CAP 6 MG (BASE EQUIVALENT)	59400018100150	Brand
FANAPT	ILOPERIDONE TAB 1 MG	59070035000310	Brand
FANAPT	ILOPERIDONE TAB 2 MG	59070035000320	Brand
FANAPT	ILOPERIDONE TAB 4 MG	59070035000340	Brand
FANAPT	ILOPERIDONE TAB 6 MG	59070035000360	Brand
FANAPT	ILOPERIDONE TAB 8 MG	59070035000380	Brand
FANAPT	ILOPERIDONE TAB 10 MG	59070035000385	Brand

FANAPT	ILOPERIDONE TAB 12 MG	59070035000390	Brand
FANAPT TITRATION PACK	ILOPERIDONE TAB 1 MG & 2 MG & 4 MG & 6 MG TITRATION PAK	59070035006320	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 10.5 MG	59400022400110	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 21 MG	59400022400115	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 42 MG	59400022400120	Brand
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 20 MG	59400023100310	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 40 MG	59400023100320	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 60 MG	59400023100330	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 80 MG	59400023100340	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 120 MG	59400023100350	Generic
REXULTI	BREXPIPRAZOLE TAB 0.25 MG	59250020000310	Brand
REXULTI	BREXPIPRAZOLE TAB 0.5 MG	59250020000320	Brand
REXULTI	BREXPIPRAZOLE TAB 1 MG	59250020000330	Brand
REXULTI	BREXPIPRAZOLE TAB 2 MG	59250020000340	Brand
REXULTI	BREXPIPRAZOLE TAB 3 MG	59250020000350	Brand
REXULTI	BREXPIPRAZOLE TAB 4 MG	59250020000360	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

OR

- **2** For Minnesota Plans Only Both of the following for continuation of care (i.e. formulary changes or new member [(as evidenced by coverage effective date of less than or equal to 90 days]):
- 2.1 The member has been treated with the drug for 90 days prior to the change

AND

2.2 Submission of medical records (e.g., chart notes) of documentation that the drug prescribed will best treat the patient's condition

Product Name: Generic Asenapine, Vraylar, Fanapt, Caplyta, Generic Lurasidone, Rexulti		
Approval Length	12/31/2039	
Guideline Type	Step Therapy - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 2.5 MG (BASE EQUIV)	59155015100710	Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 5 MG (BASE EQUIV)	59155015100720	Generic
ASENAPINE MALEATE SL	ASENAPINE MALEATE SL TAB 10 MG (BASE EQUIV)	59155015100730	Generic
VRAYLAR	CARIPRAZINE HCL CAP THERAPY PACK 1.5 MG (1) & 3 MG (6)	5940001810B220	Brand
VRAYLAR	CARIPRAZINE HCL CAP 1.5 MG (BASE EQUIVALENT)	59400018100120	Brand
VRAYLAR	CARIPRAZINE HCL CAP 3 MG (BASE EQUIVALENT)	59400018100130	Brand
VRAYLAR	CARIPRAZINE HCL CAP 4.5 MG (BASE EQUIVALENT)	59400018100140	Brand
VRAYLAR	CARIPRAZINE HCL CAP 6 MG (BASE EQUIVALENT)	59400018100150	Brand
FANAPT	ILOPERIDONE TAB 1 MG	59070035000310	Brand
FANAPT	ILOPERIDONE TAB 2 MG	59070035000320	Brand
FANAPT	ILOPERIDONE TAB 4 MG	59070035000340	Brand
FANAPT	ILOPERIDONE TAB 6 MG	59070035000360	Brand
FANAPT	ILOPERIDONE TAB 8 MG	59070035000380	Brand
FANAPT	ILOPERIDONE TAB 10 MG	59070035000385	Brand
FANAPT	ILOPERIDONE TAB 12 MG	59070035000390	Brand
FANAPT TITRATION PACK	ILOPERIDONE TAB 1 MG & 2 MG & 4 MG & 6 MG TITRATION PAK	59070035006320	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 10.5 MG	59400022400110	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 21 MG	59400022400115	Brand
CAPLYTA	LUMATEPERONE TOSYLATE CAP 42 MG	59400022400120	Brand
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 20 MG	59400023100310	Generic

LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 40 MG	59400023100320	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 60 MG	59400023100330	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 80 MG	59400023100340	Generic
LURASIDONE HYDROCHLORIDE	LURASIDONE HCL TAB 120 MG	59400023100350	Generic
REXULTI	BREXPIPRAZOLE TAB 0.25 MG	59250020000310	Brand
REXULTI	BREXPIPRAZOLE TAB 0.5 MG	59250020000320	Brand
REXULTI	BREXPIPRAZOLE TAB 1 MG	59250020000330	Brand
REXULTI	BREXPIPRAZOLE TAB 2 MG	59250020000340	Brand
REXULTI	BREXPIPRAZOLE TAB 3 MG	59250020000350	Brand
REXULTI	BREXPIPRAZOLE TAB 4 MG	59250020000360	Brand

1 - Trial and failure, contraindication, or intolerance of a preferred second-generation antipsychotic (i.e., aripiprazole, olanzapine, risperidone, quetiapine, or ziprasidone).

Product Name: Generic Aripiprazole ODT		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ARIPIPRAZOLE ODT	ARIPIPRAZOLE ORALLY DISINTEGRATING TAB 10 MG	59250015007220	Generic
ARIPIPRAZOLE ODT	ARIPIPRAZOLE ORALLY DISINTEGRATING TAB 15 MG	59250015007230	Generic

Approval Criteria

- 1 Both of the following:
 - **1.1** Trial and failure, contraindication, or intolerance of aripiprazole tablets.

AND

1.2 Trial and failure, contraindication, or intolerance of one other preferred antipsychotic (i.e., olanzapine, risperidone, quetiapine, ziprasidone).

OR

2 - For Minnesota Plans Only: If requested drug is prescribed for emotional disturbance or mental illness, approve if submission of medical records (e.g., chart notes) documenting that all equivalent drugs in the formulary were considered and it has been determined that the drug prescribed will best treat the member's condition

Product Name: Generic Aripiprazole ODT		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type	Step Therapy - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ARIPIPRAZOLE ODT	ARIPIPRAZOLE ORALLY DISINTEGRATING TAB 10 MG	59250015007220	Generic
ARIPIPRAZOLE ODT	ARIPIPRAZOLE ORALLY DISINTEGRATING TAB 15 MG	59250015007230	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

OR

- **2** For Minnesota Plans Only Both of the following for continuation of care (i.e. formulary changes or new member [(as evidenced by coverage effective date of less than or equal to 90 days]):
- **2.1** The member has been treated with the drug for 90 days prior to the change

AND

2.2 Submission of medical records (e.g., chart notes) of documentation that the drug prescribed will best treat the patient's condition

Product Name: Generic Aripiprazole ODT	
Approval Length 12/31/2039	
Guideline Type	Step Therapy - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ARIPIPRAZOLE ODT	ARIPIPRAZOLE ORALLY DISINTEGRATING TAB 10 MG	59250015007220	Generic
ARIPIPRAZOLE ODT	ARIPIPRAZOLE ORALLY DISINTEGRATING TAB 15 MG	59250015007230	Generic

Approval Criteria

1 - Trial and failure, contraindication, or intolerance of aripiprazole tablets.

AND

2 - Trial and failure, contraindication, or intolerance of one other preferred antipsychotic (i.e., olanzapine, risperidone, quetiapine, ziprasidone).

2. Revision History

Date	Notes
8/25/2023	New Program

ł	Restricted Oral Oncology Drug	
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Prior Authorization Guideline

Guideline ID	GL-145303
Guideline Name	Restricted Oral Oncology Drug
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	12/1/2018
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Akeega, Alunbrig, Ayvakit, Balversa, Braftovi, Brukinsa, Calquence, Caprelsa, Cometriq, Copiktra, Daurismo, Erleada, Fotivda, Gavreto, Iclusig, Idhifa, Inqovi, Jaypirca, Krazati, Lenvima, Lorbrena, Lumakras, Lytgobi, Mektovi, Nerlynx, Orgovyx, Orserdu, Pemazyre, Qinlock, Rezlidhia, Rydapt, Scemblix, Talzenna, Tepmetko, Tibsovo, Turalio, Vanflyta, Vitrakvi, Vizimpro, Vonjo, Welireg, Xospata, Xpovio, Yonsa Approval Length 12 month(s) Therapy Stage **Initial Authorization** Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans Product Generic Name Brand/Generic **GPI** Name YONSA ABIRATERONE ACETATE MICRONIZED TAB 125 Brand 21406010250310 MG

CALQUENCE	ACALABRUTINIB MALEATE TAB 100 MG	21532103500320	Brand
KRAZATI	ADAGRASIB TAB 200 MG	21532410000320	Brand
ERLEADA	APALUTAMIDE TAB 60 MG	21402410000320	Brand
ERLEADA	APALUTAMIDE TAB 240 MG	21402410000360	Brand
SCEMBLIX	ASCIMINIB HCL TAB 20 MG	21531806100320	Brand
SCEMBLIX	ASCIMINIB HCL TAB 40 MG	21531806100340	Brand
AYVAKIT	AVAPRITINIB TAB 25 MG	21490009000310	Brand
AYVAKIT	AVAPRITINIB TAB 50 MG	21490009000315	Brand
AYVAKIT	AVAPRITINIB TAB 100 MG	21490009000320	Brand
AYVAKIT	AVAPRITINIB TAB 200 MG	21490009000330	Brand
AYVAKIT	AVAPRITINIB TAB 300 MG	21490009000340	Brand
WELIREG	BELZUTIFAN TAB 40 MG	21421020000320	Brand
MEKTOVI	BINIMETINIB TAB 15 MG	21533520000320	Brand
ALUNBRIG	BRIGATINIB TAB INITIATION THERAPY PACK 90 MG & 180 MG	2153051000B720	Brand
ALUNBRIG	BRIGATINIB TAB 30 MG	21530510000330	Brand
ALUNBRIG	BRIGATINIB TAB 90 MG	21530510000350	Brand
ALUNBRIG	BRIGATINIB TAB 180 MG	21530510000365	Brand
COMETRIQ	CABOZANTINIB S-MALATE CAP 3 X 20 MG (60 MG DOSE) KIT	21533010106460	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 1 X 20 MG (100 DOSE) KIT	21533010106470	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 3 X 20 MG (140 DOSE) KIT	21533010106480	Brand
VIZIMPRO	DACOMITINIB TAB 15 MG	21360019000320	Brand
VIZIMPRO	DACOMITINIB TAB 30 MG	21360019000330	Brand
VIZIMPRO	DACOMITINIB TAB 45 MG	21360019000340	Brand
INQOVI	DECITABINE-CEDAZURIDINE TAB 35-100 MG	21990002250320	Brand
COPIKTRA	DUVELISIB CAP 15 MG	21538030000120	Brand
COPIKTRA	DUVELISIB CAP 25 MG	21538030000130	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 86 MG	21403720100320	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 345 MG	21403720100340	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 50 MG (BASE EQUIVALENT)	21535030200320	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 100 MG (BASE EQUIVALENT)	21535030200340	Brand

BRAFTOVI	ENCORAFENIB CAP 75 MG	21532040000130	Brand
BALVERSA	ERDAFITINIB TAB 3 MG	21532225000320	Brand
BALVERSA	ERDAFITINIB TAB 4 MG	21532225000325	Brand
BALVERSA	ERDAFITINIB TAB 5 MG	21532225000330	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (12 MG DAILY DOSE)	2153222800B720	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (16 MG DAILY DOSE)	2153222800B725	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (20 MG DAILY DOSE)	2153222800B730	Brand
XOSPATA	GILTERITINIB FUMARATE TABLET 40 MG (BASE EQUIVALENT)	21533020200320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 25 MG (BASE EQUIVALENT)	21370030300320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 100 MG (BASE EQUIVALENT)	21370030300335	Brand
TIBSOVO	IVOSIDENIB TAB 250 MG	21534940000320	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 25 MG (BASE EQUIVALENT)	21533835200120	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 100 MG (BASE EQUIVALENT)	21533835200150	Brand
VITRAKVI	LAROTRECTINIB SULFATE ORAL SOLN 20 MG/ML (BASE EQUIVALENT)	21533835202020	Brand
LENVIMA 4 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 4 MG (4 MG DAILY DOSE)	2133505420B210	Brand
LENVIMA 8 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 4 MG (8 MG DAILY DOSE)	2133505420B215	Brand
LENVIMA 10 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 MG (10 MG DAILY DOSE)	2133505420B220	Brand
LENVIMA 12MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 3 X 4 MG (12 MG DAILY DOSE)	2133505420B223	Brand
LENVIMA 20 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 10 MG (20 MG DAILY DOSE)	2133505420B230	Brand
LENVIMA 14 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 & 4 MG (14 MG DAILY DOSE)	2133505420B240	Brand
LENVIMA 18 MG DAILY DOSE	LENVATINIB CAP THER PACK 10 MG & 2 X 4 MG (18 MG DAILY DOSE)	2133505420B244	Brand

LENVIMA 24 MG DAILY DOSE	LENVATINIB CAP THER PACK 2 X 10 MG & 4 MG (24 MG DAILY DOSE)	2133505420B250	Brand
LORBRENA	LORLATINIB TAB 25 MG	21530556000320	Brand
LORBRENA	LORLATINIB TAB 100 MG	21530556000330	Brand
RYDAPT	MIDOSTAURIN CAP 25 MG	21533030000130	Brand
NERLYNX	NERATINIB MALEATE TAB 40 MG (BASE EQUIVALENT)	21533035100320	Brand
REZLIDHIA	OLUTASIDENIB CAP 150 MG	21534960000120	Brand
VONJO	PACRITINIB CITRATE CAP 100 MG	21537550100120	Brand
PEMAZYRE	PEMIGATINIB TAB 4.5 MG	21532260000320	Brand
PEMAZYRE	PEMIGATINIB TAB 9 MG	21532260000330	Brand
PEMAZYRE	PEMIGATINIB TAB 13.5 MG	21532260000340	Brand
TURALIO	PEXIDARTINIB HCL CAP 125 MG (BASE EQUIVALENT)	21533045010110	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 50 MG	21532165000320	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 100 MG	21532165000330	Brand
ICLUSIG	PONATINIB HCL TAB 10 MG (BASE EQUIV)	21531875100315	Brand
ICLUSIG	PONATINIB HCL TAB 15 MG (BASE EQUIV)	21531875100320	Brand
ICLUSIG	PONATINIB HCL TAB 30 MG (BASE EQUIV)	21531875100330	Brand
ICLUSIG	PONATINIB HCL TAB 45 MG (BASE EQUIV)	21531875100340	Brand
GAVRETO	PRALSETINIB CAP 100 MG	21535750000120	Brand
ORGOVYX	RELUGOLIX TAB 120 MG	21405570000320	Brand
QINLOCK	RIPRETINIB TAB 50 MG	21533053000320	Brand
XPOVIO 80 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (80 MG TWICE WEEKLY)	2156006000B720	Brand
XPOVIO 60 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (60 MG TWICE WEEKLY)	2156006000B755	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG ONCE WEEKLY)	2156006000B760	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG TWICE WEEKLY)	2156006000B765	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (80 MG ONCE WEEKLY)	2156006000B770	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 50 MG (100 MG ONCE WEEKLY)	2156006000B775	Brand

XPOVIO	SELINEXOR TAB THERAPY PACK 60 MG (60 MG ONCE WEEKLY)	2156006000B780	Brand
LUMAKRAS	SOTORASIB TAB 120 MG	21532480000320	Brand
LUMAKRAS	SOTORASIB TAB 320 MG	21532480000340	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.1 MG (BASE EQUIVALENT)	21535580400105	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.25 MG (BASE EQUIVALENT)	21535580400110	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.35 MG (BASE EQUIVALENT)	21535580400112	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.5 MG (BASE EQUIVALENT)	21535580400114	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.75 MG (BASE EQUIVALENT)	21535580400118	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 1 MG (BASE EQUIVALENT)	21535580400120	Brand
ТЕРМЕТКО	TEPOTINIB HCL TAB 225 MG	21533773100320	Brand
FOTIVDA	TIVOZANIB HCL CAP 0.89 MG (BASE EQUIVALENT)	21533076250120	Brand
FOTIVDA	TIVOZANIB HCL CAP 1.34 MG (BASE EQUIVALENT)	21533076250130	Brand
CAPRELSA	VANDETANIB TAB 100 MG	21533085000320	Brand
CAPRELSA	VANDETANIB TAB 300 MG	21533085000340	Brand
BRUKINSA	ZANUBRUTINIB CAP 80 MG	21532195000120	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 50-500 MG	21409902120320	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 100-500 MG	21409902120330	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 17.7 MG	21533047100320	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 26.5 MG	21533047100325	Brand
CALQUENCE	ACALABRUTINIB CAP 100 MG	21532103000120	Brand
	-)	

- 1 One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

AND

- 2 Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.

Product Name: Akeega, Alunbrig, Ayvakit, Balversa, Braftovi, Brukinsa, Calquence, Caprelsa, Cometriq, Copiktra, Daurismo, Erleada, Fotivda, Gavreto, Iclusig, Idhifa, Inqovi, Jaypirca, Krazati, Lenvima, Lorbrena, Lumakras, Lytgobi, Mektovi, Nerlynx, Orgovyx, Orserdu, Pemazyre, Qinlock, Rezlidhia, Rydapt, Scemblix, Talzenna, Tepmetko, Tibsovo, Turalio, Vanflyta, Vitrakvi, Vizimpro, Vonjo, Welireg, Xospata, Xpovio, Yonsa

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - ALL Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
YONSA	ABIRATERONE ACETATE MICRONIZED TAB 125 MG	21406010250310	Brand
CALQUENCE	ACALABRUTINIB MALEATE TAB 100 MG	21532103500320	Brand
KRAZATI	ADAGRASIB TAB 200 MG	21532410000320	Brand
ERLEADA	APALUTAMIDE TAB 60 MG	21402410000320	Brand
ERLEADA	APALUTAMIDE TAB 240 MG	21402410000360	Brand
SCEMBLIX	ASCIMINIB HCL TAB 20 MG	21531806100320	Brand
SCEMBLIX	ASCIMINIB HCL TAB 40 MG	21531806100340	Brand
AYVAKIT	AVAPRITINIB TAB 25 MG	21490009000310	Brand
AYVAKIT	AVAPRITINIB TAB 50 MG	21490009000315	Brand
AYVAKIT	AVAPRITINIB TAB 100 MG	21490009000320	Brand
AYVAKIT	AVAPRITINIB TAB 200 MG	21490009000330	Brand

AYVAKIT	AVAPRITINIB TAB 300 MG	21490009000340	Brand
WELIREG	BELZUTIFAN TAB 40 MG	21421020000320	Brand
MEKTOVI	BINIMETINIB TAB 15 MG	21533520000320	Brand
ALUNBRIG	BRIGATINIB TAB INITIATION THERAPY PACK 90 MG & 180 MG	2153051000B720	Brand
ALUNBRIG	BRIGATINIB TAB 30 MG	21530510000330	Brand
ALUNBRIG	BRIGATINIB TAB 90 MG	21530510000350	Brand
ALUNBRIG	BRIGATINIB TAB 180 MG	21530510000365	Brand
COMETRIQ	CABOZANTINIB S-MALATE CAP 3 X 20 MG (60 MG DOSE) KIT	21533010106460	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 1 X 20 MG (100 DOSE) KIT	21533010106470	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 3 X 20 MG (140 DOSE) KIT	21533010106480	Brand
VIZIMPRO	DACOMITINIB TAB 15 MG	21360019000320	Brand
VIZIMPRO	DACOMITINIB TAB 30 MG	21360019000330	Brand
VIZIMPRO	DACOMITINIB TAB 45 MG	21360019000340	Brand
INQOVI	DECITABINE-CEDAZURIDINE TAB 35-100 MG	21990002250320	Brand
COPIKTRA	DUVELISIB CAP 15 MG	21538030000120	Brand
COPIKTRA	DUVELISIB CAP 25 MG	21538030000130	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 86 MG	21403720100320	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 345 MG	21403720100340	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 50 MG (BASE EQUIVALENT)	21535030200320	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 100 MG (BASE EQUIVALENT)	21535030200340	Brand
BRAFTOVI	ENCORAFENIB CAP 75 MG	21532040000130	Brand
BALVERSA	ERDAFITINIB TAB 3 MG	21532225000320	Brand
BALVERSA	ERDAFITINIB TAB 4 MG	21532225000325	Brand
BALVERSA	ERDAFITINIB TAB 5 MG	21532225000330	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (12 MG DAILY DOSE)	2153222800B720	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (16 MG DAILY DOSE)	2153222800B725	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (20 MG DAILY DOSE)	2153222800B730	Brand
XOSPATA	GILTERITINIB FUMARATE TABLET 40 MG (BASE EQUIVALENT)	21533020200320	Brand

DAURISMO	GLASDEGIB MALEATE TAB 25 MG (BASE EQUIVALENT)	21370030300320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 100 MG (BASE EQUIVALENT)	21370030300335	Brand
TIBSOVO	IVOSIDENIB TAB 250 MG	21534940000320	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 25 MG (BASE EQUIVALENT)	21533835200120	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 100 MG (BASE EQUIVALENT)	21533835200150	Brand
VITRAKVI	LAROTRECTINIB SULFATE ORAL SOLN 20 MG/ML (BASE EQUIVALENT)	21533835202020	Brand
LENVIMA 4 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 4 MG (4 MG DAILY DOSE)	2133505420B210	Brand
LENVIMA 8 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 4 MG (8 MG DAILY DOSE)	2133505420B215	Brand
LENVIMA 10 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 MG (10 MG DAILY DOSE)	2133505420B220	Brand
LENVIMA 12MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 3 X 4 MG (12 MG DAILY DOSE)	2133505420B223	Brand
LENVIMA 20 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 10 MG (20 MG DAILY DOSE)	2133505420B230	Brand
LENVIMA 14 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 & 4 MG (14 MG DAILY DOSE)	2133505420B240	Brand
LENVIMA 18 MG DAILY DOSE	LENVATINIB CAP THER PACK 10 MG & 2 X 4 MG (18 MG DAILY DOSE)	2133505420B244	Brand
LENVIMA 24 MG DAILY DOSE	LENVATINIB CAP THER PACK 2 X 10 MG & 4 MG (24 MG DAILY DOSE)	2133505420B250	Brand
LORBRENA	LORLATINIB TAB 25 MG	21530556000320	Brand
LORBRENA	LORLATINIB TAB 100 MG	21530556000330	Brand
RYDAPT	MIDOSTAURIN CAP 25 MG	21533030000130	Brand
NERLYNX	NERATINIB MALEATE TAB 40 MG (BASE EQUIVALENT)	21533035100320	Brand
REZLIDHIA	OLUTASIDENIB CAP 150 MG	21534960000120	Brand
VONJO	PACRITINIB CITRATE CAP 100 MG	21537550100120	Brand
PEMAZYRE	PEMIGATINIB TAB 4.5 MG	21532260000320	Brand
PEMAZYRE	PEMIGATINIB TAB 9 MG	21532260000330	Brand

PEMAZYRE	PEMIGATINIB TAB 13.5 MG	21532260000340	Brand
TURALIO	PEXIDARTINIB HCL CAP 125 MG (BASE EQUIVALENT)	21533045010110	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 50 MG	21532165000320	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 100 MG	21532165000330	Brand
ICLUSIG	PONATINIB HCL TAB 10 MG (BASE EQUIV)	21531875100315	Brand
ICLUSIG	PONATINIB HCL TAB 15 MG (BASE EQUIV)	21531875100320	Brand
ICLUSIG	PONATINIB HCL TAB 30 MG (BASE EQUIV)	21531875100330	Brand
ICLUSIG	PONATINIB HCL TAB 45 MG (BASE EQUIV)	21531875100340	Brand
GAVRETO	PRALSETINIB CAP 100 MG	21535750000120	Brand
ORGOVYX	RELUGOLIX TAB 120 MG	21405570000320	Brand
QINLOCK	RIPRETINIB TAB 50 MG	21533053000320	Brand
XPOVIO 80 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (80 MG TWICE WEEKLY)	2156006000B720	Brand
XPOVIO 60 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (60 MG TWICE WEEKLY)	2156006000B755	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG ONCE WEEKLY)	2156006000B760	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG TWICE WEEKLY)	2156006000B765	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (80 MG ONCE WEEKLY)	2156006000B770	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 50 MG (100 MG ONCE WEEKLY)	2156006000B775	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 60 MG (60 MG ONCE WEEKLY)	2156006000B780	Brand
LUMAKRAS	SOTORASIB TAB 120 MG	21532480000320	Brand
LUMAKRAS	SOTORASIB TAB 320 MG	21532480000340	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.1 MG (BASE EQUIVALENT)	21535580400105	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.25 MG (BASE EQUIVALENT)	21535580400110	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.35 MG (BASE EQUIVALENT)	21535580400112	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.5 MG (BASE EQUIVALENT)	21535580400114	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.75 MG (BASE EQUIVALENT)	21535580400118	Brand

TALZENNA	TALAZOPARIB TOSYLATE CAP 1 MG (BASE EQUIVALENT)	21535580400120	Brand
ТЕРМЕТКО	TEPOTINIB HCL TAB 225 MG	21533773100320	Brand
FOTIVDA	TIVOZANIB HCL CAP 0.89 MG (BASE EQUIVALENT)	21533076250120	Brand
FOTIVDA	TIVOZANIB HCL CAP 1.34 MG (BASE EQUIVALENT)	21533076250130	Brand
CAPRELSA	VANDETANIB TAB 100 MG	21533085000320	Brand
CAPRELSA	VANDETANIB TAB 300 MG	21533085000340	Brand
BRUKINSA	ZANUBRUTINIB CAP 80 MG	21532195000120	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 50-500 MG	21409902120320	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 100-500 MG	21409902120330	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 17.7 MG	21533047100320	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 26.5 MG	21533047100325	Brand
CALQUENCE	ACALABRUTINIB CAP 100 MG	21532103000120	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes *Includes any relevant genetic testing, mutations, etc.	
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Product Name: Akeega, Alunbrig, Ayvakit, Balversa, Braftovi, Brukinsa, Calquence, Caprelsa, Cometriq, Copiktra, Daurismo, Erleada, Fotivda, Gavreto, Iclusig, Idhifa, Inqovi, Jaypirca, Krazati, Lenvima, Lorbrena, Lumakras, Lytgobi, Mektovi, Nerlynx, Orgovyx, Orserdu, Pemazyre, Qinlock, Rezlidhia, Rydapt, Scemblix, Talzenna, Tepmetko, Tibsovo, Turalio, Vanflyta, Vitrakvi, Vizimpro, Vonjo, Welireg, Xospata, Xpovio, Yonsa

Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - MN Plans

Product Name	Generic Name	GPI	Brand/Generic
YONSA	ABIRATERONE ACETATE MICRONIZED TAB 125 MG	21406010250310	Brand
CALQUENCE	ACALABRUTINIB MALEATE TAB 100 MG	21532103500320	Brand
KRAZATI	ADAGRASIB TAB 200 MG	21532410000320	Brand
ERLEADA	APALUTAMIDE TAB 60 MG	21402410000320	Brand
ERLEADA	APALUTAMIDE TAB 240 MG	21402410000360	Brand
SCEMBLIX	ASCIMINIB HCL TAB 20 MG	21531806100320	Brand
SCEMBLIX	ASCIMINIB HCL TAB 40 MG	21531806100340	Brand
AYVAKIT	AVAPRITINIB TAB 25 MG	21490009000310	Brand
AYVAKIT	AVAPRITINIB TAB 50 MG	21490009000315	Brand
AYVAKIT	AVAPRITINIB TAB 100 MG	21490009000320	Brand
AYVAKIT	AVAPRITINIB TAB 200 MG	21490009000330	Brand
AYVAKIT	AVAPRITINIB TAB 300 MG	21490009000340	Brand
WELIREG	BELZUTIFAN TAB 40 MG	21421020000320	Brand
MEKTOVI	BINIMETINIB TAB 15 MG	21533520000320	Brand
ALUNBRIG	BRIGATINIB TAB INITIATION THERAPY PACK 90 MG & 180 MG	2153051000B720	Brand
ALUNBRIG	BRIGATINIB TAB 30 MG	21530510000330	Brand
ALUNBRIG	BRIGATINIB TAB 90 MG	21530510000350	Brand
ALUNBRIG	BRIGATINIB TAB 180 MG	21530510000365	Brand
COMETRIQ	CABOZANTINIB S-MALATE CAP 3 X 20 MG (60 MG DOSE) KIT	21533010106460	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 1 X 20 MG (100 DOSE) KIT	21533010106470	Brand

CONTENTS	CARCZANTINIR C MAL CAR 4 V CO MO C C V CC MO		Donal
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 3 X 20 MG (140 DOSE) KIT	21533010106480	Brand
VIZIMPRO	DACOMITINIB TAB 15 MG	21360019000320	Brand
VIZIMPRO	DACOMITINIB TAB 30 MG	21360019000330	Brand
VIZIMPRO	DACOMITINIB TAB 45 MG	21360019000340	Brand
INQOVI	DECITABINE-CEDAZURIDINE TAB 35-100 MG	21990002250320	Brand
COPIKTRA	DUVELISIB CAP 15 MG	21538030000120	Brand
COPIKTRA	DUVELISIB CAP 25 MG	21538030000130	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 86 MG	21403720100320	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 345 MG	21403720100340	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 50 MG (BASE EQUIVALENT)	21535030200320	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 100 MG (BASE EQUIVALENT)	21535030200340	Brand
BRAFTOVI	ENCORAFENIB CAP 75 MG	21532040000130	Brand
BALVERSA	ERDAFITINIB TAB 3 MG	21532225000320	Brand
BALVERSA	ERDAFITINIB TAB 4 MG	21532225000325	Brand
BALVERSA	ERDAFITINIB TAB 5 MG	21532225000330	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (12 MG DAILY DOSE)	2153222800B720	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (16 MG DAILY DOSE)	2153222800B725	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (20 MG DAILY DOSE)	2153222800B730	Brand
XOSPATA	GILTERITINIB FUMARATE TABLET 40 MG (BASE EQUIVALENT)	21533020200320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 25 MG (BASE EQUIVALENT)	21370030300320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 100 MG (BASE EQUIVALENT)	21370030300335	Brand
TIBSOVO	IVOSIDENIB TAB 250 MG	21534940000320	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 25 MG (BASE EQUIVALENT)	21533835200120	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 100 MG (BASE EQUIVALENT)	21533835200150	Brand
VITRAKVI	LAROTRECTINIB SULFATE ORAL SOLN 20 MG/ML (BASE EQUIVALENT)	21533835202020	Brand
LENVIMA 4 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 4 MG (4 MG DAILY DOSE)	2133505420B210	Brand

LENVIMA 8 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 4 MG (8 MG DAILY DOSE)	2133505420B215	Brand
LENVIMA 10 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 MG (10 MG DAILY DOSE)	2133505420B220	Brand
LENVIMA 12MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 3 X 4 MG (12 MG DAILY DOSE)	2133505420B223	Brand
LENVIMA 20 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 10 MG (20 MG DAILY DOSE)	2133505420B230	Brand
LENVIMA 14 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 & 4 MG (14 MG DAILY DOSE)	2133505420B240	Brand
LENVIMA 18 MG DAILY DOSE	LENVATINIB CAP THER PACK 10 MG & 2 X 4 MG (18 MG DAILY DOSE)	2133505420B244	Brand
LENVIMA 24 MG DAILY DOSE	LENVATINIB CAP THER PACK 2 X 10 MG & 4 MG (24 MG DAILY DOSE)	2133505420B250	Brand
LORBRENA	LORLATINIB TAB 25 MG	21530556000320	Brand
LORBRENA	LORLATINIB TAB 100 MG	21530556000330	Brand
RYDAPT	MIDOSTAURIN CAP 25 MG	21533030000130	Brand
NERLYNX	NERATINIB MALEATE TAB 40 MG (BASE EQUIVALENT)	21533035100320	Brand
REZLIDHIA	OLUTASIDENIB CAP 150 MG	21534960000120	Brand
VONJO	PACRITINIB CITRATE CAP 100 MG	21537550100120	Brand
PEMAZYRE	PEMIGATINIB TAB 4.5 MG	21532260000320	Brand
PEMAZYRE	PEMIGATINIB TAB 9 MG	21532260000330	Brand
PEMAZYRE	PEMIGATINIB TAB 13.5 MG	21532260000340	Brand
TURALIO	PEXIDARTINIB HCL CAP 125 MG (BASE EQUIVALENT)	21533045010110	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 50 MG	21532165000320	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 100 MG	21532165000330	Brand
ICLUSIG	PONATINIB HCL TAB 10 MG (BASE EQUIV)	21531875100315	Brand
ICLUSIG	PONATINIB HCL TAB 15 MG (BASE EQUIV)	21531875100320	Brand
ICLUSIG	PONATINIB HCL TAB 30 MG (BASE EQUIV)	21531875100330	Brand
ICLUSIG	PONATINIB HCL TAB 45 MG (BASE EQUIV)	21531875100340	Brand
GAVRETO	PRALSETINIB CAP 100 MG	21535750000120	Brand
ORGOVYX	RELUGOLIX TAB 120 MG	21405570000320	Brand

QINLOCK	RIPRETINIB TAB 50 MG	21533053000320	Brand
XPOVIO 80 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (80 MG TWICE WEEKLY)	2156006000B720	Brand
XPOVIO 60 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (60 MG TWICE WEEKLY)	2156006000B755	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG ONCE WEEKLY)	2156006000B760	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG TWICE WEEKLY)	2156006000B765	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (80 MG ONCE WEEKLY)	2156006000B770	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 50 MG (100 MG ONCE WEEKLY)	2156006000B775	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 60 MG (60 MG ONCE WEEKLY)	2156006000B780	Brand
LUMAKRAS	SOTORASIB TAB 120 MG	21532480000320	Brand
LUMAKRAS	SOTORASIB TAB 320 MG	21532480000340	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.1 MG (BASE EQUIVALENT)	21535580400105	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.25 MG (BASE EQUIVALENT)	21535580400110	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.35 MG (BASE EQUIVALENT)	21535580400112	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.5 MG (BASE EQUIVALENT)	21535580400114	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.75 MG (BASE EQUIVALENT)	21535580400118	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 1 MG (BASE EQUIVALENT)	21535580400120	Brand
TEPMETKO	TEPOTINIB HCL TAB 225 MG	21533773100320	Brand
FOTIVDA	TIVOZANIB HCL CAP 0.89 MG (BASE EQUIVALENT)	21533076250120	Brand
FOTIVDA	TIVOZANIB HCL CAP 1.34 MG (BASE EQUIVALENT)	21533076250130	Brand
CAPRELSA	VANDETANIB TAB 100 MG	21533085000320	Brand
CAPRELSA	VANDETANIB TAB 300 MG	21533085000340	Brand
BRUKINSA	ZANUBRUTINIB CAP 80 MG	21532195000120	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 50-500 MG	21409902120320	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 100-500 MG	21409902120330	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 17.7 MG	21533047100320	Brand

VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 26.5 MG	21533047100325	Brand
CALQUENCE	ACALABRUTINIB CAP 100 MG	21532103000120	Brand

- 1 One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member in one of the following:*
 - United States Pharmacopeia Drug Information
 - American Hospital Formulary Service Drug Information
 - One article in a major peer-reviewed medical journal that recognizes the safety and efficacy of the requested drug, in the member's specific condition

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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Product Name: Akeega, Alunbrig, Ayvakit, Balversa, Braftovi, Brukinsa, Calquence, Caprelsa, Cometriq, Copiktra, Daurismo, Erleada, Fotivda, Gavreto, Iclusig, Idhifa, Inqovi, Jaypirca,

Krazati, Lenvima, Lorbrena, Lumakras, Lytgobi, Mektovi, Nerlynx, Orgovyx, Orserdu, Pemazyre, Qinlock, Rezlidhia, Rydapt, Scemblix, Talzenna, Tepmetko, Tibsovo, Turalio, Vanflyta, Vitrakvi, Vizimpro, Vonjo, Welireg, Xospata, Xpovio, Yonsa

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - MN Plans

Product Name	Generic Name	GPI	Brand/Generic
YONSA	ABIRATERONE ACETATE MICRONIZED TAB 125 MG	21406010250310	Brand
CALQUENCE	ACALABRUTINIB MALEATE TAB 100 MG	21532103500320	Brand
KRAZATI	ADAGRASIB TAB 200 MG	21532410000320	Brand
ERLEADA	APALUTAMIDE TAB 60 MG	21402410000320	Brand
ERLEADA	APALUTAMIDE TAB 240 MG	21402410000360	Brand
SCEMBLIX	ASCIMINIB HCL TAB 20 MG	21531806100320	Brand
SCEMBLIX	ASCIMINIB HCL TAB 40 MG	21531806100340	Brand
AYVAKIT	AVAPRITINIB TAB 25 MG	21490009000310	Brand
AYVAKIT	AVAPRITINIB TAB 50 MG	21490009000315	Brand
AYVAKIT	AVAPRITINIB TAB 100 MG	21490009000320	Brand
AYVAKIT	AVAPRITINIB TAB 200 MG	21490009000330	Brand
AYVAKIT	AVAPRITINIB TAB 300 MG	21490009000340	Brand
WELIREG	BELZUTIFAN TAB 40 MG	21421020000320	Brand
MEKTOVI	BINIMETINIB TAB 15 MG	21533520000320	Brand
ALUNBRIG	BRIGATINIB TAB INITIATION THERAPY PACK 90 MG & 180 MG	2153051000B720	Brand
ALUNBRIG	BRIGATINIB TAB 30 MG	21530510000330	Brand
ALUNBRIG	BRIGATINIB TAB 90 MG	21530510000350	Brand
ALUNBRIG	BRIGATINIB TAB 180 MG	21530510000365	Brand
COMETRIQ	CABOZANTINIB S-MALATE CAP 3 X 20 MG (60 MG DOSE) KIT	21533010106460	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 1 X 20 MG (100 DOSE) KIT	21533010106470	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 3 X 20 MG (140 DOSE) KIT	21533010106480	Brand
VIZIMPRO	DACOMITINIB TAB 15 MG	21360019000320	Brand
VIZIMPRO	DACOMITINIB TAB 30 MG	21360019000330	Brand

VIZIMPRO	DACOMITINIB TAB 45 MG	21360019000340	Brand
INQOVI	DECITABINE-CEDAZURIDINE TAB 35-100 MG	21990002250320	Brand
COPIKTRA	DUVELISIB CAP 15 MG		Brand
		21538030000120	
COPIKTRA	DUVELISIB CAP 25 MG	21538030000130	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 86 MG	21403720100320	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 345 MG	21403720100340	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 50 MG (BASE EQUIVALENT)	21535030200320	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 100 MG (BASE EQUIVALENT)	21535030200340	Brand
BRAFTOVI	ENCORAFENIB CAP 75 MG	21532040000130	Brand
BALVERSA	ERDAFITINIB TAB 3 MG	21532225000320	Brand
BALVERSA	ERDAFITINIB TAB 4 MG	21532225000325	Brand
BALVERSA	ERDAFITINIB TAB 5 MG	21532225000330	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (12 MG DAILY DOSE)	2153222800B720	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (16 MG DAILY DOSE)	2153222800B725	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (20 MG DAILY DOSE)	2153222800B730	Brand
XOSPATA	GILTERITINIB FUMARATE TABLET 40 MG (BASE EQUIVALENT)	21533020200320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 25 MG (BASE EQUIVALENT)	21370030300320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 100 MG (BASE EQUIVALENT)	21370030300335	Brand
TIBSOVO	IVOSIDENIB TAB 250 MG	21534940000320	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 25 MG (BASE EQUIVALENT)	21533835200120	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 100 MG (BASE EQUIVALENT)	21533835200150	Brand
VITRAKVI	LAROTRECTINIB SULFATE ORAL SOLN 20 MG/ML (BASE EQUIVALENT)	21533835202020	Brand
LENVIMA 4 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 4 MG (4 MG DAILY DOSE)	2133505420B210	Brand
LENVIMA 8 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 4 MG (8 MG DAILY DOSE)	2133505420B215	Brand
LENVIMA 10 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 MG (10 MG DAILY DOSE)	2133505420B220	Brand

LENVIMA 12MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 3 X 4 MG (12 MG DAILY DOSE)	2133505420B223	Brand
LENVIMA 20 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 10 MG (20 MG DAILY DOSE)	2133505420B230	Brand
LENVIMA 14 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 & 4 MG (14 MG DAILY DOSE)	2133505420B240	Brand
LENVIMA 18 MG DAILY DOSE	LENVATINIB CAP THER PACK 10 MG & 2 X 4 MG (18 MG DAILY DOSE)	2133505420B244	Brand
LENVIMA 24 MG DAILY DOSE	LENVATINIB CAP THER PACK 2 X 10 MG & 4 MG (24 MG DAILY DOSE)	2133505420B250	Brand
LORBRENA	LORLATINIB TAB 25 MG	21530556000320	Brand
LORBRENA	LORLATINIB TAB 100 MG	21530556000330	Brand
RYDAPT	MIDOSTAURIN CAP 25 MG	21533030000130	Brand
NERLYNX	NERATINIB MALEATE TAB 40 MG (BASE EQUIVALENT)	21533035100320	Brand
REZLIDHIA	OLUTASIDENIB CAP 150 MG	21534960000120	Brand
VONJO	PACRITINIB CITRATE CAP 100 MG	21537550100120	Brand
PEMAZYRE	PEMIGATINIB TAB 4.5 MG	21532260000320	Brand
PEMAZYRE	PEMIGATINIB TAB 9 MG	21532260000330	Brand
PEMAZYRE	PEMIGATINIB TAB 13.5 MG	21532260000340	Brand
TURALIO	PEXIDARTINIB HCL CAP 125 MG (BASE EQUIVALENT)	21533045010110	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 50 MG	21532165000320	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 100 MG	21532165000330	Brand
ICLUSIG	PONATINIB HCL TAB 10 MG (BASE EQUIV)	21531875100315	Brand
ICLUSIG	PONATINIB HCL TAB 15 MG (BASE EQUIV)	21531875100320	Brand
ICLUSIG	PONATINIB HCL TAB 30 MG (BASE EQUIV)	21531875100330	Brand
ICLUSIG	PONATINIB HCL TAB 45 MG (BASE EQUIV)	21531875100340	Brand
GAVRETO	PRALSETINIB CAP 100 MG	21535750000120	Brand
ORGOVYX	RELUGOLIX TAB 120 MG	21405570000320	Brand
QINLOCK	RIPRETINIB TAB 50 MG	21533053000320	Brand
XPOVIO 80 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (80 MG TWICE WEEKLY)	2156006000B720	Brand

XPOVIO 60 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (60 MG TWICE WEEKLY)	2156006000B755	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG ONCE WEEKLY)	2156006000B760	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG TWICE WEEKLY)	2156006000B765	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (80 MG ONCE WEEKLY)	2156006000B770	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 50 MG (100 MG ONCE WEEKLY)	2156006000B775	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 60 MG (60 MG ONCE WEEKLY)	2156006000B780	Brand
LUMAKRAS	SOTORASIB TAB 120 MG	21532480000320	Brand
LUMAKRAS	SOTORASIB TAB 320 MG	21532480000340	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.1 MG (BASE EQUIVALENT)	21535580400105	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.25 MG (BASE EQUIVALENT)	21535580400110	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.35 MG (BASE EQUIVALENT)	21535580400112	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.5 MG (BASE EQUIVALENT)	21535580400114	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.75 MG (BASE EQUIVALENT)	21535580400118	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 1 MG (BASE EQUIVALENT)	21535580400120	Brand
TEPMETKO	TEPOTINIB HCL TAB 225 MG	21533773100320	Brand
FOTIVDA	TIVOZANIB HCL CAP 0.89 MG (BASE EQUIVALENT)	21533076250120	Brand
FOTIVDA	TIVOZANIB HCL CAP 1.34 MG (BASE EQUIVALENT)	21533076250130	Brand
CAPRELSA	VANDETANIB TAB 100 MG	21533085000320	Brand
CAPRELSA	VANDETANIB TAB 300 MG	21533085000340	Brand
BRUKINSA	ZANUBRUTINIB CAP 80 MG	21532195000120	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 50-500 MG	21409902120320	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 100-500 MG	21409902120330	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 17.7 MG	21533047100320	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 26.5 MG	21533047100325	Brand
CALQUENCE	ACALABRUTINIB CAP 100 MG	21532103000120	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member in one of the following*:
 - United States Pharmacopeia Drug Information
 - American Hospital Formulary Service Drug Information
 - One article in a major peer-reviewed medical journal that recognizes the safety and efficacy of the requested drug, in the member's specific condition

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes *Includes any relevant genetic testing, mutations, etc.

Product Name: Akeega, Alunbrig, Ayvakit, Balversa, Braftovi, Brukinsa, Calquence, Caprelsa, Cometriq, Copiktra, Daurismo, Erleada, Fotivda, Gavreto, Iclusig, Idhifa, Inqovi, Jaypirca, Krazati, Lenvima, Lorbrena, Lumakras, Lytgobi, Mektovi, Nerlynx, Orgovyx, Orserdu, Pemazyre, Qinlock, Rezlidhia, Rydapt, Scemblix, Talzenna, Tepmetko, Tibsovo, Turalio, Vanflyta, Vitrakvi, Vizimpro, Vonjo, Welireg, Xospata, Xpovio, Yonsa

Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL Plans

Product Name	Generic Name	GPI	Brand/Generic
YONSA	ABIRATERONE ACETATE MICRONIZED TAB 125 MG	21406010250310	Brand
CALQUENCE	ACALABRUTINIB MALEATE TAB 100 MG	21532103500320	Brand
KRAZATI	ADAGRASIB TAB 200 MG	21532410000320	Brand
ERLEADA	APALUTAMIDE TAB 60 MG	21402410000320	Brand
ERLEADA	APALUTAMIDE TAB 240 MG	21402410000360	Brand
SCEMBLIX	ASCIMINIB HCL TAB 20 MG	21531806100320	Brand
SCEMBLIX	ASCIMINIB HCL TAB 40 MG	21531806100340	Brand
AYVAKIT	AVAPRITINIB TAB 25 MG	21490009000310	Brand
AYVAKIT	AVAPRITINIB TAB 50 MG	21490009000315	Brand
AYVAKIT	AVAPRITINIB TAB 100 MG	21490009000320	Brand
AYVAKIT	AVAPRITINIB TAB 200 MG	21490009000330	Brand
AYVAKIT	AVAPRITINIB TAB 300 MG	21490009000340	Brand
WELIREG	BELZUTIFAN TAB 40 MG	21421020000320	Brand
MEKTOVI	BINIMETINIB TAB 15 MG	21533520000320	Brand
ALUNBRIG	BRIGATINIB TAB INITIATION THERAPY PACK 90 MG & 180 MG	2153051000B720	Brand
ALUNBRIG	BRIGATINIB TAB 30 MG	21530510000330	Brand
ALUNBRIG	BRIGATINIB TAB 90 MG	21530510000350	Brand
ALUNBRIG	BRIGATINIB TAB 180 MG	21530510000365	Brand
COMETRIQ	CABOZANTINIB S-MALATE CAP 3 X 20 MG (60 MG DOSE) KIT	21533010106460	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 1 X 20 MG (100 DOSE) KIT	21533010106470	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 3 X 20 MG (140 DOSE) KIT	21533010106480	Brand
VIZIMPRO	DACOMITINIB TAB 15 MG	21360019000320	Brand
VIZIMPRO	DACOMITINIB TAB 30 MG	21360019000330	Brand
VIZIMPRO	DACOMITINIB TAB 45 MG	21360019000340	Brand
INQOVI	DECITABINE-CEDAZURIDINE TAB 35-100 MG	21990002250320	Brand

COPIKTRA	DUVELISIB CAP 15 MG	21538030000120	Brand
COPIKTRA	DUVELISIB CAP 25 MG	21538030000130	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 86 MG	21403720100320	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 345 MG	21403720100340	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 50 MG (BASE EQUIVALENT)	21535030200320	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 100 MG (BASE EQUIVALENT)	21535030200340	Brand
BRAFTOVI	ENCORAFENIB CAP 75 MG	21532040000130	Brand
BALVERSA	ERDAFITINIB TAB 3 MG	21532225000320	Brand
BALVERSA	ERDAFITINIB TAB 4 MG	21532225000325	Brand
BALVERSA	ERDAFITINIB TAB 5 MG	21532225000330	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (12 MG DAILY DOSE)	2153222800B720	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (16 MG DAILY DOSE)	2153222800B725	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (20 MG DAILY DOSE)	2153222800B730	Brand
XOSPATA	GILTERITINIB FUMARATE TABLET 40 MG (BASE EQUIVALENT)	21533020200320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 25 MG (BASE EQUIVALENT)	21370030300320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 100 MG (BASE EQUIVALENT)	21370030300335	Brand
TIBSOVO	IVOSIDENIB TAB 250 MG	21534940000320	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 25 MG (BASE EQUIVALENT)	21533835200120	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 100 MG (BASE EQUIVALENT)	21533835200150	Brand
VITRAKVI	LAROTRECTINIB SULFATE ORAL SOLN 20 MG/ML (BASE EQUIVALENT)	21533835202020	Brand
LENVIMA 4 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 4 MG (4 MG DAILY DOSE)	2133505420B210	Brand
LENVIMA 8 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 4 MG (8 MG DAILY DOSE)	2133505420B215	Brand
LENVIMA 10 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 MG (10 MG DAILY DOSE)	2133505420B220	Brand
LENVIMA 12MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 3 X 4 MG (12 MG DAILY DOSE)	2133505420B223	Brand

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LENVIMA 20 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 10 MG (20 MG DAILY DOSE)	2133505420B230	Brand
LENVIMA 14 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 & 4 MG (14 MG DAILY DOSE)	2133505420B240	Brand
LENVIMA 18 MG DAILY DOSE	LENVATINIB CAP THER PACK 10 MG & 2 X 4 MG (18 MG DAILY DOSE)	2133505420B244	Brand
LENVIMA 24 MG DAILY DOSE	LENVATINIB CAP THER PACK 2 X 10 MG & 4 MG (24 MG DAILY DOSE)	2133505420B250	Brand
LORBRENA	LORLATINIB TAB 25 MG	21530556000320	Brand
LORBRENA	LORLATINIB TAB 100 MG	21530556000330	Brand
RYDAPT	MIDOSTAURIN CAP 25 MG	21533030000130	Brand
NERLYNX	NERATINIB MALEATE TAB 40 MG (BASE EQUIVALENT)	21533035100320	Brand
REZLIDHIA	OLUTASIDENIB CAP 150 MG	21534960000120	Brand
VONJO	PACRITINIB CITRATE CAP 100 MG	21537550100120	Brand
PEMAZYRE	PEMIGATINIB TAB 4.5 MG	21532260000320	Brand
PEMAZYRE	PEMIGATINIB TAB 9 MG	21532260000330	Brand
PEMAZYRE	PEMIGATINIB TAB 13.5 MG	21532260000340	Brand
TURALIO	PEXIDARTINIB HCL CAP 125 MG (BASE EQUIVALENT)	21533045010110	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 50 MG	21532165000320	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 100 MG	21532165000330	Brand
ICLUSIG	PONATINIB HCL TAB 10 MG (BASE EQUIV)	21531875100315	Brand
ICLUSIG	PONATINIB HCL TAB 15 MG (BASE EQUIV)	21531875100320	Brand
ICLUSIG	PONATINIB HCL TAB 30 MG (BASE EQUIV)	21531875100330	Brand
ICLUSIG	PONATINIB HCL TAB 45 MG (BASE EQUIV)	21531875100340	Brand
GAVRETO	PRALSETINIB CAP 100 MG	21535750000120	Brand
ORGOVYX	RELUGOLIX TAB 120 MG	21405570000320	Brand
QINLOCK	RIPRETINIB TAB 50 MG	21533053000320	Brand
XPOVIO 80 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (80 MG TWICE WEEKLY)	2156006000B720	Brand
XPOVIO 60 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (60 MG TWICE WEEKLY)	2156006000B755	Brand

XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG ONCE WEEKLY) 2156006000B760		Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG 2156006000) TWICE WEEKLY)		Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (80 MG ONCE WEEKLY)	2156006000B770	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 50 MG (100 MG ONCE WEEKLY)	2156006000B775	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 60 MG (60 MG ONCE WEEKLY)	2156006000B780	Brand
LUMAKRAS	SOTORASIB TAB 120 MG	21532480000320	Brand
LUMAKRAS	SOTORASIB TAB 320 MG	21532480000340	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.1 MG (BASE EQUIVALENT)	21535580400105	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.25 MG (BASE EQUIVALENT)	21535580400110	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.35 MG (BASE EQUIVALENT)	21535580400112	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.5 MG (BASE EQUIVALENT)	21535580400114	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.75 MG (BASE EQUIVALENT)	21535580400118	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 1 MG (BASE EQUIVALENT)	21535580400120	Brand
TEPMETKO	TEPOTINIB HCL TAB 225 MG	21533773100320	Brand
FOTIVDA	TIVOZANIB HCL CAP 0.89 MG (BASE EQUIVALENT)	21533076250120	Brand
FOTIVDA	TIVOZANIB HCL CAP 1.34 MG (BASE EQUIVALENT)	21533076250130	Brand
CAPRELSA	VANDETANIB TAB 100 MG	21533085000320	Brand
CAPRELSA	VANDETANIB TAB 300 MG	21533085000340	Brand
BRUKINSA	ZANUBRUTINIB CAP 80 MG	21532195000120	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 50-500 MG	21409902120320	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 100-500 MG	21409902120330	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 17.7 MG	21533047100320	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 26.5 MG	21533047100325	Brand
CALQUENCE	ACALABRUTINIB CAP 100 MG	21532103000120	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member in one of the following*:
 - American Hospital Formulary Service Drug Information
 - Thompson Micromedex's Drug Dex
 - Elsevier Gold Standard's Clinical Pharmacology
 - Two articles in a major peer-reviewed medical journal from the United States or Great Britain that recognizes the safety and efficacy of the requested drug, in the member's specific condition

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes | *Includes any relevant genetic testing, mutations, etc.

Product Name: Akeega, Alunbrig, Ayvakit, Balversa, Braftovi, Brukinsa, Calquence, Caprelsa, Cometriq, Copiktra, Daurismo, Erleada, Fotivda, Gavreto, Iclusig, Idhifa, Inqovi, Jaypirca, Krazati, Lenvima, Lorbrena, Lumakras, Lytgobi, Mektovi, Nerlynx, Orgovyx, Orserdu, Pemazyre, Qinlock, Rezlidhia, Rydapt, Scemblix, Talzenna, Tepmetko, Tibsovo, Turalio, Vanflyta, Vitrakvi, Vizimpro, Vonjo, Welireg, Xospata, Xpovio, Yonsa

Approval Length 12 month(s)

Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL Plans

Product Name	Generic Name	GPI	Brand/Generic
YONSA	ABIRATERONE ACETATE MICRONIZED TAB 125 MG	21406010250310	Brand
CALQUENCE	ACALABRUTINIB MALEATE TAB 100 MG	21532103500320	Brand
KRAZATI	ADAGRASIB TAB 200 MG	21532410000320	Brand
ERLEADA	APALUTAMIDE TAB 60 MG	21402410000320	Brand
ERLEADA	APALUTAMIDE TAB 240 MG	21402410000360	Brand
SCEMBLIX	ASCIMINIB HCL TAB 20 MG	21531806100320	Brand
SCEMBLIX	ASCIMINIB HCL TAB 40 MG	21531806100340	Brand
AYVAKIT	AVAPRITINIB TAB 25 MG	21490009000310	Brand
AYVAKIT	AVAPRITINIB TAB 50 MG	21490009000315	Brand
AYVAKIT	AVAPRITINIB TAB 100 MG	21490009000320	Brand
AYVAKIT	AVAPRITINIB TAB 200 MG	21490009000330	Brand
AYVAKIT	AVAPRITINIB TAB 300 MG	21490009000340	Brand
WELIREG	BELZUTIFAN TAB 40 MG	21421020000320	Brand
MEKTOVI	BINIMETINIB TAB 15 MG	21533520000320	Brand
ALUNBRIG	BRIGATINIB TAB INITIATION THERAPY PACK 90 MG & 180 MG	2153051000B720	Brand
ALUNBRIG	BRIGATINIB TAB 30 MG	21530510000330	Brand
ALUNBRIG	BRIGATINIB TAB 90 MG	21530510000350	Brand
ALUNBRIG	BRIGATINIB TAB 180 MG	21530510000365	Brand
COMETRIQ	CABOZANTINIB S-MALATE CAP 3 X 20 MG (60 MG DOSE) KIT	21533010106460	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 1 X 20 MG (100 DOSE) KIT	21533010106470	Brand
COMETRIQ	CABOZANTINIB S-MAL CAP 1 X 80 MG & 3 X 20 MG (140 DOSE) KIT	21533010106480	Brand
VIZIMPRO	DACOMITINIB TAB 15 MG	21360019000320	Brand
VIZIMPRO	DACOMITINIB TAB 30 MG	21360019000330	Brand
VIZIMPRO	DACOMITINIB TAB 45 MG	21360019000340	Brand
INQOVI	DECITABINE-CEDAZURIDINE TAB 35-100 MG	21990002250320	Brand
COPIKTRA	DUVELISIB CAP 15 MG	21538030000120	Brand

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COPIKTRA	DUVELISIB CAP 25 MG	21538030000130	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 86 MG	21403720100320	Brand
ORSERDU	ELACESTRANT HYDROCHLORIDE TAB 345 MG	21403720100340	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 50 MG (BASE EQUIVALENT)	21535030200320	Brand
IDHIFA	ENASIDENIB MESYLATE TAB 100 MG (BASE EQUIVALENT)	21535030200340	Brand
BRAFTOVI	ENCORAFENIB CAP 75 MG	21532040000130	Brand
BALVERSA	ERDAFITINIB TAB 3 MG	21532225000320	Brand
BALVERSA	ERDAFITINIB TAB 4 MG	21532225000325	Brand
BALVERSA	ERDAFITINIB TAB 5 MG	21532225000330	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (12 MG DAILY DOSE)	2153222800B720	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (16 MG DAILY DOSE)	2153222800B725	Brand
LYTGOBI	FUTIBATINIB TAB THERAPY PACK 4 MG (20 MG DAILY DOSE)	2153222800B730	Brand
XOSPATA	GILTERITINIB FUMARATE TABLET 40 MG (BASE EQUIVALENT)	21533020200320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 25 MG (BASE EQUIVALENT)	21370030300320	Brand
DAURISMO	GLASDEGIB MALEATE TAB 100 MG (BASE EQUIVALENT)	21370030300335	Brand
TIBSOVO	IVOSIDENIB TAB 250 MG	21534940000320	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 25 MG (BASE EQUIVALENT)	21533835200120	Brand
VITRAKVI	LAROTRECTINIB SULFATE CAP 100 MG (BASE EQUIVALENT)	21533835200150	Brand
VITRAKVI	LAROTRECTINIB SULFATE ORAL SOLN 20 MG/ML (BASE EQUIVALENT)	21533835202020	Brand
LENVIMA 4 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 4 MG (4 MG DAILY DOSE)	2133505420B210	Brand
LENVIMA 8 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 4 MG (8 MG DAILY DOSE)	2133505420B215	Brand
LENVIMA 10 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 MG (10 MG DAILY DOSE)	2133505420B220	Brand
LENVIMA 12MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 3 X 4 MG (12 MG DAILY DOSE)	2133505420B223	Brand

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LENVIMA 20 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 2 X 10 MG (20 MG DAILY DOSE)	2133505420B230	Brand
LENVIMA 14 MG DAILY DOSE	LENVATINIB CAP THERAPY PACK 10 & 4 MG (14 MG DAILY DOSE)	2133505420B240	Brand
LENVIMA 18 MG DAILY DOSE	LENVATINIB CAP THER PACK 10 MG & 2 X 4 MG (18 MG DAILY DOSE)	2133505420B244	Brand
LENVIMA 24 MG DAILY DOSE	LENVATINIB CAP THER PACK 2 X 10 MG & 4 MG (24 MG DAILY DOSE)	2133505420B250	Brand
LORBRENA	LORLATINIB TAB 25 MG	21530556000320	Brand
LORBRENA	LORLATINIB TAB 100 MG	21530556000330	Brand
RYDAPT	MIDOSTAURIN CAP 25 MG	21533030000130	Brand
NERLYNX	NERATINIB MALEATE TAB 40 MG (BASE EQUIVALENT)	21533035100320	Brand
REZLIDHIA	OLUTASIDENIB CAP 150 MG	21534960000120	Brand
VONJO	PACRITINIB CITRATE CAP 100 MG	21537550100120	Brand
PEMAZYRE	PEMIGATINIB TAB 4.5 MG	21532260000320	Brand
PEMAZYRE	PEMIGATINIB TAB 9 MG	21532260000330	Brand
PEMAZYRE	PEMIGATINIB TAB 13.5 MG	21532260000340	Brand
TURALIO	PEXIDARTINIB HCL CAP 125 MG (BASE EQUIVALENT)	21533045010110	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 50 MG	21532165000320	Brand
JAYPIRCA	PIRTOBRUTINIB TAB 100 MG	21532165000330	Brand
ICLUSIG	PONATINIB HCL TAB 10 MG (BASE EQUIV)	21531875100315	Brand
ICLUSIG	PONATINIB HCL TAB 15 MG (BASE EQUIV)	21531875100320	Brand
ICLUSIG	PONATINIB HCL TAB 30 MG (BASE EQUIV)	21531875100330	Brand
ICLUSIG	PONATINIB HCL TAB 45 MG (BASE EQUIV)	21531875100340	Brand
GAVRETO	PRALSETINIB CAP 100 MG	21535750000120	Brand
ORGOVYX	RELUGOLIX TAB 120 MG	21405570000320	Brand
QINLOCK	RIPRETINIB TAB 50 MG	21533053000320	Brand
XPOVIO 80 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (80 MG TWICE WEEKLY)	2156006000B720	Brand
XPOVIO 60 MG TWICE WEEKLY	SELINEXOR TAB THERAPY PACK 20 MG (60 MG TWICE WEEKLY)	2156006000B755	Brand

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XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG ONCE WEEKLY)	2156006000B760	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (40 MG TWICE WEEKLY)		Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 40 MG (80 MG ONCE WEEKLY)	2156006000B770	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 50 MG (100 MG ONCE WEEKLY)	2156006000B775	Brand
XPOVIO	SELINEXOR TAB THERAPY PACK 60 MG (60 MG ONCE WEEKLY)	2156006000B780	Brand
LUMAKRAS	SOTORASIB TAB 120 MG	21532480000320	Brand
LUMAKRAS	SOTORASIB TAB 320 MG	21532480000340	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.1 MG (BASE EQUIVALENT)	21535580400105	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.25 MG (BASE EQUIVALENT)	21535580400110	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.35 MG (BASE EQUIVALENT)	21535580400112	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.5 MG (BASE EQUIVALENT)	21535580400114	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 0.75 MG (BASE EQUIVALENT)	21535580400118	Brand
TALZENNA	TALAZOPARIB TOSYLATE CAP 1 MG (BASE EQUIVALENT)	21535580400120	Brand
TEPMETKO	TEPOTINIB HCL TAB 225 MG	21533773100320	Brand
FOTIVDA	TIVOZANIB HCL CAP 0.89 MG (BASE EQUIVALENT)	21533076250120	Brand
FOTIVDA	TIVOZANIB HCL CAP 1.34 MG (BASE EQUIVALENT)	21533076250130	Brand
CAPRELSA	VANDETANIB TAB 100 MG	21533085000320	Brand
CAPRELSA	VANDETANIB TAB 300 MG	21533085000340	Brand
BRUKINSA	ZANUBRUTINIB CAP 80 MG	21532195000120	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 50-500 MG	21409902120320	Brand
AKEEGA	NIRAPARIB TOSYLATE-ABIRATERONE ACETATE TAB 100-500 MG	21409902120330	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 17.7 MG	21533047100320	Brand
VANFLYTA	QUIZARTINIB DIHYDROCHLORIDE TAB 26.5 MG	21533047100325	Brand
CALQUENCE	ACALABRUTINIB CAP 100 MG	21532103000120	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member in one of the following*:
 - American Hospital Formulary Service Drug Information
 - Thompson Micromedex's Drug Dex
 - Elsevier Gold Standard's Clinical Pharmacology
 - Two articles in a major peer-reviewed medical journal from the United States or Great Britain that recognizes the safety and efficacy of the requested drug, in the member's specific condition

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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2. Revision History

Date	Notes
4/17/2024	Updated guideline

Restricted Oral Oncology Drugs Quartz Specialty Pharmacy Network				
(3) The Proposition of Trading to Section and county of the Section Se				

Prior Authorization Guideline

Guideline ID	GL-145307	
Guideline Name	Restricted Oral Oncology Drugs Quartz Specialty Pharmacy Network	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	9/1/2018
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Alecensa, Cotellic, Erivedge, Exkivity, Generic abiraterone, Generic lapatinib, Generic lenalidomide, Gilotrif, Hycamtin, Ibrance, Imbruvica, Kisqali, Kisqali Femara, Koselugo, Lonsurf, Mekinist, Ninlaro, Odomzo, Onureg, Pomalyst, Rozlytrek, Stivarga, Tabrecta, Tafinlar, Tukysa, Venclexta, Verzenio, Zolinza, Zydelig				
Approval Lengt	th 12 month(s)			
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization - ALL Plans Except IL and MN Plans		
Product Name	Generio	c Name	GPI	Brand/Generic
GILOTRIF	AFATINIB DIMALEATE TAB 20 MG (BASE EQUIVALENT)		21360006100320	Brand

GILOTRIF	AFATINIB DIMALEATE TAB 30 MG (BASE EQUIVALENT)	21360006100330	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 40 MG (BASE EQUIVALENT)	21360006100340	Brand
ALECENSA	ALECTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21530507100120	Brand
ONUREG	AZACITIDINE TAB 200 MG	21300003000320	Brand
ONUREG	AZACITIDINE TAB 300 MG	21300003000330	Brand
TABRECTA	CAPMATINIB HCL TAB 150 MG	21533716200320	Brand
TABRECTA	CAPMATINIB HCL TAB 200 MG	21533716200330	Brand
COTELLIC	COBIMETINIB FUMARATE TAB 20 MG (BASE EQUIVALENT)	21533530200320	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 50 MG (BASE EQUIVALENT)	21532025100120	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 75 MG (BASE EQUIVALENT)	21532025100130	Brand
TAFINLAR	DABRAFENIB MESYLATE TAB FOR ORAL SUSP 10 MG (BASE EQUIV)	21532025107320	Brand
ROZLYTREK	ENTRECTINIB CAP 100 MG	21533820000120	Brand
ROZLYTREK	ENTRECTINIB CAP 200 MG	21533820000130	Brand
IMBRUVICA	IBRUTINIB CAP 70 MG	21532133000110	Brand
IMBRUVICA	IBRUTINIB CAP 140 MG	21532133000120	Brand
IMBRUVICA	IBRUTINIB TAB 420 MG	21532133000340	Brand
IMBRUVICA	IBRUTINIB ORAL SUSP 70 MG/ML	21532133001820	Brand
ZYDELIG	IDELALISIB TAB 100 MG	21538040000320	Brand
ZYDELIG	IDELALISIB TAB 150 MG	21538040000330	Brand
NINLARO	IXAZOMIB CITRATE CAP 2.3 MG (BASE EQUIVALENT)	21536045100120	Brand
NINLARO	IXAZOMIB CITRATE CAP 3 MG (BASE EQUIVALENT)	21536045100130	Brand
NINLARO	IXAZOMIB CITRATE CAP 4 MG (BASE EQUIVALENT)	21536045100140	Brand
LAPATINIB DITOSYLATE	LAPATINIB DITOSYLATE TAB 250 MG (BASE EQUIV)	21533026100320	Generic
LENALIDOMIDE	LENALIDOMIDE CAPS 2.5 MG	99394050000110	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 5 MG	99394050000120	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 10 MG	99394050000130	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 15 MG	99394050000140	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 20 MG	99394050000145	Generic

LENALIDOMIDE	LENALIDOMIDE CAP 25 MG	99394050000150	Generic
EXKIVITY	MOBOCERTINIB SUCCINATE CAP 40 MG	21360050600120	Brand
POMALYST	POMALIDOMIDE CAP 1 MG	21450080000110	Brand
POMALYST	POMALIDOMIDE CAP 2 MG	21450080000115	Brand
POMALYST	POMALIDOMIDE CAP 3 MG	21450080000120	Brand
POMALYST	POMALIDOMIDE CAP 4 MG	21450080000125	Brand
STIVARGA	REGORAFENIB TAB 40 MG	21533050000320	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 10 MG	21533565500110	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 25 MG	21533565500125	Brand
ODOMZO	SONIDEGIB PHOSPHATE CAP 200 MG (BASE EQUIVALENT)	21370060200120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 0.25 MG (BASE EQUIV)	21550080100120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 1 MG (BASE EQUIV)	21550080100140	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 0.5 MG (BASE EQUIVALENT)	21533570100310	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 2 MG (BASE EQUIVALENT)	21533570100330	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE FOR SOLN 0.05 MG/ML (BASE EQ)	21533570102120	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 15-6.14 MG	21990002750320	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 20-8.19 MG	21990002750330	Brand
TUKYSA	TUCATINIB TAB 50 MG	21170080000320	Brand
TUKYSA	TUCATINIB TAB 150 MG	21170080000340	Brand
VENCLEXTA STARTING PACK	VENETOCLAX TAB THERAPY STARTER PACK 10 & 50 & 100 MG	2147008000B720	Brand
VENCLEXTA	VENETOCLAX TAB 10 MG	21470080000320	Brand
VENCLEXTA	VENETOCLAX TAB 50 MG	21470080000340	Brand
VENCLEXTA	VENETOCLAX TAB 100 MG	21470080000360	Brand
ERIVEDGE	VISMODEGIB CAP 150 MG	21370070000120	Brand
ZOLINZA	VORINOSTAT CAP 100 MG	21531575000120	Brand
IBRANCE	PALBOCICLIB CAP 75 MG	21531060000120	Brand
IBRANCE	PALBOCICLIB CAP 100 MG	21531060000130	Brand
IBRANCE	PALBOCICLIB CAP 125 MG	21531060000140	Brand
IBRANCE	PALBOCICLIB TAB 75 MG	21531060000320	Brand

IBRANCE	PALBOCICLIB TAB 100 MG	21531060000330	Brand
IBRANCE	PALBOCICLIB TAB 125 MG	21531060000340	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 200 MG DAILY DOSE	2153107050B720	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 400 MG DAILY DOSE (200 MG TAB)	2153107050B740	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 600 MG DAILY DOSE (200 MG TAB)	2153107050B760	Brand
KISQALI FEMARA 200 DOSE	RIBOCICLIB 200 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B730	Brand
KISQALI FEMARA 400 DOSE	RIBOCICLIB 400 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B740	Brand
KISQALI FEMARA 600 DOSE	RIBOCICLIB 600 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B760	Brand
VERZENIO	ABEMACICLIB TAB 50 MG	21531010000305	Brand
VERZENIO	ABEMACICLIB TAB 100 MG	21531010000310	Brand
VERZENIO	ABEMACICLIB TAB 150 MG	21531010000315	Brand
VERZENIO	ABEMACICLIB TAB 200 MG	21531010000320	Brand
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 250 MG	21406010200320	Generic
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 500 MG	21406010200330	Generic
ROZLYTREK	ENTRECTINIB PELLET PACK 50 MG	21533820003020	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

AND

- 2 Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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Product Name: Alecensa, Cotellic, Erivedge, Exkivity, Generic abiraterone, Generic lapatinib, Generic lenalidomide, Gilotrif, Hycamtin, Ibrance, Imbruvica, Kisqali, Kisqali Femara, Koselugo, Lonsurf, Mekinist, Ninlaro, Odomzo, Onureg, Pomalyst, Rozlytrek, Stivarga, Tabrecta, Tafinlar, Tukysa, Venclexta, Verzenio, Zolinza, Zydelig

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - ALL Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
GILOTRIF	AFATINIB DIMALEATE TAB 20 MG (BASE EQUIVALENT)	21360006100320	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 30 MG (BASE EQUIVALENT)	21360006100330	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 40 MG (BASE EQUIVALENT)	21360006100340	Brand
ALECENSA	ALECTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21530507100120	Brand
ONUREG	AZACITIDINE TAB 200 MG	21300003000320	Brand
ONUREG	AZACITIDINE TAB 300 MG	21300003000330	Brand
TABRECTA	CAPMATINIB HCL TAB 150 MG	21533716200320	Brand
TABRECTA	CAPMATINIB HCL TAB 200 MG	21533716200330	Brand
COTELLIC	COBIMETINIB FUMARATE TAB 20 MG (BASE EQUIVALENT)	21533530200320	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 50 MG (BASE EQUIVALENT)	21532025100120	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 75 MG (BASE EQUIVALENT)	21532025100130	Brand

TAFINLAR	DABRAFENIB MESYLATE TAB FOR ORAL SUSP 10 MG (BASE EQUIV)	21532025107320	Brand
ROZLYTREK	ENTRECTINIB CAP 100 MG	21533820000120	Brand
ROZLYTREK	ENTRECTINIB CAP 200 MG	21533820000130	Brand
IMBRUVICA	IBRUTINIB CAP 70 MG	21532133000110	Brand
IMBRUVICA	IBRUTINIB CAP 140 MG	21532133000120	Brand
IMBRUVICA	IBRUTINIB TAB 420 MG	21532133000340	Brand
IMBRUVICA	IBRUTINIB ORAL SUSP 70 MG/ML	21532133001820	Brand
ZYDELIG	IDELALISIB TAB 100 MG	21538040000320	Brand
ZYDELIG	IDELALISIB TAB 150 MG	21538040000330	Brand
NINLARO	IXAZOMIB CITRATE CAP 2.3 MG (BASE EQUIVALENT)	21536045100120	Brand
NINLARO	IXAZOMIB CITRATE CAP 3 MG (BASE EQUIVALENT)	21536045100130	Brand
NINLARO	IXAZOMIB CITRATE CAP 4 MG (BASE EQUIVALENT)	21536045100140	Brand
LAPATINIB DITOSYLATE	LAPATINIB DITOSYLATE TAB 250 MG (BASE EQUIV)	21533026100320	Generic
LENALIDOMIDE	LENALIDOMIDE CAPS 2.5 MG	99394050000110	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 5 MG	99394050000120	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 10 MG	99394050000130	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 15 MG	99394050000140	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 20 MG	99394050000145	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 25 MG	99394050000150	Generic
EXKIVITY	MOBOCERTINIB SUCCINATE CAP 40 MG	21360050600120	Brand
POMALYST	POMALIDOMIDE CAP 1 MG	21450080000110	Brand
POMALYST	POMALIDOMIDE CAP 2 MG	21450080000115	Brand
POMALYST	POMALIDOMIDE CAP 3 MG	21450080000120	Brand
POMALYST	POMALIDOMIDE CAP 4 MG	21450080000125	Brand
STIVARGA	REGORAFENIB TAB 40 MG	21533050000320	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 10 MG	21533565500110	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 25 MG	21533565500125	Brand
ODOMZO	SONIDEGIB PHOSPHATE CAP 200 MG (BASE EQUIVALENT)	21370060200120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 0.25 MG (BASE EQUIV)	21550080100120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 1 MG (BASE EQUIV)	21550080100140	Brand

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MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 0.5 MG (BASE EQUIVALENT)	21533570100310	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 2 MG (BASE EQUIVALENT)	21533570100330	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE FOR SOLN 0.05 MG/ML (BASE EQ)	21533570102120	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 15-6.14 MG	21990002750320	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 20-8.19 MG	21990002750330	Brand
TUKYSA	TUCATINIB TAB 50 MG	21170080000320	Brand
TUKYSA	TUCATINIB TAB 150 MG	21170080000340	Brand
VENCLEXTA STARTING PACK	VENETOCLAX TAB THERAPY STARTER PACK 10 & 50 & 100 MG	2147008000B720	Brand
VENCLEXTA	VENETOCLAX TAB 10 MG	21470080000320	Brand
VENCLEXTA	VENETOCLAX TAB 50 MG	21470080000340	Brand
VENCLEXTA	VENETOCLAX TAB 100 MG	21470080000360	Brand
ERIVEDGE	VISMODEGIB CAP 150 MG	21370070000120	Brand
ZOLINZA	VORINOSTAT CAP 100 MG	21531575000120	Brand
IBRANCE	PALBOCICLIB CAP 75 MG	21531060000120	Brand
IBRANCE	PALBOCICLIB CAP 100 MG	21531060000130	Brand
IBRANCE	PALBOCICLIB CAP 125 MG	21531060000140	Brand
IBRANCE	PALBOCICLIB TAB 75 MG	21531060000320	Brand
IBRANCE	PALBOCICLIB TAB 100 MG	21531060000330	Brand
IBRANCE	PALBOCICLIB TAB 125 MG	21531060000340	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 200 MG DAILY DOSE	2153107050B720	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 400 MG DAILY DOSE (200 MG TAB)	2153107050B740	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 600 MG DAILY DOSE (200 MG TAB)	2153107050B760	Brand
KISQALI FEMARA 200 DOSE	RIBOCICLIB 200 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B730	Brand
KISQALI FEMARA 400 DOSE	RIBOCICLIB 400 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B740	Brand
KISQALI FEMARA 600 DOSE	RIBOCICLIB 600 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B760	Brand
VERZENIO	ABEMACICLIB TAB 50 MG	21531010000305	Brand

VERZENIO	ABEMACICLIB TAB 100 MG	21531010000310	Brand
VERZENIO	ABEMACICLIB TAB 150 MG	21531010000315	Brand
VERZENIO	ABEMACICLIB TAB 200 MG	21531010000320	Brand
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 250 MG	21406010200320	Generic
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 500 MG	21406010200330	Generic
ROZLYTREK	ENTRECTINIB PELLET PACK 50 MG	21533820003020	Brand

- 1 One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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Product Name: Alecensa, Bosulif, Cabometyx, Cotellic, Erivedge, Exkivity, Generic abiraterone, Generic erlotinib, Generic everolimus, Generic gefitinib, Generic lapatinib, Generic lenalidomide, Generic sorafenib, Generic sunitinib, Gilotrif, Hycamtin, Ibrance, Imbruvica, Inlyta, Inrebic, Jakafi, Kisqali, Kisqali Femara, Koselugo, Lonsurf, Lynparza, Mekinist, Ninlaro, Nubeqa, Odomzo, Onureg, Generic pazopanib, Piqray, Pomalyst, Retevmo, Rozlytrek, Rubraca, Sprycel, Stivarga, Tabrecta, Tafinlar, Tagrisso, Tasigna, Tazverik, Tukysa, Venclexta, Verzenio, Xalkori, Xtandi, Zejula, Zelboraf, Zolinza, Zydelig, Zykadia

Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL Plans

Product Name	Generic Name	GPI	Brand/Generic
GILOTRIF	AFATINIB DIMALEATE TAB 20 MG (BASE EQUIVALENT)	21360006100320	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 30 MG (BASE EQUIVALENT)	21360006100330	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 40 MG (BASE EQUIVALENT)	21360006100340	Brand
ALECENSA	ALECTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21530507100120	Brand
ONUREG	AZACITIDINE TAB 200 MG	21300003000320	Brand
ONUREG	AZACITIDINE TAB 300 MG	21300003000330	Brand
TABRECTA	CAPMATINIB HCL TAB 150 MG	21533716200320	Brand
TABRECTA	CAPMATINIB HCL TAB 200 MG	21533716200330	Brand
COTELLIC	COBIMETINIB FUMARATE TAB 20 MG (BASE EQUIVALENT)	21533530200320	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 50 MG (BASE EQUIVALENT)	21532025100120	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 75 MG (BASE EQUIVALENT)	21532025100130	Brand
TAFINLAR	DABRAFENIB MESYLATE TAB FOR ORAL SUSP 10 MG (BASE EQUIV)	21532025107320	Brand
ROZLYTREK	ENTRECTINIB CAP 100 MG	21533820000120	Brand
ROZLYTREK	ENTRECTINIB CAP 200 MG	21533820000130	Brand
IMBRUVICA	IBRUTINIB CAP 70 MG	21532133000110	Brand
IMBRUVICA	IBRUTINIB CAP 140 MG	21532133000120	Brand
IMBRUVICA	IBRUTINIB TAB 420 MG	21532133000340	Brand
IMBRUVICA	IBRUTINIB ORAL SUSP 70 MG/ML	21532133001820	Brand
ZYDELIG	IDELALISIB TAB 100 MG	21538040000320	Brand
ZYDELIG	IDELALISIB TAB 150 MG	21538040000330	Brand
NINLARO	IXAZOMIB CITRATE CAP 2.3 MG (BASE EQUIVALENT)	21536045100120	Brand
NINLARO	IXAZOMIB CITRATE CAP 3 MG (BASE EQUIVALENT)	21536045100130	Brand
NINLARO	IXAZOMIB CITRATE CAP 4 MG (BASE EQUIVALENT)	21536045100140	Brand

LADATINES	LABATINIB BITOOM ATE TAB OF AND (5.05		. 1
LAPATINIB DITOSYLATE	LAPATINIB DITOSYLATE TAB 250 MG (BASE EQUIV)	21533026100320	Generic
LENALIDOMIDE	LENALIDOMIDE CAPS 2.5 MG	99394050000110	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 5 MG	99394050000120	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 10 MG	99394050000130	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 15 MG	99394050000140	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 20 MG	99394050000145	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 25 MG	99394050000150	Generic
EXKIVITY	MOBOCERTINIB SUCCINATE CAP 40 MG	21360050600120	Brand
POMALYST	POMALIDOMIDE CAP 1 MG	21450080000110	Brand
POMALYST	POMALIDOMIDE CAP 2 MG	21450080000115	Brand
POMALYST	POMALIDOMIDE CAP 3 MG	21450080000120	Brand
POMALYST	POMALIDOMIDE CAP 4 MG	21450080000125	Brand
STIVARGA	REGORAFENIB TAB 40 MG	21533050000320	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 10 MG	21533565500110	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 25 MG	21533565500125	Brand
ODOMZO	SONIDEGIB PHOSPHATE CAP 200 MG (BASE EQUIVALENT)	21370060200120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 0.25 MG (BASE EQUIV)	21550080100120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 1 MG (BASE EQUIV)	21550080100140	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 0.5 MG (BASE EQUIVALENT)	21533570100310	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 2 MG (BASE EQUIVALENT)	21533570100330	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE FOR SOLN 0.05 MG/ML (BASE EQ)	21533570102120	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 15-6.14 MG	21990002750320	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 20-8.19 MG	21990002750330	Brand
TUKYSA	TUCATINIB TAB 50 MG	21170080000320	Brand
TUKYSA	TUCATINIB TAB 150 MG	21170080000340	Brand
VENCLEXTA STARTING PACK	VENETOCLAX TAB THERAPY STARTER PACK 10 & 50 & 100 MG	2147008000B720	Brand
VENCLEXTA	VENETOCLAX TAB 10 MG	21470080000320	Brand
VENCLEXTA	VENETOCLAX TAB 50 MG	21470080000340	Brand
VENCLEXTA	VENETOCLAX TAB 100 MG	21470080000360	Brand
ERIVEDGE	VISMODEGIB CAP 150 MG	21370070000120	Brand

ZOLINZA	VORINOSTAT CAP 100 MG	21531575000120	Brand
IBRANCE	PALBOCICLIB CAP 75 MG	21531060000120	Brand
IBRANCE	PALBOCICLIB CAP 100 MG	21531060000130	Brand
IBRANCE	PALBOCICLIB CAP 125 MG	21531060000140	Brand
IBRANCE	PALBOCICLIB TAB 75 MG	21531060000320	Brand
IBRANCE	PALBOCICLIB TAB 100 MG	21531060000330	Brand
IBRANCE	PALBOCICLIB TAB 125 MG	21531060000340	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 200 MG DAILY DOSE	2153107050B720	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 400 MG DAILY DOSE (200 MG TAB)	2153107050B740	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 600 MG DAILY DOSE (200 MG TAB)	2153107050B760	Brand
KISQALI FEMARA 200 DOSE	RIBOCICLIB 200 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B730	Brand
KISQALI FEMARA 400 DOSE	RIBOCICLIB 400 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B740	Brand
KISQALI FEMARA 600 DOSE	RIBOCICLIB 600 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B760	Brand
VERZENIO	ABEMACICLIB TAB 50 MG	21531010000305	Brand
VERZENIO	ABEMACICLIB TAB 100 MG	21531010000310	Brand
VERZENIO	ABEMACICLIB TAB 150 MG	21531010000315	Brand
VERZENIO	ABEMACICLIB TAB 200 MG	21531010000320	Brand
PIQRAY 200MG DAILY DOSE	ALPELISIB TAB THERAPY PACK 200 MG DAILY DOSE	2153801000B720	Brand
PIQRAY 250MG DAILY DOSE	ALPELISIB TAB PACK 250 MG DAILY DOSE (200 MG & 50 MG TABS)	2153801000B725	Brand
PIQRAY 300MG DAILY DOSE	ALPELISIB TAB PACK 300 MG DAILY DOSE (2X150 MG TAB)	2153801000B730	Brand
INLYTA	AXITINIB TAB 1 MG	21335013000320	Brand
INLYTA	AXITINIB TAB 5 MG	21335013000340	Brand
BOSULIF	BOSUTINIB TAB 100 MG	21531812000320	Brand
BOSULIF	BOSUTINIB TAB 400 MG	21531812000327	Brand
BOSULIF	BOSUTINIB TAB 500 MG	21531812000340	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 20 MG (BASE EQUIVALENT)	21533010100320	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 40 MG (BASE EQUIVALENT)	21533010100330	Brand

CABOMETYX	CABOZANTINIB S-MALATE TAB 60 MG (BASE EQUIVALENT)	21533010100340	Brand
ZYKADIA	CERITINIB TAB 150 MG	21530514000330	Brand
XALKORI	CRIZOTINIB CAP 200 MG	21530517000120	Brand
XALKORI	CRIZOTINIB CAP 250 MG	21530517000125	Brand
NUBEQA	DAROLUTAMIDE TAB 300 MG	21402425000320	Brand
SPRYCEL	DASATINIB TAB 20 MG	21531820000320	Brand
SPRYCEL	DASATINIB TAB 50 MG	21531820000340	Brand
SPRYCEL	DASATINIB TAB 70 MG	21531820000350	Brand
SPRYCEL	DASATINIB TAB 80 MG	21531820000354	Brand
SPRYCEL	DASATINIB TAB 100 MG	21531820000360	Brand
SPRYCEL	DASATINIB TAB 140 MG	21531820000380	Brand
XTANDI	ENZALUTAMIDE CAP 40 MG	21402430000120	Brand
XTANDI	ENZALUTAMIDE TAB 40 MG	21402430000320	Brand
XTANDI	ENZALUTAMIDE TAB 80 MG	21402430000340	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Generic
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Generic
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Generic
EVEROLIMUS	EVEROLIMUS TAB 2.5 MG	21532530000310	Generic
EVEROLIMUS	EVEROLIMUS TAB 5 MG	21532530000320	Generic
EVEROLIMUS	EVEROLIMUS TAB 7.5 MG	21532530000325	Generic
EVEROLIMUS	EVEROLIMUS TAB 10 MG	21532530000330	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Generic
INREBIC	FEDRATINIB HCL CAP 100 MG	21537520200120	Brand
GEFITINIB	GEFITINIB TAB 250 MG	21360030000320	Generic
TASIGNA	NILOTINIB HCL CAP 50 MG (BASE EQUIVALENT)	21531860200110	Brand
TASIGNA	NILOTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21531860200115	Brand
TASIGNA	NILOTINIB HCL CAP 200 MG (BASE EQUIVALENT)	21531860200125	Brand

ZEJULA	NIRAPARIB TOSYLATE TAB 100 MG (BASE EQUIVALENT)	21535550200320	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21535550200330	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 300 MG (BASE EQUIVALENT)	21535550200340	Brand
LYNPARZA	OLAPARIB TAB 100 MG	21535560000330	Brand
LYNPARZA	OLAPARIB TAB 150 MG	21535560000340	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 40 MG (BASE EQUIVALENT)	21360068200320	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 80 MG (BASE EQUIVALENT)	21360068200330	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 200 MG (BASE EQUIVALENT)	21535570200320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 250 MG (BASE EQUIVALENT)	21535570200325	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 300 MG (BASE EQUIVALENT)	21535570200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 5 MG (BASE EQUIVALENT)	21537560200310	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	21537560200320	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 15 MG (BASE EQUIVALENT)	21537560200325	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 20 MG (BASE EQUIVALENT)	21537560200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 25 MG (BASE EQUIVALENT)	21537560200335	Brand
RETEVMO	SELPERCATINIB CAP 40 MG	21535779000120	Brand
RETEVMO	SELPERCATINIB CAP 80 MG	21535779000140	Brand
SORAFENIB	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SORAFENIB TOSYLATE	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Generic
TAZVERIK	TAZEMETOSTAT HBR TAB 200 MG	21533675200320	Brand

ZELBORAF	VEMURAFENIB TAB 240 MG	21532080000320	Brand
BOSULIF	BOSUTINIB CAP 100 MG	21531812000130	Brand
BOSULIF	BOSUTINIB CAP 50 MG	21531812000120	Brand
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 250 MG	21406010200320	Generic
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 500 MG	21406010200330	Generic
ROZLYTREK	ENTRECTINIB PELLET PACK 50 MG	21533820003020	Brand
PAZOPANIB HYDROCHLORIDE	PAZOPANIB HCL TAB 200 MG (BASE EQUIV)	21533042100320	Generic
XALKORI	CRIZOTINIB CAP SPRINKLE 20 MG	21530517006820	Brand
XALKORI	CRIZOTINIB CAP SPRINKLE 50 MG	21530517006830	Brand
XALKORI	CRIZOTINIB CAP SPRINKLE 150 MG	21530517006850	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member* in one of the following:
 - American Hospital Formulary Service Drug Information
 - Thompson Micromedex's Drug Dex
 - Elsevier Gold Standard's Clinical Pharmacology
 - Two articles in peer-reviewed professional medical journals from the United States or Great Britain that recognize the safety and efficacy of the requested drug in the member's specific condition

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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Product Name: Alecensa, Bosulif, Cabometyx, Cotellic, Erivedge, Exkivity, Generic abiraterone, Generic erlotinib, Generic everolimus, Generic gefitinib, Generic lapatinib, Generic lenalidomide, Generic sorafenib, Generic sunitinib, Gilotrif, Hycamtin, Ibrance, Imbruvica, Inlyta, Inrebic, Jakafi, Kisqali, Kisqali Femara, Koselugo, Lonsurf, Lynparza, Mekinist, Ninlaro, Nubeqa, Odomzo, Onureg, Generic pazopanib, Piqray, Pomalyst, Retevmo, Rozlytrek, Rubraca, Sprycel, Stivarga, Tabrecta, Tafinlar, Tagrisso, Tasigna, Tazverik, Tukysa, Venclexta, Verzenio, Xalkori, Xtandi, Zejula, Zelboraf, Zolinza, Zydelig, Zykadia

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL Plans

Product Name	Generic Name	GPI	Brand/Generic
GILOTRIF	AFATINIB DIMALEATE TAB 20 MG (BASE EQUIVALENT)	21360006100320	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 30 MG (BASE EQUIVALENT)	21360006100330	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 40 MG (BASE EQUIVALENT)	21360006100340	Brand
ALECENSA	ALECTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21530507100120	Brand
ONUREG	AZACITIDINE TAB 200 MG	21300003000320	Brand
ONUREG	AZACITIDINE TAB 300 MG	21300003000330	Brand
TABRECTA	CAPMATINIB HCL TAB 150 MG	21533716200320	Brand
TABRECTA	CAPMATINIB HCL TAB 200 MG	21533716200330	Brand
COTELLIC	COBIMETINIB FUMARATE TAB 20 MG (BASE EQUIVALENT)	21533530200320	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 50 MG (BASE EQUIVALENT)	21532025100120	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 75 MG (BASE EQUIVALENT)	21532025100130	Brand

TAFINLAR	DABRAFENIB MESYLATE TAB FOR ORAL SUSP 10 MG (BASE EQUIV)	21532025107320	Brand
ROZLYTREK	ENTRECTINIB CAP 100 MG	21533820000120	Brand
ROZLYTREK	ENTRECTINIB CAP 200 MG	21533820000130	Brand
IMBRUVICA	IBRUTINIB CAP 70 MG	21532133000110	Brand
IMBRUVICA	IBRUTINIB CAP 140 MG	21532133000120	Brand
IMBRUVICA	IBRUTINIB TAB 420 MG	21532133000340	Brand
IMBRUVICA	IBRUTINIB ORAL SUSP 70 MG/ML	21532133001820	Brand
ZYDELIG	IDELALISIB TAB 100 MG	21538040000320	Brand
ZYDELIG	IDELALISIB TAB 150 MG	21538040000330	Brand
NINLARO	IXAZOMIB CITRATE CAP 2.3 MG (BASE EQUIVALENT)	21536045100120	Brand
NINLARO	IXAZOMIB CITRATE CAP 3 MG (BASE EQUIVALENT)	21536045100130	Brand
NINLARO	IXAZOMIB CITRATE CAP 4 MG (BASE EQUIVALENT)	21536045100140	Brand
LAPATINIB DITOSYLATE	LAPATINIB DITOSYLATE TAB 250 MG (BASE EQUIV)	21533026100320	Generic
LENALIDOMIDE	LENALIDOMIDE CAPS 2.5 MG	99394050000110	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 5 MG	99394050000120	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 10 MG	99394050000130	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 15 MG	99394050000140	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 20 MG	99394050000145	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 25 MG	99394050000150	Generic
EXKIVITY	MOBOCERTINIB SUCCINATE CAP 40 MG	21360050600120	Brand
POMALYST	POMALIDOMIDE CAP 1 MG	21450080000110	Brand
POMALYST	POMALIDOMIDE CAP 2 MG	21450080000115	Brand
POMALYST	POMALIDOMIDE CAP 3 MG	21450080000120	Brand
POMALYST	POMALIDOMIDE CAP 4 MG	21450080000125	Brand
STIVARGA	REGORAFENIB TAB 40 MG	21533050000320	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 10 MG	21533565500110	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 25 MG	21533565500125	Brand
ODOMZO	SONIDEGIB PHOSPHATE CAP 200 MG (BASE EQUIVALENT)	21370060200120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 0.25 MG (BASE EQUIV)	21550080100120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 1 MG (BASE EQUIV)	21550080100140	Brand

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MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 0.5 MG (BASE EQUIVALENT)	21533570100310	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 2 MG (BASE EQUIVALENT)	21533570100330	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE FOR SOLN 0.05 MG/ML (BASE EQ)	21533570102120	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 15-6.14 MG	21990002750320	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 20-8.19 MG	21990002750330	Brand
TUKYSA	TUCATINIB TAB 50 MG	21170080000320	Brand
TUKYSA	TUCATINIB TAB 150 MG	21170080000340	Brand
VENCLEXTA STARTING PACK	VENETOCLAX TAB THERAPY STARTER PACK 10 & 50 & 100 MG	2147008000B720	Brand
VENCLEXTA	VENETOCLAX TAB 10 MG	21470080000320	Brand
VENCLEXTA	VENETOCLAX TAB 50 MG	21470080000340	Brand
VENCLEXTA	VENETOCLAX TAB 100 MG	21470080000360	Brand
ERIVEDGE	VISMODEGIB CAP 150 MG	21370070000120	Brand
ZOLINZA	VORINOSTAT CAP 100 MG	21531575000120	Brand
IBRANCE	PALBOCICLIB CAP 75 MG	21531060000120	Brand
IBRANCE	PALBOCICLIB CAP 100 MG	21531060000130	Brand
IBRANCE	PALBOCICLIB CAP 125 MG	21531060000140	Brand
IBRANCE	PALBOCICLIB TAB 75 MG	21531060000320	Brand
IBRANCE	PALBOCICLIB TAB 100 MG	21531060000330	Brand
IBRANCE	PALBOCICLIB TAB 125 MG	21531060000340	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 200 MG DAILY DOSE	2153107050B720	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 400 MG DAILY DOSE (200 MG TAB)	2153107050B740	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 600 MG DAILY DOSE (200 MG TAB)	2153107050B760	Brand
KISQALI FEMARA 200 DOSE	RIBOCICLIB 200 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B730	Brand
KISQALI FEMARA 400 DOSE	RIBOCICLIB 400 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B740	Brand
KISQALI FEMARA 600 DOSE	RIBOCICLIB 600 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B760	Brand
VERZENIO	ABEMACICLIB TAB 50 MG	21531010000305	Brand
VERZENIO	ABEMACICLIB TAB 100 MG	21531010000310	Brand
VERZENIO	ABEMACICLIB TAB 150 MG	21531010000315	Brand

VERZENIO	ABEMACICLIB TAB 200 MG	21531010000320	Brand
PIQRAY 200MG	ALPELISIB TAB THERAPY PACK 200 MG	21538010000320 2153801000B720	Brand
DAILY DOSE	DAILY DOSE	2133001000B720	Dialia
PIQRAY 250MG DAILY DOSE	ALPELISIB TAB PACK 250 MG DAILY DOSE (200 MG & 50 MG TABS)	2153801000B725	Brand
PIQRAY 300MG DAILY DOSE	ALPELISIB TAB PACK 300 MG DAILY DOSE (2X150 MG TAB)	2153801000B730	Brand
INLYTA	AXITINIB TAB 1 MG	21335013000320	Brand
INLYTA	AXITINIB TAB 5 MG	21335013000340	Brand
BOSULIF	BOSUTINIB TAB 100 MG	21531812000320	Brand
BOSULIF	BOSUTINIB TAB 400 MG	21531812000327	Brand
BOSULIF	BOSUTINIB TAB 500 MG	21531812000340	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 20 MG (BASE EQUIVALENT)	21533010100320	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 40 MG (BASE EQUIVALENT)	21533010100330	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 60 MG (BASE EQUIVALENT)	21533010100340	Brand
ZYKADIA	CERITINIB TAB 150 MG	21530514000330	Brand
XALKORI	CRIZOTINIB CAP 200 MG	21530517000120	Brand
XALKORI	CRIZOTINIB CAP 250 MG	21530517000125	Brand
NUBEQA	DAROLUTAMIDE TAB 300 MG	21402425000320	Brand
SPRYCEL	DASATINIB TAB 20 MG	21531820000320	Brand
SPRYCEL	DASATINIB TAB 50 MG	21531820000340	Brand
SPRYCEL	DASATINIB TAB 70 MG	21531820000350	Brand
SPRYCEL	DASATINIB TAB 80 MG	21531820000354	Brand
SPRYCEL	DASATINIB TAB 100 MG	21531820000360	Brand
SPRYCEL	DASATINIB TAB 140 MG	21531820000380	Brand
XTANDI	ENZALUTAMIDE CAP 40 MG	21402430000120	Brand
XTANDI	ENZALUTAMIDE TAB 40 MG	21402430000320	Brand
XTANDI	ENZALUTAMIDE TAB 80 MG	21402430000340	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Generic
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Generic
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Generic

EVEROLIMUS	EVEROLIMUS TAB 2.5 MG	21532530000310	Generic
EVEROLIMUS	EVEROLIMUS TAB 5 MG	21532530000320	Generic
EVEROLIMUS	EVEROLIMUS TAB 7.5 MG	21532530000325	Generic
EVEROLIMUS	EVEROLIMUS TAB 10 MG	21532530000330	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Generic
INREBIC	FEDRATINIB HCL CAP 100 MG	21537520200120	Brand
GEFITINIB	GEFITINIB TAB 250 MG	21360030000320	Generic
TASIGNA	NILOTINIB HCL CAP 50 MG (BASE EQUIVALENT)	21531860200110	Brand
TASIGNA	NILOTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21531860200115	Brand
TASIGNA	NILOTINIB HCL CAP 200 MG (BASE EQUIVALENT)	21531860200125	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 100 MG (BASE EQUIVALENT)	21535550200320	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21535550200330	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 300 MG (BASE EQUIVALENT)	21535550200340	Brand
LYNPARZA	OLAPARIB TAB 100 MG	21535560000330	Brand
LYNPARZA	OLAPARIB TAB 150 MG	21535560000340	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 40 MG (BASE EQUIVALENT)	21360068200320	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 80 MG (BASE EQUIVALENT)	21360068200330	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 200 MG (BASE EQUIVALENT)	21535570200320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 250 MG (BASE EQUIVALENT)	21535570200325	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 300 MG (BASE EQUIVALENT)	21535570200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 5 MG (BASE EQUIVALENT)	21537560200310	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	21537560200320	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 15 MG (BASE EQUIVALENT)	21537560200325	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 20 MG (BASE EQUIVALENT)	21537560200330	Brand

JAKAFI	RUXOLITINIB PHOSPHATE TAB 25 MG (BASE EQUIVALENT)	21537560200335	Brand
RETEVMO	SELPERCATINIB CAP 40 MG	21535779000120	Brand
RETEVMO	SELPERCATINIB CAP 80 MG	21535779000140	Brand
SORAFENIB	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SORAFENIB TOSYLATE	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Generic
TAZVERIK	TAZEMETOSTAT HBR TAB 200 MG	21533675200320	Brand
ZELBORAF	VEMURAFENIB TAB 240 MG	21532080000320	Brand
BOSULIF	BOSUTINIB CAP 50 MG	21531812000120	Brand
BOSULIF	BOSUTINIB CAP 100 MG	21531812000130	Brand
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 250 MG	21406010200320	Generic
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 500 MG	21406010200330	Generic
ROZLYTREK	ENTRECTINIB PELLET PACK 50 MG	21533820003020	Brand
PAZOPANIB HYDROCHLORIDE	PAZOPANIB HCL TAB 200 MG (BASE EQUIV)	21533042100320	Generic
Xalkori	CRIZOTINIB CAP SPRINKLE 20 MG		
Xalkori	CRIZOTINIB CAP SPRINKLE 50 MG		
Xalkori	CRIZOTINIB CAP SPRINKLE 150 MG	21530517006850	Brand

- 1 One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member* in one of the following:
 - American Hospital Formulary Service Drug Information
 - Thompson Micromedex's Drug Dex
 - Elsevier Gold Standard's Clinical Pharmacology
 - Two articles in peer-reviewed professional medical journals from the United States or Great Britain that recognize the safety and efficacy of the requested drug in the member's specific condition

AND

- 2 Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.

Product Name: Alecensa, Cotellic, Erivedge, Exkivity, Generic abiraterone, Generic Iapatinib, Generic Ienalidomide, Gilotrif, Hycamtin, Ibrance, Imbruvica, Kisqali, Kisqali Femara, Koselugo, Lonsurf, Mekinist, Ninlaro, Odomzo, Onureg, Pomalyst, Rozlytrek, Stivarga, Tabrecta, Tafinlar, Tukysa, Venclexta, Verzenio, Zolinza, Zydelig

Approval Length 12 month(s)

Therapy Stage Initial Authorization

Guideline Type Prior Authorization - MN Plans

Product Reneric Name GPI Brand/Generic Name

GILOTRIF	AFATINIB DIMALEATE TAB 20 MG (BASE EQUIVALENT)	21360006100320	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 30 MG (BASE EQUIVALENT)	21360006100330	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 40 MG (BASE EQUIVALENT)	21360006100340	Brand
ALECENSA	ALECTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21530507100120	Brand
ONUREG	AZACITIDINE TAB 200 MG	21300003000320	Brand
ONUREG	AZACITIDINE TAB 300 MG	21300003000330	Brand
TABRECTA	CAPMATINIB HCL TAB 150 MG	21533716200320	Brand
TABRECTA	CAPMATINIB HCL TAB 200 MG	21533716200330	Brand
COTELLIC	COBIMETINIB FUMARATE TAB 20 MG (BASE EQUIVALENT)	21533530200320	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 50 MG (BASE EQUIVALENT)	21532025100120	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 75 MG (BASE EQUIVALENT)	21532025100130	Brand
TAFINLAR	DABRAFENIB MESYLATE TAB FOR ORAL SUSP 10 MG (BASE EQUIV)	21532025107320	Brand
ROZLYTREK	ENTRECTINIB CAP 100 MG	21533820000120	Brand
ROZLYTREK	ENTRECTINIB CAP 200 MG	21533820000130	Brand
IMBRUVICA	IBRUTINIB CAP 70 MG	21532133000110	Brand
IMBRUVICA	IBRUTINIB CAP 140 MG	21532133000120	Brand
IMBRUVICA	IBRUTINIB TAB 420 MG	21532133000340	Brand
IMBRUVICA	IBRUTINIB ORAL SUSP 70 MG/ML	21532133001820	Brand
ZYDELIG	IDELALISIB TAB 100 MG	21538040000320	Brand
ZYDELIG	IDELALISIB TAB 150 MG	21538040000330	Brand
NINLARO	IXAZOMIB CITRATE CAP 2.3 MG (BASE EQUIVALENT)	21536045100120	Brand
NINLARO	IXAZOMIB CITRATE CAP 3 MG (BASE EQUIVALENT)	21536045100130	Brand
NINLARO	IXAZOMIB CITRATE CAP 4 MG (BASE EQUIVALENT)	21536045100140	Brand
LAPATINIB DITOSYLATE	LAPATINIB DITOSYLATE TAB 250 MG (BASE EQUIV)	21533026100320	Generic
LENALIDOMIDE	LENALIDOMIDE CAPS 2.5 MG	99394050000110	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 5 MG	99394050000120	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 10 MG	99394050000130	Generic

LENALIDOMIDE	LENALIDOMIDE CAP 15 MG	99394050000140	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 20 MG	99394050000145	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 25 MG	99394050000150	Generic
EXKIVITY	MOBOCERTINIB SUCCINATE CAP 40 MG	21360050600120	Brand
POMALYST	POMALIDOMIDE CAP 1 MG	21450080000110	Brand
POMALYST	POMALIDOMIDE CAP 2 MG	21450080000115	Brand
POMALYST	POMALIDOMIDE CAP 3 MG	21450080000120	Brand
POMALYST	POMALIDOMIDE CAP 4 MG	21450080000125	Brand
STIVARGA	REGORAFENIB TAB 40 MG	21533050000320	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 10 MG	21533565500110	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 25 MG	21533565500125	Brand
ODOMZO	SONIDEGIB PHOSPHATE CAP 200 MG (BASE EQUIVALENT)	21370060200120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 0.25 MG (BASE EQUIV)	21550080100120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 1 MG (BASE EQUIV)	21550080100140	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 0.5 MG (BASE EQUIVALENT)	21533570100310	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 2 MG (BASE EQUIVALENT)	21533570100330	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE FOR SOLN 0.05 MG/ML (BASE EQ)	21533570102120	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 15-6.14 MG	21990002750320	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 20-8.19 MG	21990002750330	Brand
TUKYSA	TUCATINIB TAB 50 MG	21170080000320	Brand
TUKYSA	TUCATINIB TAB 150 MG	21170080000340	Brand
VENCLEXTA STARTING PACK	VENETOCLAX TAB THERAPY STARTER PACK 10 & 50 & 100 MG	2147008000B720	Brand
VENCLEXTA	VENETOCLAX TAB 10 MG	21470080000320	Brand
VENCLEXTA	VENETOCLAX TAB 50 MG	21470080000340	Brand
VENCLEXTA	VENETOCLAX TAB 100 MG	21470080000360	Brand
ERIVEDGE	VISMODEGIB CAP 150 MG	21370070000120	Brand
ZOLINZA	VORINOSTAT CAP 100 MG	21531575000120	Brand
IBRANCE	PALBOCICLIB CAP 75 MG	21531060000120	Brand
IBRANCE	PALBOCICLIB CAP 100 MG	21531060000130	Brand

IBRANCE	PALBOCICLIB CAP 125 MG	21531060000140	Brand
IBRANCE	PALBOCICLIB TAB 75 MG	21531060000320	Brand
IBRANCE	PALBOCICLIB TAB 100 MG	21531060000330	Brand
IBRANCE	PALBOCICLIB TAB 125 MG	21531060000340	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 200 MG DAILY DOSE	2153107050B720	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 400 MG DAILY DOSE (200 MG TAB)	2153107050B740	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 600 MG DAILY DOSE (200 MG TAB)	2153107050B760	Brand
KISQALI FEMARA 200 DOSE	RIBOCICLIB 200 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B730	Brand
KISQALI FEMARA 400 DOSE	RIBOCICLIB 400 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B740	Brand
KISQALI FEMARA 600 DOSE	RIBOCICLIB 600 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B760	Brand
VERZENIO	ABEMACICLIB TAB 50 MG	21531010000305	Brand
VERZENIO	ABEMACICLIB TAB 100 MG	21531010000310	Brand
VERZENIO	ABEMACICLIB TAB 150 MG	21531010000315	Brand
VERZENIO	ABEMACICLIB TAB 200 MG	21531010000320	Brand
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 250 MG	21406010200320	Generic
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 500 MG	21406010200330	Generic
ROZLYTREK	ENTRECTINIB PELLET PACK 50 MG	21533820003020	Brand

- 1 One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2

recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member* in one of the following:
 - United States Pharmacopeia Drug Information
 - American Hospital Formulary Service Drug Information
 - One article in a major peer- reviewed medical journal that recognizes the safety and efficacy of the requested drug in the member's specific condition

AND

- 2 Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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Product Name: Alecensa, Cotellic, Erivedge, Exkivity, Generic abiraterone, Generic lapatinib, Generic lenalidomide, Gilotrif, Hycamtin, Ibrance, Imbruvica, Kisqali, Kisqali Femara, Koselugo, Lonsurf, Mekinist, Ninlaro, Odomzo, Onureg, Pomalyst, Rozlytrek, Stivarga, Tabrecta, Tafinlar, Tukysa, Venclexta, Verzenio, Zolinza, Zydelig

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - MN Plans

Product Name	Generic Name	GPI	Brand/Generic
GILOTRIF	AFATINIB DIMALEATE TAB 20 MG (BASE EQUIVALENT)	21360006100320	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 30 MG (BASE EQUIVALENT)	21360006100330	Brand
GILOTRIF	AFATINIB DIMALEATE TAB 40 MG (BASE EQUIVALENT)	21360006100340	Brand
ALECENSA	ALECTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21530507100120	Brand

ONUREG	AZACITIDINE TAB 200 MG	21300003000320	Brand
ONUREG	AZACITIDINE TAB 300 MG	21300003000330	Brand
TABRECTA	CAPMATINIB HCL TAB 150 MG	21533716200320	Brand
TABRECTA	CAPMATINIB HCL TAB 200 MG	21533716200330	Brand
COTELLIC	COBIMETINIB FUMARATE TAB 20 MG (BASE EQUIVALENT)	21533530200320	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 50 MG (BASE EQUIVALENT)	21532025100120	Brand
TAFINLAR	DABRAFENIB MESYLATE CAP 75 MG (BASE EQUIVALENT)	21532025100130	Brand
TAFINLAR	DABRAFENIB MESYLATE TAB FOR ORAL SUSP 10 MG (BASE EQUIV)	21532025107320	Brand
ROZLYTREK	ENTRECTINIB CAP 100 MG	21533820000120	Brand
ROZLYTREK	ENTRECTINIB CAP 200 MG	21533820000130	Brand
IMBRUVICA	IBRUTINIB CAP 70 MG	21532133000110	Brand
IMBRUVICA	IBRUTINIB CAP 140 MG	21532133000120	Brand
IMBRUVICA	IBRUTINIB TAB 420 MG	21532133000340	Brand
IMBRUVICA	IBRUTINIB ORAL SUSP 70 MG/ML	21532133001820	Brand
ZYDELIG	IDELALISIB TAB 100 MG	21538040000320	Brand
ZYDELIG	IDELALISIB TAB 150 MG	21538040000330	Brand
NINLARO	IXAZOMIB CITRATE CAP 2.3 MG (BASE EQUIVALENT)	21536045100120	Brand
NINLARO	IXAZOMIB CITRATE CAP 3 MG (BASE EQUIVALENT)	21536045100130	Brand
NINLARO	IXAZOMIB CITRATE CAP 4 MG (BASE EQUIVALENT)	21536045100140	Brand
LAPATINIB DITOSYLATE	LAPATINIB DITOSYLATE TAB 250 MG (BASE EQUIV)	21533026100320	Generic
LENALIDOMIDE	LENALIDOMIDE CAPS 2.5 MG	99394050000110	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 5 MG	99394050000120	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 10 MG	99394050000130	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 15 MG	99394050000140	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 20 MG	99394050000145	Generic
LENALIDOMIDE	LENALIDOMIDE CAP 25 MG	99394050000150	Generic
EXKIVITY	MOBOCERTINIB SUCCINATE CAP 40 MG	21360050600120	Brand
POMALYST	POMALIDOMIDE CAP 1 MG	21450080000110	Brand
POMALYST	POMALIDOMIDE CAP 2 MG	21450080000115	Brand

POMALYST	POMALIDOMIDE CAP 3 MG	21450080000120	Brand
POMALYST	POMALIDOMIDE CAP 4 MG	21450080000125	Brand
STIVARGA	REGORAFENIB TAB 40 MG	21533050000320	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 10 MG	21533565500110	Brand
KOSELUGO	SELUMETINIB SULFATE CAP 25 MG	21533565500125	Brand
ODOMZO	SONIDEGIB PHOSPHATE CAP 200 MG (BASE EQUIVALENT)	21370060200120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 0.25 MG (BASE EQUIV)	21550080100120	Brand
HYCAMTIN	TOPOTECAN HCL CAP 1 MG (BASE EQUIV)	21550080100140	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 0.5 MG (BASE EQUIVALENT)	21533570100310	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE TAB 2 MG (BASE EQUIVALENT)	21533570100330	Brand
MEKINIST	TRAMETINIB DIMETHYL SULFOXIDE FOR SOLN 0.05 MG/ML (BASE EQ)	21533570102120	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 15-6.14 MG	21990002750320	Brand
LONSURF	TRIFLURIDINE-TIPIRACIL TAB 20-8.19 MG	21990002750330	Brand
TUKYSA	TUCATINIB TAB 50 MG	21170080000320	Brand
TUKYSA	TUCATINIB TAB 150 MG	21170080000340	Brand
VENCLEXTA STARTING PACK	VENETOCLAX TAB THERAPY STARTER PACK 10 & 50 & 100 MG	2147008000B720	Brand
VENCLEXTA	VENETOCLAX TAB 10 MG	21470080000320	Brand
VENCLEXTA	VENETOCLAX TAB 50 MG	21470080000340	Brand
VENCLEXTA	VENETOCLAX TAB 100 MG	21470080000360	Brand
ERIVEDGE	VISMODEGIB CAP 150 MG	21370070000120	Brand
ZOLINZA	VORINOSTAT CAP 100 MG	21531575000120	Brand
IBRANCE	PALBOCICLIB CAP 75 MG	21531060000120	Brand
IBRANCE	PALBOCICLIB CAP 100 MG	21531060000130	Brand
IBRANCE	PALBOCICLIB CAP 125 MG	21531060000140	Brand
IBRANCE	PALBOCICLIB TAB 75 MG	21531060000320	Brand
IBRANCE	PALBOCICLIB TAB 100 MG	21531060000330	Brand
IBRANCE	PALBOCICLIB TAB 125 MG	21531060000340	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 200 MG DAILY DOSE	2153107050B720	Brand
KISQALI	RIBOCICLIB SUCCINATE TAB PACK 400 MG DAILY DOSE (200 MG TAB)	2153107050B740	Brand

KISQALI	RIBOCICLIB SUCCINATE TAB PACK 600 MG DAILY DOSE (200 MG TAB)	2153107050B760	Brand
KISQALI FEMARA 200 DOSE	RIBOCICLIB 200 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B730	Brand
KISQALI FEMARA 400 DOSE	RIBOCICLIB 400 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B740	Brand
KISQALI FEMARA 600 DOSE	RIBOCICLIB 600 MG DOSE (200 MG TAB) & LETROZOLE 2.5 MG TBPK	2199000260B760	Brand
VERZENIO	ABEMACICLIB TAB 50 MG	21531010000305	Brand
VERZENIO	ABEMACICLIB TAB 100 MG	21531010000310	Brand
VERZENIO	ABEMACICLIB TAB 150 MG	21531010000315	Brand
VERZENIO	ABEMACICLIB TAB 200 MG	21531010000320	Brand
BOSULIF	BOSUTINIB CAP 50 MG	21531812000120	Brand
BOSULIF	BOSUTINIB CAP 100 MG	21531812000130	Brand
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 250 MG	21406010200320	Generic
ABIRATERONE ACETATE	ABIRATERONE ACETATE TAB 500 MG	21406010200330	Generic
ROZLYTREK	ENTRECTINIB PELLET PACK 50 MG	21533820003020	Brand

- 1 One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member* in one of the following:
 - United States Pharmacopeia Drug Information
 - American Hospital Formulary Service Drug Information
 - One article in a major peer- reviewed medical journal that recognizes the safety and efficacy of the requested drug in the member's specific condition

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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2. Revision History

Date	Notes
4/15/2024	Add generic pazopanib

Restricted Oral Oncology Drugs Split Fi		
	State Company and the September September (1994) and the September (199	

Prior Authorization Guideline

Guideline ID	GL-141303
Guideline Name	Restricted Oral Oncology Drugs Split Fill
Formulary	Quartz

Guideline Note:

Effective Date:	2/15/2024
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1. Criteria

Product Name: Bosulif, Cabometyx, Generic erlotinib, Generic everolimus, Generic gefitinib, Generic sorafenib, Generic sunitinib, Inlyta, Inrebic, Jakafi, Lynparza, Nubeqa, Piqray, Retevmo, Rubraca, Sprycel, Tagrisso, Tasigna, Tazverik, Votrient, Xalkori, Xtandi, Zejula, Zelboraf, Zykadia				
Approval Length	th 12 month(s)			
Therapy Stage	Therapy Stage Initial Authorization			
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans		ns		
Product Name	Gene	eric Name	GPI	Brand/Generic
PIQRAY 200MG DAILY DOSE		LISIB TAB THERAPY PACK 200 MG / DOSE	2153801000B720	Brand
PIQRAY 250MG DAILY DOSE	ALPELISIB TAB PACK 250 MG DAILY DOSE (200 MG & 50 MG TABS)		2153801000B725	Brand
PIQRAY 300MG DAILY DOSE		LISIB TAB PACK 300 MG DAILY DOSE 0 MG TAB)	2153801000B730	Brand

INLYTA	AXITINIB TAB 1 MG	21335013000320	Brand
INLYTA	AXITINIB TAB 5 MG	21335013000340	Brand
BOSULIF	BOSUTINIB TAB 100 MG	21531812000320	Brand
BOSULIF	BOSUTINIB TAB 400 MG	21531812000327	Brand
BOSULIF	BOSUTINIB TAB 500 MG	21531812000340	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 20 MG (BASE EQUIVALENT)	21533010100320	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 40 MG (BASE EQUIVALENT)	21533010100330	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 60 MG (BASE EQUIVALENT)	21533010100340	Brand
ZYKADIA	CERITINIB TAB 150 MG	21530514000330	Brand
XALKORI	CRIZOTINIB CAP 200 MG	21530517000120	Brand
XALKORI	CRIZOTINIB CAP 250 MG	21530517000125	Brand
NUBEQA	DAROLUTAMIDE TAB 300 MG	21402425000320	Brand
SPRYCEL	DASATINIB TAB 20 MG	21531820000320	Brand
SPRYCEL	DASATINIB TAB 50 MG	21531820000340	Brand
SPRYCEL	DASATINIB TAB 70 MG	21531820000350	Brand
SPRYCEL	DASATINIB TAB 80 MG	21531820000354	Brand
SPRYCEL	DASATINIB TAB 100 MG	21531820000360	Brand
SPRYCEL	DASATINIB TAB 140 MG	21531820000380	Brand
XTANDI	ENZALUTAMIDE CAP 40 MG	21402430000120	Brand
XTANDI	ENZALUTAMIDE TAB 40 MG	21402430000320	Brand
XTANDI	ENZALUTAMIDE TAB 80 MG	21402430000340	Brand
TARCEVA	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Generic
TARCEVA	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Generic
TARCEVA	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Generic
EVEROLIMUS	EVEROLIMUS TAB 2.5 MG	21532530000310	Generic

AFINITOR	EVEROLIMUS TAB 2.5 MG	21532530000310	Brand
EVEROLIMUS	EVEROLIMUS TAB 5 MG	21532530000320	Generic
AFINITOR	EVEROLIMUS TAB 5 MG	21532530000320	Brand
EVEROLIMUS	EVEROLIMUS TAB 7.5 MG	21532530000325	Generic
AFINITOR	EVEROLIMUS TAB 7.5 MG	21532530000325	Brand
EVEROLIMUS	EVEROLIMUS TAB 10 MG	21532530000330	Generic
AFINITOR	EVEROLIMUS TAB 10 MG	21532530000330	Brand
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Generic
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Generic
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Generic
INREBIC	FEDRATINIB HCL CAP 100 MG	21537520200120	Brand
IRESSA	GEFITINIB TAB 250 MG	21360030000320	Brand
GEFITINIB	GEFITINIB TAB 250 MG	21360030000320	Generic
TASIGNA	NILOTINIB HCL CAP 50 MG (BASE EQUIVALENT)	21531860200110	Brand
TASIGNA	NILOTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21531860200115	Brand
TASIGNA	NILOTINIB HCL CAP 200 MG (BASE EQUIVALENT)	21531860200125	Brand
ZEJULA	NIRAPARIB TOSYLATE CAP 100 MG (BASE EQUIVALENT)	21535550200120	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 100 MG (BASE EQUIVALENT)	21535550200320	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21535550200330	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 300 MG (BASE EQUIVALENT)	21535550200340	Brand
LYNPARZA	OLAPARIB TAB 100 MG	21535560000330	Brand
LYNPARZA	OLAPARIB TAB 150 MG	21535560000340	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 40 MG (BASE EQUIVALENT)	21360068200320	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 80 MG (BASE EQUIVALENT)	21360068200330	Brand

VOTRIENT	PAZOPANIB HCL TAB 200 MG (BASE EQUIV)	21533042100320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 200 MG (BASE EQUIVALENT)	21535570200320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 250 MG (BASE EQUIVALENT)	21535570200325	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 300 MG (BASE EQUIVALENT)	21535570200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 5 MG (BASE EQUIVALENT)	21537560200310	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	21537560200320	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 15 MG (BASE EQUIVALENT)	21537560200325	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 20 MG (BASE EQUIVALENT)	21537560200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 25 MG (BASE EQUIVALENT)	21537560200335	Brand
RETEVMO	SELPERCATINIB CAP 40 MG	21535779000120	Brand
RETEVMO	SELPERCATINIB CAP 80 MG	21535779000140	Brand
NEXAVAR	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Brand
SORAFENIB	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SORAFENIB TOSYLATE	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SUTENT	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Generic
SUTENT	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Generic
SUTENT	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Generic
SUTENT	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Generic
TAZVERIK	TAZEMETOSTAT HBR TAB 200 MG	21533675200320	Brand
ZELBORAF	VEMURAFENIB TAB 240 MG	21532080000320	Brand

BOSULIF	BOSUTINIB CAP 50 MG	21531812000120	Brand
BOSULIF	BOSUTINIB CAP 100 MG	21531812000130	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.

Product Name: Bosulif, Cabometyx, Generic erlotinib, Generic everolimus, Generic gefitinib, Generic sorafenib, Generic sunitinib, Inlyta, Inrebic, Jakafi, Lynparza, Nubeqa, Piqray, Retevmo, Rubraca, Sprycel, Sutent, Tagrisso, Tasigna, Tarceva, Tazverik, Votrient, Xalkori, Xtandi, Zejula, Zelboraf, Zykadia

Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - ALL Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
PIQRAY 200MG DAILY DOSE	ALPELISIB TAB THERAPY PACK 200 MG DAILY DOSE	2153801000B720	Brand

PIQRAY 250MG DAILY DOSE	ALPELISIB TAB PACK 250 MG DAILY DOSE (200 MG & 50 MG TABS)	2153801000B725	Brand
PIQRAY 300MG DAILY DOSE	ALPELISIB TAB PACK 300 MG DAILY DOSE (2X150 MG TAB)	2153801000B730	Brand
INLYTA	AXITINIB TAB 1 MG	21335013000320	Brand
INLYTA	AXITINIB TAB 5 MG	21335013000340	Brand
BOSULIF	BOSUTINIB TAB 100 MG	21531812000320	Brand
BOSULIF	BOSUTINIB TAB 400 MG	21531812000327	Brand
BOSULIF	BOSUTINIB TAB 500 MG	21531812000340	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 20 MG (BASE EQUIVALENT)	21533010100320	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 40 MG (BASE EQUIVALENT)	21533010100330	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 60 MG (BASE EQUIVALENT)	21533010100340	Brand
ZYKADIA	CERITINIB TAB 150 MG	21530514000330	Brand
XALKORI	CRIZOTINIB CAP 200 MG	21530517000120	Brand
XALKORI	CRIZOTINIB CAP 250 MG	21530517000125	Brand
NUBEQA	DAROLUTAMIDE TAB 300 MG	21402425000320	Brand
SPRYCEL	DASATINIB TAB 20 MG	21531820000320	Brand
SPRYCEL	DASATINIB TAB 50 MG	21531820000340	Brand
SPRYCEL	DASATINIB TAB 70 MG	21531820000350	Brand
SPRYCEL	DASATINIB TAB 80 MG	21531820000354	Brand
SPRYCEL	DASATINIB TAB 100 MG	21531820000360	Brand
SPRYCEL	DASATINIB TAB 140 MG	21531820000380	Brand
XTANDI	ENZALUTAMIDE CAP 40 MG	21402430000120	Brand
XTANDI	ENZALUTAMIDE TAB 40 MG	21402430000320	Brand
XTANDI	ENZALUTAMIDE TAB 80 MG	21402430000340	Brand
TARCEVA	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Generic
TARCEVA	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Generic
TARCEVA	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Brand

ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Generic
EVEROLIMUS	EVEROLIMUS TAB 2.5 MG	21532530000310	Generic
AFINITOR	EVEROLIMUS TAB 2.5 MG	21532530000310	Brand
EVEROLIMUS	EVEROLIMUS TAB 5 MG	21532530000320	Generic
AFINITOR	EVEROLIMUS TAB 5 MG	21532530000320	Brand
EVEROLIMUS	EVEROLIMUS TAB 7.5 MG	21532530000325	Generic
AFINITOR	EVEROLIMUS TAB 7.5 MG	21532530000325	Brand
EVEROLIMUS	EVEROLIMUS TAB 10 MG	21532530000330	Generic
AFINITOR	EVEROLIMUS TAB 10 MG	21532530000330	Brand
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Generic
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Generic
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Generic
INREBIC	FEDRATINIB HCL CAP 100 MG	21537520200120	Brand
IRESSA	GEFITINIB TAB 250 MG	21360030000320	Brand
GEFITINIB	GEFITINIB TAB 250 MG	21360030000320	Generic
TASIGNA	NILOTINIB HCL CAP 50 MG (BASE EQUIVALENT)	21531860200110	Brand
TASIGNA	NILOTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21531860200115	Brand
TASIGNA	NILOTINIB HCL CAP 200 MG (BASE EQUIVALENT)	21531860200125	Brand
ZEJULA	NIRAPARIB TOSYLATE CAP 100 MG (BASE EQUIVALENT)	21535550200120	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 100 MG (BASE EQUIVALENT)	21535550200320	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21535550200330	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 300 MG (BASE EQUIVALENT)	21535550200340	Brand
LYNPARZA	OLAPARIB TAB 100 MG	21535560000330	Brand
LYNPARZA	OLAPARIB TAB 150 MG	21535560000340	Brand

TAGRISSO	OSIMERTINIB MESYLATE TAB 40 MG (BASE EQUIVALENT)	21360068200320	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 80 MG (BASE EQUIVALENT)	21360068200330	Brand
VOTRIENT	PAZOPANIB HCL TAB 200 MG (BASE EQUIV)	21533042100320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 200 MG (BASE EQUIVALENT)	21535570200320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 250 MG (BASE EQUIVALENT)	21535570200325	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 300 MG (BASE EQUIVALENT)	21535570200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 5 MG (BASE EQUIVALENT)	21537560200310	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	21537560200320	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 15 MG (BASE EQUIVALENT)	21537560200325	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 20 MG (BASE EQUIVALENT)	21537560200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 25 MG (BASE EQUIVALENT)	21537560200335	Brand
RETEVMO	SELPERCATINIB CAP 40 MG	21535779000120	Brand
RETEVMO	SELPERCATINIB CAP 80 MG	21535779000140	Brand
NEXAVAR	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Brand
SORAFENIB	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SORAFENIB TOSYLATE	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SUTENT	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Generic
SUTENT	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Generic
SUTENT	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Generic
SUTENT	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Brand

SUNITINIB MALATE	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Generic
TAZVERIK	TAZEMETOSTAT HBR TAB 200 MG	21533675200320	Brand
ZELBORAF	VEMURAFENIB TAB 240 MG	21532080000320	Brand
BOSULIF	BOSUTINIB CAP 50 MG	21531812000120	Brand
BOSULIF	BOSUTINIB CAP 100 MG	21531812000130	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

AND

- **2** Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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Product Name: Bosulif, Cabometyx, Generic erlotinib, Generic everolimus, Generic gefitinib, Generic sorafenib, Generic sunitinib, Inlyta, Inrebic, Jakafi, Lynparza, Nubeqa, Piqray, Retevmo, Rubraca, Sprycel, Tagrisso, Tasigna, Tarceva, Tazverik, Votrient, Xalkori, Xtandi, Zejula, Zelboraf, Zykadia

Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - MN Plans

Product Name	Generic Name	GPI	Brand/Generic
PIQRAY 200MG DAILY DOSE	ALPELISIB TAB THERAPY PACK 200 MG DAILY DOSE	2153801000B720	Brand
PIQRAY 250MG DAILY DOSE	ALPELISIB TAB PACK 250 MG DAILY DOSE (200 MG & 50 MG TABS)	2153801000B725	Brand
PIQRAY 300MG DAILY DOSE	ALPELISIB TAB PACK 300 MG DAILY DOSE (2X150 MG TAB)	2153801000B730	Brand
INLYTA	AXITINIB TAB 1 MG	21335013000320	Brand
INLYTA	AXITINIB TAB 5 MG	21335013000340	Brand
BOSULIF	BOSUTINIB TAB 100 MG	21531812000320	Brand
BOSULIF	BOSUTINIB TAB 400 MG	21531812000327	Brand
BOSULIF	BOSUTINIB TAB 500 MG	21531812000340	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 20 MG (BASE EQUIVALENT)	21533010100320	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 40 MG (BASE EQUIVALENT)	21533010100330	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 60 MG (BASE EQUIVALENT)	21533010100340	Brand
ZYKADIA	CERITINIB TAB 150 MG	21530514000330	Brand
XALKORI	CRIZOTINIB CAP 200 MG	21530517000120	Brand
XALKORI	CRIZOTINIB CAP 250 MG	21530517000125	Brand
NUBEQA	DAROLUTAMIDE TAB 300 MG	21402425000320	Brand
SPRYCEL	DASATINIB TAB 20 MG	21531820000320	Brand
SPRYCEL	DASATINIB TAB 50 MG	21531820000340	Brand
SPRYCEL	DASATINIB TAB 70 MG	21531820000350	Brand
SPRYCEL	DASATINIB TAB 80 MG	21531820000354	Brand
SPRYCEL	DASATINIB TAB 100 MG	21531820000360	Brand
SPRYCEL	DASATINIB TAB 140 MG	21531820000380	Brand
XTANDI	ENZALUTAMIDE CAP 40 MG	21402430000120	Brand
XTANDI	ENZALUTAMIDE TAB 40 MG	21402430000320	Brand
XTANDI	ENZALUTAMIDE TAB 80 MG	21402430000340	Brand
TARCEVA	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Generic
TARCEVA	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Brand

ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Generic
TARCEVA	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Generic
EVEROLIMUS	EVEROLIMUS TAB 2.5 MG	21532530000310	Generic
AFINITOR	EVEROLIMUS TAB 2.5 MG	21532530000310	Brand
EVEROLIMUS	EVEROLIMUS TAB 5 MG	21532530000320	Generic
AFINITOR	EVEROLIMUS TAB 5 MG	21532530000320	Brand
EVEROLIMUS	EVEROLIMUS TAB 7.5 MG	21532530000325	Generic
AFINITOR	EVEROLIMUS TAB 7.5 MG	21532530000325	Brand
EVEROLIMUS	EVEROLIMUS TAB 10 MG	21532530000330	Generic
AFINITOR	EVEROLIMUS TAB 10 MG	21532530000330	Brand
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Generic
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Generic
AFINITOR DISPERZ	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Brand
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Generic
INREBIC	FEDRATINIB HCL CAP 100 MG	21537520200120	Brand
IRESSA	GEFITINIB TAB 250 MG	21360030000320	Brand
GEFITINIB	GEFITINIB TAB 250 MG	21360030000320	Generic
TASIGNA	NILOTINIB HCL CAP 50 MG (BASE EQUIVALENT)	21531860200110	Brand
TASIGNA	NILOTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21531860200115	Brand
TASIGNA	NILOTINIB HCL CAP 200 MG (BASE EQUIVALENT)	21531860200125	Brand
ZEJULA	NIRAPARIB TOSYLATE CAP 100 MG (BASE EQUIVALENT)	21535550200120	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 100 MG (BASE EQUIVALENT)	21535550200320	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21535550200330	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 300 MG (BASE EQUIVALENT)	21535550200340	Brand

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LYNPARZA	OLAPARIB TAB 100 MG	21535560000330	Brand
LYNPARZA	OLAPARIB TAB 150 MG	21535560000340	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 40 MG (BASE EQUIVALENT)	21360068200320	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 80 MG (BASE EQUIVALENT)	21360068200330	Brand
VOTRIENT	PAZOPANIB HCL TAB 200 MG (BASE EQUIV)	21533042100320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 200 MG (BASE EQUIVALENT)	21535570200320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 250 MG (BASE EQUIVALENT)	21535570200325	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 300 MG (BASE EQUIVALENT)	21535570200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 5 MG (BASE EQUIVALENT)	21537560200310	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	21537560200320	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 15 MG (BASE EQUIVALENT)	21537560200325	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 20 MG (BASE EQUIVALENT)	21537560200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 25 MG (BASE EQUIVALENT)	21537560200335	Brand
RETEVMO	SELPERCATINIB CAP 40 MG	21535779000120	Brand
RETEVMO	SELPERCATINIB CAP 80 MG	21535779000140	Brand
NEXAVAR	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Brand
SORAFENIB	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SORAFENIB TOSYLATE	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SUTENT	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Generic
SUTENT	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Generic
SUTENT	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Generic

SUTENT	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Brand
SUNITINIB MALATE	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Generic
TAZVERIK	TAZEMETOSTAT HBR TAB 200 MG	21533675200320	Brand
ZELBORAF	VEMURAFENIB TAB 240 MG	21532080000320	Brand
BOSULIF	BOSUTINIB CAP 50 MG	21531812000120	Brand
BOSULIF	BOSUTINIB CAP 100 MG	21531812000130	Brand

- **1** One of the following:
- **1.1** The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member in one of the following:*
 - United States Pharmacopeia Drug Information
 - American Hospital Formulary Service Drug Information
 - One article in a major peer-reviewed medical journal that recognizes the safety and efficacy of the requested drug in the member's specific condition

AND

- 2 Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist

other specialist in the treatment of malignancy

Notes

*Includes any relevant genetic testing, mutations, etc.

Product Name: Bosulif, Cabometyx, Generic erlotinib, Generic everolimus, Generic gefitinib, Generic sorafenib, Generic sunitinib, Inlyta, Inrebic, Jakafi, Lynparza, Nubeqa, Piqray, Retevmo, Rubraca, Sprycel, Tagrisso, Tasigna, Tarceva, Tazverik, Votrient, Xalkori, Xtandi, Zejula, Zelboraf, Zykadia

Approval Length 12 month(s)

Therapy Stage Reauthorization

Guideline Type Prior Authorization - MN Plans

Product Name	Generic Name	GPI	Brand/Generic
PIQRAY 200MG DAILY DOSE	ALPELISIB TAB THERAPY PACK 200 MG DAILY DOSE	2153801000B720	Brand
PIQRAY 250MG DAILY DOSE	ALPELISIB TAB PACK 250 MG DAILY DOSE (200 MG & 50 MG TABS)	2153801000B725	Brand
PIQRAY 300MG DAILY DOSE	ALPELISIB TAB PACK 300 MG DAILY DOSE (2X150 MG TAB)	2153801000B730	Brand
INLYTA	AXITINIB TAB 1 MG	21335013000320	Brand
INLYTA	AXITINIB TAB 5 MG	21335013000340	Brand
BOSULIF	BOSUTINIB TAB 100 MG	21531812000320	Brand
BOSULIF	BOSUTINIB TAB 400 MG	21531812000327	Brand
BOSULIF	BOSUTINIB TAB 500 MG	21531812000340	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 20 MG (BASE EQUIVALENT)	21533010100320	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 40 MG (BASE EQUIVALENT)	21533010100330	Brand
CABOMETYX	CABOZANTINIB S-MALATE TAB 60 MG (BASE EQUIVALENT)	21533010100340	Brand
ZYKADIA	CERITINIB TAB 150 MG	21530514000330	Brand
XALKORI	CRIZOTINIB CAP 200 MG	21530517000120	Brand
XALKORI	CRIZOTINIB CAP 250 MG	21530517000125	Brand
NUBEQA	DAROLUTAMIDE TAB 300 MG	21402425000320	Brand
SPRYCEL	DASATINIB TAB 20 MG	21531820000320	Brand
SPRYCEL	DASATINIB TAB 50 MG	21531820000340	Brand
SPRYCEL	DASATINIB TAB 70 MG	21531820000350	Brand
SPRYCEL	DASATINIB TAB 80 MG	21531820000354	Brand

SPRYCEL	DASATINIB TAB 100 MG	21531820000360	Brand
SPRYCEL	DASATINIB TAB 140 MG	21531820000380	Brand
XTANDI	ENZALUTAMIDE CAP 40 MG	21402430000120	Brand
XTANDI	ENZALUTAMIDE TAB 40 MG	21402430000320	Brand
XTANDI	ENZALUTAMIDE TAB 80 MG	21402430000340	Brand
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 25 MG (BASE EQUIVALENT)	21360025100320	Generic
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 100 MG (BASE EQUIVALENT)	21360025100330	Generic
ERLOTINIB HYDROCHLORIDE	ERLOTINIB HCL TAB 150 MG (BASE EQUIVALENT)	21360025100360	Generic
EVEROLIMUS	EVEROLIMUS TAB 2.5 MG	21532530000310	Generic
EVEROLIMUS	EVEROLIMUS TAB 5 MG	21532530000320	Generic
EVEROLIMUS	EVEROLIMUS TAB 7.5 MG	21532530000325	Generic
EVEROLIMUS	EVEROLIMUS TAB 10 MG	21532530000330	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 2 MG	21532530007310	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 3 MG	21532530007320	Generic
EVEROLIMUS	EVEROLIMUS TAB FOR ORAL SUSP 5 MG	21532530007340	Generic
INREBIC	FEDRATINIB HCL CAP 100 MG	21537520200120	Brand
GEFITINIB	GEFITINIB TAB 250 MG	21360030000320	Generic
TASIGNA	NILOTINIB HCL CAP 50 MG (BASE EQUIVALENT)	21531860200110	Brand
TASIGNA	NILOTINIB HCL CAP 150 MG (BASE EQUIVALENT)	21531860200115	Brand
TASIGNA	NILOTINIB HCL CAP 200 MG (BASE EQUIVALENT)	21531860200125	Brand
ZEJULA	NIRAPARIB TOSYLATE CAP 100 MG (BASE EQUIVALENT)	21535550200120	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 100 MG (BASE EQUIVALENT)	21535550200320	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21535550200330	Brand
ZEJULA	NIRAPARIB TOSYLATE TAB 300 MG (BASE EQUIVALENT)	21535550200340	Brand
LYNPARZA	OLAPARIB TAB 100 MG	21535560000330	Brand
LYNPARZA	OLAPARIB TAB 150 MG	21535560000340	Brand
TAGRISSO	OSIMERTINIB MESYLATE TAB 40 MG (BASE EQUIVALENT)	21360068200320	Brand

TAGRISSO	OSIMERTINIB MESYLATE TAB 80 MG (BASE EQUIVALENT)	21360068200330	Brand
VOTRIENT	PAZOPANIB HCL TAB 200 MG (BASE EQUIV)	21533042100320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 200 MG (BASE EQUIVALENT)	21535570200320	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 250 MG (BASE EQUIVALENT)	21535570200325	Brand
RUBRACA	RUCAPARIB CAMSYLATE TAB 300 MG (BASE EQUIVALENT)	21535570200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 5 MG (BASE EQUIVALENT)	21537560200310	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 10 MG (BASE EQUIVALENT)	21537560200320	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 15 MG (BASE EQUIVALENT)	21537560200325	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 20 MG (BASE EQUIVALENT)	21537560200330	Brand
JAKAFI	RUXOLITINIB PHOSPHATE TAB 25 MG (BASE EQUIVALENT)	21537560200335	Brand
RETEVMO	SELPERCATINIB CAP 40 MG	21535779000120	Brand
RETEVMO	SELPERCATINIB CAP 80 MG	21535779000140	Brand
SORAFENIB	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SORAFENIB TOSYLATE	SORAFENIB TOSYLATE TAB 200 MG (BASE EQUIVALENT)	21533060400320	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 12.5 MG (BASE EQUIVALENT)	21533070300120	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 25 MG (BASE EQUIVALENT)	21533070300130	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 37.5 MG (BASE EQUIVALENT)	21533070300135	Generic
SUNITINIB MALATE	SUNITINIB MALATE CAP 50 MG (BASE EQUIVALENT)	21533070300140	Generic
TAZVERIK	TAZEMETOSTAT HBR TAB 200 MG	21533675200320	Brand
ZELBORAF	VEMURAFENIB TAB 240 MG	21532080000320	Brand
BOSULIF	BOSUTINIB CAP 50 MG	21531812000120	Brand
BOSULIF	BOSUTINIB CAP 100 MG	21531812000130	Brand

1 - One of the following:

1.1 The requested drug is being used alone or in a combination regimen that is FDA-labeled for the treatment of the specific condition the member presents with*

OR

1.2 The requested drug is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) in the specific condition of the member*

OR

- **1.3** The requested drug is being used alone or in a combination regimen that is recommended for use in the specific condition of the member in one of the following:*
 - United States Pharmacopeia Drug Information
 - American Hospital Formulary Service Drug Information
 - One article in a major peer-reviewed medical journal that recognizes the safety and efficacy of the requested drug in the member's specific condition

AND

- 2 Prescribed by or in consultation with one of the following:
 - oncologist
 - hematologist
 - · other specialist in the treatment of malignancy

Notes	*Includes any relevant genetic testing, mutations, etc.
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2. Revision History

Date	Notes
2/14/2024	Update program – Bosulif capsules added criteria

Restricted Paroxetine				
The state of the s				

Guideline ID	GL-131421
Guideline Name	Restricted Paroxetine
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Paroxetine mesylate		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
PAROXETINE	PAROXETINE MESYLATE CAP 7.5 MG (BASE EQUIV)	62226060300110	Generic

Approval Criteria

1 - Diagnosis of vasomotor symptoms due to menopause

AND

2 - Failure of a trial of generic paroxetine (Paxil generic) at an equivalent dose

AND

3 - The prescriber provides an evidence-based clinical rationale for why the requested formulation would provide different results.

Product Name: Paroxetine mesylate			
Approval Length 12 month(s)			
Therapy Stage	Therapy Stage Reauthorization		
Guideline Type Prior Authorization-IL and MN Plans Only			

Product Name	Generic Name	GPI	Brand/Generic
PAROXETINE	PAROXETINE MESYLATE CAP 7.5 MG (BASE EQUIV)	62226060300110	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Paroxetine mesylate			
Approval Length 12/31/2039			
Guideline Type Prior Authorization-All plans except IL and MN			

Product Name	Generic Name	GPI	Brand/Generic
PAROXETINE	PAROXETINE MESYLATE CAP 7.5 MG (BASE EQUIV)	62226060300110	Generic

Approval Criteria

1 - Diagnosis of vasomotor symptoms due to menopause

AND

2 - Failure of a trial of generic paroxetine (Paxil generic) at an equivalent dose

AND

3 - The prescriber provides an evidence-based clinical rationale for why the requested formulation would provide different results.

2. Revision History

Date	Notes
10/16/2023	New program

Restricted	Restricted Phosphate Binders				
The behalf longs content to displayed. The flowing has been record, on					

Guideline ID	GL-131422
Guideline Name	Restricted Phosphate Binders
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Velphoro		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
VELPHORO	SUCROFERRIC OXYHYDROXIDE CHEW TAB 500 MG	52800080100520	Brand

Approval Criteria

1 - Diagnosis of chronic kidney disease (CKD) requiring dialysis

AND

2 - Trial and failure, contraindication, or intolerance to BOTH a sevelamer product (e.g. Renagel, Renvela) and lanthanum (Fosrenol)

Product Name: Velphoro		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
VELPHORO	SUCROFERRIC OXYHYDROXIDE CHEW TAB 500 MG	52800080100520	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Velphoro	
Approval Length	12/31/2039
Guideline Type	Prior Authorization-All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
VELPHORO	SUCROFERRIC OXYHYDROXIDE CHEW TAB 500 MG	52800080100520	Brand

Approval Criteria

1 - Diagnosis of chronic kidney disease (CKD) requiring dialysis

AND

2 - Trial and failure, contraindication, or intolerance to BOTH a sevelamer product (e.g. Renagel, Renvela) and lanthanum (Fosrenol)

2. Revision History

Date	Notes
10/9/2023	New Program

Restricted Progesterone		
The black longs connect to display. The fire may have been record, varieties, and details labely that the loss points in the connectific and long	in .	

Guideline ID	GL-144934
Guideline Name	Restricted Progesterone
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	1/20/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Crinone, Endometrin, progesterone injection				
Diagnosis		Pregnancy		
Approval Length		12 month(s)		
Guideline Type		Prior Authorization - IL and MN Plans Only		
Product Name	Gene	ric Name	GPI	Brand/Generic
CRINONE	PROG	ESTERONE VAGINAL GEL 4%	55370060004010	Brand
CRINONE	PROG	ESTERONE VAGINAL GEL 8%	55370060004020	Brand
ENDOMETRIN	PROG	ESTERONE VAGINAL INSERT 100 MG	55370060009910	Brand
PROGESTERONE	PROG	ESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic

- 1 One of the following:
- **1.1** For members in the 1st trimester of pregnancy, ALL of the following:
 - Submission of medical records (e.g., chart notes) documenting member is pregnant
 - Prescriber determines that progesterone is to maintain pregnancy

OR

- **1.2** For members in the 2nd trimester of pregnancy, ALL of the following:
 - Submission of medical records (e.g., chart notes) documenting a singleton pregnancy
 - Submission of medical records (e.g., chart notes) documenting member has a history of preterm birth

OR

1.3 Person is currently pregnant and continuation of prior therapy with progesterone, verified by paid claims, medical records (e.g. chart notes), or provider attestation

Product Name: Crinone, Endometrin, progesterone injection	
Diagnosis Pregnancy	
Approval Length	1st trimester use = 4 months. 2nd trimester use = 6 months.
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
CRINONE	PROGESTERONE VAGINAL GEL 4%	55370060004010	Brand
CRINONE	PROGESTERONE VAGINAL GEL 8%	55370060004020	Brand
ENDOMETRIN	PROGESTERONE VAGINAL INSERT 100 MG	55370060009910	Brand
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic

Approval Criteria

- 1 One of the following:
 - **1.1** For members in the 1st trimester of pregnancy, ALL of the following:
 - Submission of medical records (e.g., chart notes) documenting member is pregnant
 - Prescriber determines that progesterone is to maintain pregnancy

OR

- **1.2** For members in the 2nd trimester of pregnancy, ALL of the following:
 - Submission of medical records (e.g., chart notes) documenting a singleton pregnancy
 - Submission of medical records (e.g., chart notes) documenting member has a history of preterm birth

OR

1.3 Person is currently pregnant and continuation of prior therapy with progesterone, verified by paid claims, medical records (e.g. chart notes), or provider attestation

Product Name: Crinone, Endometrin, progesterone injection	
Diagnosis	Infertility
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL Plan

Product Name	Generic Name	GPI	Brand/Generic
CRINONE	PROGESTERONE VAGINAL GEL 4%	55370060004010	Brand
CRINONE	PROGESTERONE VAGINAL GEL 8%	55370060004020	Brand
ENDOMETRIN	PROGESTERONE VAGINAL INSERT 100 MG	55370060009910	Brand
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic

Approval Criteria

1 - Quartz plan issued in the state of Illinois

AND

2 - Provider attests patient has infertility coverage as outlined in Illinois Insurance Code 215 ILCS 5/356m

ILCS 5/356m	
Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Name: Crinone, Endometrin, progesterone injection		
Diagnosis	Infertility	
Approval Length	12 month(s)	
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL Plan		

Product Name	Generic Name	GPI	Brand/Generic
CRINONE	PROGESTERONE VAGINAL GEL 4%	55370060004010	Brand
CRINONE	PROGESTERONE VAGINAL GEL 8%	55370060004020	Brand
ENDOMETRIN	PROGESTERONE VAGINAL INSERT 100 MG	55370060009910	Brand
PROGESTERONE	PROGESTERONE IM IN OIL 50 MG/ML	26000040001705	Generic

Approval Criteria

1 - Quartz plan issued in the state of Illinois

AND

2 - Provider attests patient has infertility coverage as outlined in Illinois Insurance Code 215 ILCS 5/356m

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

2. Revision History

Date	Notes
3/27/2024	Guideline update

R	estricted Tacrolimus Formul	ations
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Guideline ID	GL-129869
Guideline Name	Restricted Tacrolimus Formulations
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Prograf granule packets		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
PROGRAF	TACROLIMUS PACKET FOR SUSP 0.2 MG	99404080003010	Brand
PROGRAF	TACROLIMUS PACKET FOR SUSP 1 MG	99404080003030	Brand

Approval Criteria

1 - Member has swallowing impairment or other medical condition that prevents use of solid dose forms

AND

- 2 One of the following:
- **2.1** Trial and failure, contraindication, or intolerance to an adequate trial of an alternative (e.g. sirolimus, cyclosporine)

OR

2.2 Submission of medical records (e.g., chart notes) documenting evidence-based rationale for why the alternatives would not be medically appropriate for the member's condition

Product Name: Astagraf XL, Envarsus XR	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 0.5 MG	99404080007005	Brand
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 1 MG	99404080007010	Brand
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 5 MG	99404080007020	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 0.75 MG	99404080007510	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 1 MG	99404080007515	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 4 MG	99404080007520	Brand

Approval Criteria

1 - Trial and failure (documented inability to achieve goal trough drug levels despite

appropriate dose adjustment and teaching/adherence interventions from a pharmacist and other health care providers) or intolerance of immediate release tacrolimus

Product Name: Prograf granule packets, Astagraf XL, Envarsus XR		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
PROGRAF	TACROLIMUS PACKET FOR SUSP 0.2 MG	99404080003010	Brand
PROGRAF	TACROLIMUS PACKET FOR SUSP 1 MG	99404080003030	Brand
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 0.5 MG	99404080007005	Brand
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 1 MG	99404080007010	Brand
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 5 MG	99404080007020	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 0.75 MG	99404080007510	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 1 MG	99404080007515	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 4 MG	99404080007520	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Prograf granule packets		
Approval Length 12/31/2039		
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
PROGRAF	TACROLIMUS PACKET FOR SUSP 0.2 MG	99404080003010	Brand
PROGRAF	TACROLIMUS PACKET FOR SUSP 1 MG	99404080003030	Brand

1 - Member has swallowing impairment or other medical condition that prevents use of solid dose forms

AND

- 2 One of the following:
- **2.1** Trial and failure, contraindication, or intolerance to an adequate trial of an alternative (e.g. sirolimus, cyclosporine)

OR

2.2 Submission of medical records (e.g., chart notes) documenting evidence-based rationale for why the alternatives would not be medically appropriate for the member's condition

Product Name: Astagraf XL, Envarsus XR			
Approval Length 12/31/2039			
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 0.5 MG	99404080007005	Brand
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 1 MG	99404080007010	Brand
ASTAGRAF XL	TACROLIMUS CAP ER 24HR 5 MG	99404080007020	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 0.75 MG	99404080007510	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 1 MG	99404080007515	Brand
ENVARSUS XR	TACROLIMUS TAB ER 24HR 4 MG	99404080007520	Brand

Approval Criteria

1 - Trial and failure (documented inability to achieve goal trough drug levels despite appropriate dose adjustment and teaching/adherence interventions from a pharmacist and other health care providers) or intolerance of immediate release tacrolimus

2. Revision History

Date	Notes
10/12/2023	2024 New Implementation

Reti	Retinoid Products						
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Guideline ID	GL-131450
Guideline Name	Retinoid Products
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Tretinoin, OTC adapalene, Brand Avita					
Approval Length		12/31/2039			
Guideline Type		Prior Authorization - All plans except IL and MN Plans			
Product Name	Generic Name		GPI	Brand/Generic	
DIFFERIN	ADAPALENE GEL 0.1%		90050003004010	Brand	
CVS ADAPALENE	ADAPALENE GEL 0.1%		90050003004010	Generic	
ADAPALENE	ADAPALENE GEL 0.1%		90050003004010	Generic	
ADAPALENE TREATMENT	ADAPALENE GEL 0.1%		90050003004010	Generic	
ALTRENO	TRETINOIN	LOTION 0.05%	90050030004130	Brand	
TRETINOIN	TRETINOIN	CREAM 0.025%	90050030003703	Generic	

TRETINOIN	TRETINOIN CREAM 0.05% 90050030003705 Generic		Generic
TRETINOIN	TRETINOIN CREAM 0.1% 90050030003710 Generic		Generic
TRETINOIN	TRETINOIN GEL 0.01% 90050030004005 Generic		Generic
TRETINOIN	N TRETINOIN GEL 0.025% 90050030004010 Gener		Generic
TRETINOIN	TRETINOIN GEL 0.05%	90050030004015 Generic	
AVITA	TRETINOIN CREAM 0.025%	90050030003703	Brand

1 - Diagnosis of acne or rosacea

Product Name: Tretinoin, OTC adapalene, Brand Avita		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
DIFFERIN	ADAPALENE GEL 0.1%	90050003004010	Brand
CVS ADAPALENE	ADAPALENE GEL 0.1%	90050003004010	Generic
ADAPALENE	ADAPALENE GEL 0.1%	90050003004010	Generic
ADAPALENE TREATMENT	ADAPALENE GEL 0.1%	90050003004010	Generic
ALTRENO	TRETINOIN LOTION 0.05%	90050030004130	Brand
TRETINOIN	TRETINOIN CREAM 0.025%	90050030003703	Generic
TRETINOIN	TRETINOIN CREAM 0.05%	90050030003705	Generic
TRETINOIN	TRETINOIN CREAM 0.1%	90050030003710	Generic
TRETINOIN	TRETINOIN GEL 0.01%	90050030004005	Generic
TRETINOIN	TRETINOIN GEL 0.025%	90050030004010	Generic
TRETINOIN	TRETINOIN GEL 0.05%	90050030004015	Generic
AVITA	TRETINOIN CREAM 0.025%	90050030003703	Brand

Approval Criteria

1 - Diagnosis of acne or rosacea

Product Name: Aklief		
Approval Length 12/31/2039		
Guideline Type	Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
AKLIEF	TRIFAROTENE CREAM 0.005%	90050035003720	Brand

Approval Criteria

1 - Diagnosis of acne or rosacea

AND

- 2 Trial and failure, contraindication, or intolerance to BOTH of the following:
 - preferred tretinoin
 - adapalene agent

Product Name: Aklief		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
AKLIEF	TRIFAROTENE CREAM 0.005%	90050035003720	Brand

Approval Criteria

1 - Diagnosis of acne or rosacea

AND

- **2** Trial and failure, contraindication, or intolerance to BOTH of the following:
 - preferred tretinoin
 - adapalene agent

Product Name: Prescription adapalene products		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
ADAPALENE	ADAPALENE SOLN 0.1%	90050003002010	Brand
ADAPALENE	ADAPALENE CREAM 0.1%	90050003003710	Generic
ADAPALENE	ADAPALENE GEL 0.3%	90050003004030	Generic
ADAPALENE PUMP	ADAPALENE GEL 0.3%	90050003004030	Generic
ADAPALENE	ADAPALENE PADS 0.1%	90050003004310	Brand

Approval Criteria

1 - Diagnosis of acne or rosacea

AND

2 - Trial and failure, contraindication, or intolerance to adapalene 0.1% gel

Product Name: Prescription adapalene products		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ADAPALENE	ADAPALENE SOLN 0.1%	90050003002010	Brand
ADAPALENE	ADAPALENE CREAM 0.1%	90050003003710	Generic
ADAPALENE	ADAPALENE GEL 0.3%	90050003004030	Generic
ADAPALENE PUMP	ADAPALENE GEL 0.3%	90050003004030	Generic
ADAPALENE	ADAPALENE PADS 0.1%	90050003004310	Brand

1 - Diagnosis of acne or rosacea

AND

2 - Trial and failure, contraindication, or intolerance to adapalene 0.1% gel

Product Name: Tazarotene products	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TAZAROTENE	TAZAROTENE (ACNE) FOAM 0.1%	90050027003930	Brand
FABIOR	TAZAROTENE (ACNE) FOAM 0.1%	90050027003930	Brand
TAZAROTENE	TAZAROTENE CREAM 0.1%	90250070003730	Generic
TAZAROTENE	TAZAROTENE GEL 0.05%	90250070004020	Generic
TAZAROTENE	TAZAROTENE GEL 0.1%	90250070004030	Generic
TAZORAC	TAZAROTENE CREAM 0.05%	90250070003720	Brand

Approval Criteria

1 - Diagnosis of psoriasis

OR

- 2 Both of the following:
- 2.1 Diagnosis of acne or rosacea

AND

- **2.2** Trial and failure, contraindication, or intolerance to BOTH of the following:
 - preferred tretinoin
 - adapalene agent

Product Name: Tazarotene products		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TAZAROTENE	TAZAROTENE (ACNE) FOAM 0.1%	90050027003930	Brand
FABIOR	TAZAROTENE (ACNE) FOAM 0.1%	90050027003930	Brand
TAZAROTENE	TAZAROTENE CREAM 0.1%	90250070003730	Generic
TAZAROTENE	TAZAROTENE GEL 0.05%	90250070004020	Generic
TAZAROTENE	TAZAROTENE GEL 0.1%	90250070004030	Generic
TAZORAC	TAZAROTENE CREAM 0.05%	90250070003720	Brand

Approval Criteria

1 - Diagnosis of psoriasis

OR

- **2** Both of the following:
- 2.1 Diagnosis of acne or rosacea

AND

- 2.2 Trial and failure, contraindication, or intolerance to BOTH of the following:
 - preferred tretinoin
 - adapalene agent

Product Name: Duobrii	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
DUOBRII	HALOBETASOL PROPIONATE-TAZAROTENE LOTION 0.01-0.045%	90559902484120	Brand

Approval Criteria

1 - Diagnosis of psoriasis

AND

2 - Trial and failure, contraindication, or intolerance to one preferred high or super-high potency topical corticosteroid

Product Na	Product Name: Duobrii			
Approval L	ength	12 month(s)		
Therapy St	age	Initial Authorization		
Guideline Type Prior Authorization - IL a		Prior Authorization - IL and MN Plan	IS	
Product Name	Generic Na	me	GPI	Brand/Generic

DUOBRII	HALOBETASOL PROPIONATE-TAZAROTENE LOTION 0.01-0.045%	90559902484120	Brand
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1 - Diagnosis of psoriasis

AND

2 - Trial and failure, contraindication, or intolerance to one preferred high or super-high potency topical corticosteroid

Product Name: Clindamycin/tretinoin products			
Approval Length 12/31/2039			
Guideline Type	Prior Authorization - All plans excep	t IL and MN Plans	
Product Name	Generic Name	GPI	Brand/Generic
CLINDAMYCIN PHOSPHATE/TRETINOIN	CLINDAMYCIN PHOSPHATE-TRETINOIN GEL 1.2-0.025%	90059902654020	Generic

Approval Criteria

1 - Diagnosis of acne or rosacea

AND

2 - Trial and failure of concurrent use of the individual products (topical clindamycin and preferred tretinoin)

Product Name: Clindamycin/tretinoin products			
Approval Length	12 month(s)		
Therapy Stage	Initial Authorization		
Guideline Type	Prior Authorization - IL and MN Plans		
Product Name	ame Generic Name GPI Brand/Generic		

CLINDAMYCIN PHOSPHATE/TRETINOIN	CLINDAMYCIN PHOSPHATE-TRETINOIN GEL 1.2-0.025%	90059902654020	Generic
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1 - Diagnosis of acne or rosacea

AND

2 - Trial and failure of concurrent use of the individual products (topical clindamycin and preferred tretinoin)

Product Name: All Products Listed Above			
Approval Length	12 month(s)		
Therapy Stage	Reauthorization		
Guideline Type	Prior Authorization - IL and MN Plar	าร	
Product Name	Generic Name	GPI	Brand/Generic
ADAPALENE	ADAPALENE SOLN 0.1%	90050003002010	Brand
ADAPALENE	ADAPALENE CREAM 0.1%	90050003003710	Generic
DIFFERIN	ADAPALENE GEL 0.1%	90050003004010	Brand
CVS ADAPALENE	ADAPALENE GEL 0.1%	90050003004010	Generic
ADAPALENE	ADAPALENE GEL 0.1%	90050003004010	Generic
ADAPALENE TREATMENT	ADAPALENE GEL 0.1%	90050003004010	Generic
ADAPALENE	ADAPALENE GEL 0.3%	90050003004030	Generic
ADAPALENE PUMP	ADAPALENE GEL 0.3%	90050003004030	Generic
TAZAROTENE	TAZAROTENE (ACNE) FOAM 0.1%	90050027003930	Brand
FABIOR	TAZAROTENE (ACNE) FOAM 0.1%	90050027003930	Brand
TAZAROTENE	TAZAROTENE CREAM 0.1%	90250070003730	Generic
TAZAROTENE	TAZAROTENE GEL 0.05%	90250070004020	Generic
TAZAROTENE	TAZAROTENE GEL 0.1%	90250070004030	Generic
ALTRENO	TRETINOIN LOTION 0.05%	90050030004130	Brand
TRETINOIN	TRETINOIN CREAM 0.025%	90050030003703	Generic

TRETINOIN	TRETINOIN CREAM 0.05%	90050030003705	Generic
TRETINOIN	TRETINOIN CREAM 0.1%	90050030003710	Generic
TRETINOIN	TRETINOIN GEL 0.01%	90050030004005	Generic
TRETINOIN	TRETINOIN GEL 0.025%	90050030004010	Generic
TRETINOIN	TRETINOIN GEL 0.05%	90050030004015	Generic
AKLIEF	TRIFAROTENE CREAM 0.005%	90050035003720	Brand
CLINDAMYCIN PHOSPHATE/TRETINOIN	CLINDAMYCIN PHOSPHATE-TRETINOIN GEL 1.2-0.025%	90059902654020	Generic
DUOBRII	HALOBETASOL PROPIONATE- TAZAROTENE LOTION 0.01-0.045%	90559902484120	Brand
AVITA	TRETINOIN CREAM 0.025%	90050030003703	Brand
TAZORAC	TAZAROTENE CREAM 0.05%	90250070003720	Brand
ADAPALENE	ADAPALENE PAD 0.1% SWAB	90050003004310	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

2. Revision History

Date	Notes
10/31/2023	2024 New Implementation

Revcovi (elapegademase)
The State Surgeries of States, the state most, conduct and technical parameters are considered as the States of States and States of Sta

Guideline ID GL-129217	
Guideline Name	Revcovi (elapegademase)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Revcovi	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
REVCOVI	ELAPEGADEMASE-LVLR IM SOLN 2.4 MG/1.5ML (1.6 MG/ML)	30902030202020	Brand

Approval Criteria

1 - Both of the following:

1.1 Diagnosis of adenosine deaminase severe combined immune deficiency (ADA-SCID)

AND

1.2 Prescribed by, or in consultation with, an expert in the treatment of immune deficiencies

Product Name: Ro	Product Name: Revcovi			
Approval Length	12 month(s)			
Therapy Stage	Reauthorization	Reauthorization		
Guideline Type	Prior Authorization			
Product Gener Name	ic Name	GPI	Brand/Generic	

ELAPEGADEMASE-LVLR IM SOLN 2.4 MG/1.5ML (1.6

Approval Criteria

REVCOVI

1 - The prescriber provides recent clinical documentation from the past 6 months of a trough plasma ADA activity ≥ 30 mmol/hr/L and a trough erythrocyte dAXP level below 0.02 mmol/L

2. Revision History

MG/ML)

Date	Notes
8/9/2023	New program

Brand

30902030202020

ŀ	Rezurock (belumosudil mesylate)		
•	Thinking were shippy in "both on tentor most weak what out he fall in pass the enrolled reads."		

Guideline ID	ine ID GL-128187	
Guideline Name	Rezurock (belumosudil mesylate)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Rezurock*	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
REZUROCK	BELUMOSUDIL MESYLATE TAB 200 MG	99398510500320	Brand

Approval Criteria

1 - Diagnosis of chronic graft-versus-host disease (chronic GVHD)

AND

2 - Prescribed by or in consultation by a specialist with experience in the treatment of GVHD (e.g.hematologist, oncologist, immunologist, etc.)

AND

3 - The requested is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) for the treatment of chronic GVHD

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through reauthorization criteria

Product Name: Rezurock*		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
REZUROCK	BELUMOSUDIL MESYLATE TAB 200 MG	99398510500320	Brand

Approval Criteria

1 - Diagnosis of chronic graft-versus-host disease (chronic GVHD)

AND

2 - Prescribed by or in consultation by a specialist with experience in the treatment of GVHD (e.g.hematologist, oncologist, immunologist, etc.)

AND

3 - The requested is being used alone or in a combination regimen that has a class 1 or 2 recommendation for use from the National Comprehensive Cancer Network (NCCN) for the treatment of chronic GVHD

AND

4 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through
reauthorization criteria

Date	Notes
9/7/2023	New Program

Rinvoq (upadacitinib)			
(g) the best department of the tense to the contract of the co			

Prior Authorization Guideline

Guideline ID	GL-145362	
Guideline Name	Rinvoq (upadacitinib)	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Rinvoq				
Diagnosis		Psoriatic Arthritis (PsA)		
Approval Length		12/31/2039		
Guideline Type		Prior Authorization - All Plans except IL and MN Plans		
Product Name	Generic Name GPI Brand/Gene		Brand/Generic	
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG		66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG 66603072007530 Br		Brand	
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG 66603072007540 Brand		Brand	

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq	
Diagnosis Psoriatic Arthritis (PsA)	
Approval Length	12 month(s)

Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG		66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG		66603072007530	Brand
RINVOQ	UPADACITIN	IB TAB ER 24HR 45 MG	66603072007540	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- 3.1.1 Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - · actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq	
Diagnosis Moderate to Severely Active Rheumatoid Arthritis (RA)	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a minimum duration of a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)*
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Notes	* Absolute contraindications to methotrexate are pregnancy, nursing,	
	alcoholism, alcoholic liver disease or other chronic liver disease, immu	
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm	
	bocytopenia or significant anemia, or hypersensitivity to methotrexate.	

Product Name: Rinvoq	
Diagnosis Moderate to Severely Active Rheumatoid Arthritis (RA)	
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a minimum duration of a 3-month trial and failure, contraindication, or intolerance to ONE of the following:
 - methotrexate (MTX)*
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Notes	* Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Rinvoq		
Diagnosis	Ankylosing Spondylitis (AS), Non-radiographic axial spondyloarthritis (nr-axSpA)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

- 1 Diagnosis of one of the following:
 - Ankylosing spondylitis (AS)
 - Active non-radiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation present (i.e. lab C-reactive protein elevated, imaging scans indicate inflammation)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq		
Diagnosis	Ankylosing Spondylitis (AS), Non-radiographic axial spondyloarthritis (nr-axSpA)	

Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

- 1 Diagnosis of one of the following:
 - Ankylosing spondylitis (AS)
 - Active non-radiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation present (i.e. lab C-reactive protein elevated, imaging scans indicate inflammation)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq		
Diagnosis	Moderate to Severely Active Ulcerative Colitis (UC)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Member is considered high-risk based on ONE of the following characteristics:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High CRP and ESR

- Steroid-requiring disease
- History of hospitalization
- C. difficile infection
- CMV infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

AND

3.1.3 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq		
Diagnosis	Moderate to Severely Active Ulcerative Colitis (UC)	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AN	D
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2 - Prescribed by or in consultation with a gastroenterologist

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** Member is considered high-risk based on ONE of the following characteristics:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High CRP and ESR
 - Steroid-requiring disease
 - History of hospitalization
 - C. difficile infection
 - CMV infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

AND

3.1.3 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq

Diagnosis	Atopic Dermatitis (AD)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

1 - Diagnosis of moderate to severe atopic dermatitis (AD) based on body surface area (greater than 10%), extent of disability, extent of pruritus, or impact on sleep, quality of life, current use of systemic immunomodulators)

AND

2 - Prescribed by or in consultation with a dermatologist, allergist, or immunologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to ONE of the following:
 - Biologics used in AD (e.g., dupilumab, tralokinumab, abrocitinib)
 - Other systemic agents (e.g. methotrexate, cyclosporine, azathioprine, etc.)

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq	
Diagnosis	Atopic Dermatitis (AD)

Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

1 - Diagnosis of moderate to severe atopic dermatitis (AD) based on body surface area (greater than 10%), extent of disability, extent of pruritus, or impact on sleep, quality of life, current use of systemic immunomodulators)

AND

2 - Prescribed by or in consultation with a dermatologist, allergist, or immunologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to ONE of the following:
 - Biologics used in AD (e.g., dupilumab, tralokinumab, abrocitinib)
 - Other systemic agents (e.g. methotrexate, cyclosporine, azathioprine, etc.)

OR

3.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq	
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)
Approval Length	12/31/2039

Guideline Type		Prior Authorization - Ali Plans except il and Min Plans		
Product Name	Generic Name		GPI	Brand/Generic
RINVOQ	UPADACITINIB TAB ER 24HR 15 MG		66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG		66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG		66603072007540	Brand

1 - Diagnosis of moderate to severely active Crohn's disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

3 - Member is 18 years of age or older

AND

- 4 One of the following:
- **4.1** All of the following:
- **4.1.1** One of the following:
- **4.1.1.1** Member is considered high-risk based on ONE of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease

• Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)

OR

- **4.1.1.2** Both of the following:
- **4.1.1.2.1** Member is considered low-risk

AND

4.1.1.2.2 One of the following:

- Trial and failure, contraindication, or intolerance to one conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with one conventional therapy
- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

AND

4.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

4.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Rinvoq			
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)		
Approval Length	12 month(s)		
Guideline Type	Prior Authorization - IL and MN Plans		
Product Generic Name Name		GPI	Brand/Generic

RINVOQ	UPADACITINIB TAB ER 24HR 15 MG	66603072007520	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 30 MG	66603072007530	Brand
RINVOQ	UPADACITINIB TAB ER 24HR 45 MG	66603072007540	Brand

1 - Diagnosis of moderate to severely active Crohn's disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

3 - Member is 18 years of age or older

AND

- 4 One of the following:
- **4.1** All of the following:
- **4.1.1** One of the following:
- **4.1.1.1** Member is considered high-risk based on ONE of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease
 - Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)

OR

4.1.1.2 Both of the following:

4.1.1.2.1 Member is considered low-risk

AND

4.1.1.2.2 One of the following:

- Trial and failure, contraindication, or intolerance to one conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with one conventional therapy
- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

AND

4.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

4.2 Continuation of prior therapy with upadacitinib, verified by paid claims or medical records (e.g. chart notes)

Date	Notes
4/15/2024	Update COT Language

Rytary	(Carbidopa/Levodopa)	
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Prior Authorization Guideline

Guideline ID	GL-128987
Guideline Name	Rytary (Carbidopa/Levodopa)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Rytary	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
RYTARY	CARBIDOPA & LEVODOPA CAP ER 23.75-95 MG	73209902100220	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 36.25-145 MG	73209902100230	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 48.75-195 MG	73209902100240	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 61.25-245 MG	73209902100250	Brand

1 - Have a diagnosis of Parkinson's disease, post-encephalitic parkinsonism, or parkinsonism following intoxication from carbon monoxide or manganese

AND

2 - Prescribed by, or in consultation with, a Neurologist

AND

3 - Have experienced breakthrough symptoms despite titrated treatment with CONCURRENT immediate-release and extended-release carbidopa/levodopa generics

Product Name: Rytary	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
RYTARY	CARBIDOPA & LEVODOPA CAP ER 23.75-95 MG	73209902100220	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 36.25-145 MG	73209902100230	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 48.75-195 MG	73209902100240	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 61.25-245 MG	73209902100250	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Rytary		
Approval Length 12/31/2039		
Guideline Type	Prior Authorization - All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
RYTARY	CARBIDOPA & LEVODOPA CAP ER 23.75-95 MG	73209902100220	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 36.25-145 MG	73209902100230	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 48.75-195 MG	73209902100240	Brand
RYTARY	CARBIDOPA & LEVODOPA CAP ER 61.25-245 MG	73209902100250	Brand

- **1** All of the following:
- **1.1** Have a diagnosis of Parkinson's disease, post-encephalitic parkinsonism, or parkinsonism following intoxication from carbon monoxide or manganese

AND

1.2 Prescribed by, or in consultation with, a Neurologist

AND

1.3 Have experienced breakthrough symptoms despite titrated treatment with CONCURRENT immediate-release and extended-release carbidopa/levodopa generics

OR

2 - Person is new to the plan (within the past 90 days) and submission of medical notes (e.g., chart notes) from the past 12 months that the person is continuing therapy with the requested medication

Date	Notes
9/20/2023	New Program

Sa	Samsca (Tolvaptan)			
Decidado	nagerunnen terdigeligent. Derfile meg have have neural, sowered, e	classed. Verify that the halo pushes to the connectific and invades.		

Prior Authorization Guideline

Guideline ID	GL-131950
Guideline Name Samsca (Tolvaptan)	
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic: Tolvaptan	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only	

	roduct ame	Generic Name	GPI	Brand/Generic
TO	OLVAPTAN	TOLVAPTAN TAB 15 MG	30454060000320	Generic
TO	OLVAPTAN	TOLVAPTAN TAB 30 MG	30454060000330	Generic

Approval Criteria

1 - Diagnosis of severe hypervolemic or euvolemic hyponatremia (sodium level < 125 mEq/L) OR symptomatic less severe hyponatremia (sodium level 125 mEq/L -134 mEq/L)

AND

2 - Current hospitalization for hyponatremia

AND

3 - Symptoms or low serum sodium levels persist despite supervised fluid restriction and appropriate sodium supplementation

Product Name: Generic: Tolvaptan		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
TOLVAPTAN	TOLVAPTAN TAB 15 MG	30454060000320	Generic
TOLVAPTAN	TOLVAPTAN TAB 30 MG	30454060000330	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Generic: Tolvaptan		
Approval Length	12/31/2039	
Guideline Type	Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
TOLVAPTAN	TOLVAPTAN TAB 15 MG	30454060000320	Generic
TOLVAPTAN	TOLVAPTAN TAB 30 MG	30454060000330	Generic

1 - Diagnosis of severe hypervolemic or euvolemic hyponatremia (sodium level < 125 mEq/L) OR symptomatic less severe hyponatremia (sodium level 125 mEq/L -134 mEq/L)

AND

2 - Current hospitalization for hyponatremia

AND

3 - Symptoms or low serum sodium levels persist despite supervised fluid restriction and appropriate sodium supplementation

Date	Notes
10/31/2023	New program

Sarafem (Fluoxetine 10 mg Tablet)		
[3] The Mandalings are contributions. The first in the contribution contribution are contributed as a contribution of the cont		

Prior Authorization Guideline

Guideline ID	GL-137190
Guideline Name	Sarafem (Fluoxetine 10 mg Tablet)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Fluoxetine 10 mg Tablet		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FLUOXETINE HYDROCHLORIDE	FLUOXETINE HCL TAB 10 MG	58160040000310	Generic
FLUOXETINE HYDROCHLORIDE	FLUOXETINE HCL (PMDD) TAB 10 MG	62206040000310	Generic

Approval Criteria

1 - Person requires a dose that cannot be met using the preferred fluoxetine capsule formulations (5 or 15 mg per day

AND

2 - An adequate trial of daily dosing at 10 mg and 20 mg capsule did not control symptoms or caused intolerable side effects

Product Name: Fluoxetine 10 mg Tablet		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FLUOXETINE HYDROCHLORIDE	FLUOXETINE HCL TAB 10 MG	58160040000310	Generic
FLUOXETINE HYDROCHLORIDE	FLUOXETINE HCL (PMDD) TAB 10 MG	62206040000310	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Fluoxetine 10 mg Tablet		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
FLUOXETINE HYDROCHLORIDE	FLUOXETINE HCL TAB 10 MG	58160040000310	Generic
FLUOXETINE HYDROCHLORIDE	FLUOXETINE HCL (PMDD) TAB 10 MG	62206040000310	Generic

Approval Criteria

1 - Person requires a dose that cannot be met using the preferred fluoxetine capsule formulations (5 or 15 mg per day

AND

2 - An adequate trial of daily dosing at 10 mg and 20 mg capsule did not control symptoms or caused intolerable side effects

Product Name: Fluoxetine 10 mg Tablet				
Guideline Type Quantity limit				
Product Name	ct Name Generic Name		GPI	Brand/Generic
FLUOXETINE HYDROCHLORIDE		XETINE HCL TAB 10 MG	58160040000310	Generic
FLUOXETINE HCL (PMDD) TAB 10 MG HYDROCHLORIDE		62206040000310	Generic	

Approval Criteria

1 - Doses greater than 15mg (1.5 tablets) per day should be denied. Doses greater than 15mg (1.5 tablets) per day require use of the preferred fluoxetine capsule (ie. fluoxetine 10mg capsule, fluoxetine 20mg capsule, fluoxetine 40mg capsule).

Date	Notes
11/30/2023	Update Program

Savella (milnacipran)				
The behavior and the second to				

Prior Authorization Guideline

Guideline ID	GL-129647
Guideline Name	Savella (milnacipran)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Savella		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
SAVELLA	MILNACIPRAN HCL TAB 12.5 MG	62504050100320	Brand
SAVELLA	MILNACIPRAN HCL TAB 25 MG	62504050100330	Brand
SAVELLA	MILNACIPRAN HCL TAB 50 MG	62504050100340	Brand
SAVELLA	MILNACIPRAN HCL TAB 100 MG	62504050100350	Brand
SAVELLA TITRATION PACK	MILNACIPRAN HCL TAB 12.5 MG (5) & 25 MG (8) & 50 MG (42) PAK	62504050106320	Brand

1 - Trial of at least 2 preferred treatment options for fibromyalgia: amitriptyline, cyclobenzaprine, venlafaxine, duloxetine, gabapentin, pregabalin

Product Name: Savella		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
SAVELLA	MILNACIPRAN HCL TAB 12.5 MG	62504050100320	Brand
SAVELLA	MILNACIPRAN HCL TAB 25 MG	62504050100330	Brand
SAVELLA	MILNACIPRAN HCL TAB 50 MG	62504050100340	Brand
SAVELLA	MILNACIPRAN HCL TAB 100 MG	62504050100350	Brand
SAVELLA TITRATION PACK	MILNACIPRAN HCL TAB 12.5 MG (5) & 25 MG (8) & 50 MG (42) PAK	62504050106320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Savella	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
SAVELLA	MILNACIPRAN HCL TAB 12.5 MG	62504050100320	Brand
SAVELLA	MILNACIPRAN HCL TAB 25 MG	62504050100330	Brand
SAVELLA	MILNACIPRAN HCL TAB 50 MG	62504050100340	Brand
SAVELLA	MILNACIPRAN HCL TAB 100 MG	62504050100350	Brand

	MILNACIPRAN HCL TAB 12.5 MG (5) & 25 MG (8) & 50 MG (42) PAK	62504050106320	Brand
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1 - Trial of at least 2 preferred treatment options for fibromyalgia: amitriptyline, cyclobenzaprine, venlafaxine, duloxetine, gabapentin, pregabalin

Date	Notes
10/6/2023	New Program

Secuado (asenapine patches)	
2 The bindings among to single, or Track was have two most, resident, and death belt parties the controlls and harden	

Prior Authorization Guideline

Guideline ID	GL-128186
Guideline Name	Secuado (asenapine patches)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Secuado	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SECUADO	ASENAPINE TD PATCH 24 HR 3.8 MG/24HR	59155015008520	Brand
SECUADO	ASENAPINE TD PATCH 24 HR 5.7 MG/24HR	59155015008530	Brand
SECUADO	ASENAPINE TD PATCH 24 HR 7.6 MG/24HR	59155015008540	Brand

Approval Criteria

- **1** One of the following:
- **1.1** Trial and failure or intolerance to the sublingual tablet formulation at an equivalent dose

OR

1.2 Person with swallowing impairment or other medical condition that prevents use of solid dose forms

OR

- 2 For Minnesota Plans One of the following:
- **2.1** When prescribed for emotional disturbance or mental illness, approve if prescriber provides submission of medial records (e.g. chart notes) that all equivalent drugs in the formulary were considered and it has been determined that the drug prescribed will best treat the member's condition

OR

- **2.2** Both of the following for continuation of care: (i.e. formulary changes or new member [as evidenced by coverage effective date of less than or equal to 90 days]):
 - 2.2.1 Member has been treated with the drug for 90 days prior to the change

AND

2.2.2 Prescriber provides submission of medical records (e.g., chart notes) that the drug prescribed will best treat the member's condition

Product Name: Secuado				
Approval Le	ength	12 month(s)		
Therapy Sta	age	Reauthorization		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic

SECUADO	ASENAPINE TD PATCH 24 HR 3.8 MG/24HR	59155015008520	Brand
SECUADO	ASENAPINE TD PATCH 24 HR 5.7 MG/24HR	59155015008530	Brand
SECUADO	ASENAPINE TD PATCH 24 HR 7.6 MG/24HR	59155015008540	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Secuado	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
SECUADO	ASENAPINE TD PATCH 24 HR 3.8 MG/24HR	59155015008520	Brand
SECUADO	ASENAPINE TD PATCH 24 HR 5.7 MG/24HR	59155015008530	Brand
SECUADO	ASENAPINE TD PATCH 24 HR 7.6 MG/24HR	59155015008540	Brand

Approval Criteria

1 - Trial and failure or intolerance to the sublingual tablet formulation at an equivalent dose

OR

2 - Person with swallowing impairment or other medical condition that prevents use of solid dose forms

Date	Notes
9/7/2023	New Program

Serotonin Modulating Antidepre	essant
(3) The foreign proving the finite transcription and the state of the finite transcription control or control	

Prior Authorization Guideline

Guideline ID	GL-127881
Guideline Name	Serotonin Modulating Antidepressants
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Trintellix	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Step Therapy - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TRINTELLIX	VORTIOXETINE HBR TAB 5 MG (BASE EQUIV)	58120093100310	Brand
TRINTELLIX	VORTIOXETINE HBR TAB 10 MG (BASE EQUIV)	58120093100320	Brand
TRINTELLIX	VORTIOXETINE HBR TAB 20 MG (BASE EQUIV)	58120093100340	Brand

Approval Criteria

- **1** Trial and failure, contraindication, or intolerance of at least two preferred antidepressants within the Selective Serotonin Reuptake inhibitor (SSRI) or Serotonin Norepinephrine Reuptake inhibitor (SNRI) drug classes
 - citalopram
 - escitalopram
 - sertraline
 - paroxetine
 - fluoxetine
 - venlafaxine
 - duloxetine

Product Name: Trintellix	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Step Therapy - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TRINTELLIX	VORTIOXETINE HBR TAB 5 MG (BASE EQUIV)	58120093100310	Brand
TRINTELLIX	VORTIOXETINE HBR TAB 10 MG (BASE EQUIV)	58120093100320	Brand
TRINTELLIX	VORTIOXETINE HBR TAB 20 MG (BASE EQUIV)	58120093100340	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Trintellix	
Approval Length 12/31/2039	
Guideline Type Step Therapy - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TRINTELLIX	VORTIOXETINE HBR TAB 5 MG (BASE EQUIV)	58120093100310	Brand
TRINTELLIX	VORTIOXETINE HBR TAB 10 MG (BASE EQUIV)	58120093100320	Brand
TRINTELLIX	VORTIOXETINE HBR TAB 20 MG (BASE EQUIV)	58120093100340	Brand

- **1** Trial and failure, contraindication, or intolerance of at least two preferred antidepressants within the Selective Serotonin Reuptake inhibitor (SSRI) or Serotonin Norepinephrine Reuptake inhibitor (SNRI) drug classes
 - citalopram
 - escitalopram
 - sertraline
 - paroxetine
 - fluoxetine
 - venlafaxine
 - duloxetine

Date	Notes
8/21/2023	New Program

S	Signifor (Pasireotide Diasparte)				
•	Sections provided with the section and case case of the big decision and decision.				

Prior Authorization Guideline

Guideline ID	GL-131411
Guideline Name	Signifor (Pasireotide Diasparte)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Signifor	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.3 MG/ML (BASE EQUIV)	30170075202020	Brand
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.6 MG/ML (BASE EQUIV)	30170075202030	Brand
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.9 MG/ML (BASE EQUIV)	30170075202040	Brand

- 1 Diagnosis of Cushing disease
- 1.1 Ongoing symptoms despite pituitary surgery or nonsurgical candidate

AND

2 - Age greater than or equal to 18 years

AND

3 - Trial and failure, contraindication, or intolerance to octreotide

OR

4 - Other FDA labeled indications

Product Name: Signifor	
Approval Length 12 month(s)	
Therapy Stage	Reauthorization
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.3 MG/ML (BASE EQUIV)	30170075202020	Brand
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.6 MG/ML (BASE EQUIV)	30170075202030	Brand
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.9 MG/ML (BASE EQUIV)	30170075202040	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Signifor	
Approval Length	12/31/2039
Guideline Type Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.3 MG/ML (BASE EQUIV)	30170075202020	Brand
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.6 MG/ML (BASE EQUIV)	30170075202030	Brand
SIGNIFOR	PASIREOTIDE DIASPARTATE INJ 0.9 MG/ML (BASE EQUIV)	30170075202040	Brand

- 1 Diagnosis of Cushing disease
- 1.1 Ongoing symptoms despite pituitary surgery or nonsurgical candidate

AND

2 - Age greater than or equal to 18 years

AND

3 - Trial and failure, contraindication, or intolerance to octreotide

OR

4 - Other FDA labeled indications

2. Revision History

Date	Notes
10/24/2023	New program

Simponi (golimumab)
Control of the contro

Prior Authorization Guideline

Guideline ID	GL-145260
Guideline Name	Simponi (golimumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Simponi			
Diagnosis Psoriatic Arthritis (PsA)			
Approval Length 12/31/2039			
Guideline	Prior Authorization - All plans except IL and MN Plans		
Product Name	Generic Name GPI Brand/Gener		Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML 6627004000D520 Brand		Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML 6627004000D540 Brand		Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED 6627004000E520 Brand SYRINGE 50 MG/0.5ML		

SIMPONI GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand
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1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a Dermatologist or Rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - · active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

Product Name: Simponi				
Diagnosis		Psoriatic Arthritis (PsA)		
Approval L	ength	12 month(s)		
Guideline Type Prior Authorization - IL and MN Plans				
Product Name	Generic Name		GPI	Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML		6627004000D520	Brand

SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML	6627004000D540	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.5ML	6627004000E520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a Dermatologist or Rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

Product Name: Simponi	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML	6627004000D520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML	6627004000D540	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.5ML	6627004000E520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand

1 - Diagnosis of moderate to severely active Rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - Methotrexate (MTX)*
 - Leflunomide
 - Hydroxychloroquine
 - Sulfasalazine

OR

Notes	*Absolute contraindications to methotrexate are pregnancy, nursing, a
	Icoholism, alcoholic liver disease or other chronic liver disease, immun
	odeficiency syndromes, bone marrow hyperplasia, leukopenia, thromb
	ocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Simponi	
Diagnosis Moderate to Severely Active Rheumatoid Arthritis	
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML	6627004000D520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML	6627004000D540	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.5ML	6627004000E520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand

1 - Diagnosis of moderate to severely active Rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a Rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - Methotrexate (MTX)*
 - Leflunomide
 - Hydroxychloroquine
 - Sulfasalazine

OR

3.2 Continuation of pr (e.g. chart notes)	ior therapy with golimumab, verified by paid claims or medical records
Notes	*Absolute contraindications to methotrexate are pregnancy, nursing, a lcoholism, alcoholic liver disease or other chronic liver disease, immun odeficiency syndromes, bone marrow hyperplasia, leukopenia, thromb ocytopenia or significant anemia, or hypersensitivity to methotrexate.

Product Name: Simponi	
Diagnosis	Ankylosing spondylitis (AS)
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML	6627004000D520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML	6627004000D540	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.5ML	6627004000E520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand

1 - Diagnosis of Ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a Rheumatologist

- 3 One of the following:
- **3.1** Trial and failure to a 1-month trial of scheduled prescription doses of two different NSAIDs (e.g., naproxen, nabumetone, diclofenac, etc.)

OR

3.2 Continuation of prior therapy with golimumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Simponi	
Diagnosis	Ankylosing spondylitis (AS)
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML	6627004000D520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML	6627004000D540	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.5ML	6627004000E520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand

Approval Criteria

1 - Diagnosis of Ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a Rheumatologist

- 3 One of the following:
- **3.1** Trial and failure to a 1-month trial of scheduled prescription doses of two different NSAIDs (e.g., naproxen, nabumetone, diclofenac, etc.)

OR

3.2 Continuation of prior therapy with golimumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Simponi	
Diagnosis	Ulcerative Colitis (UC)
Approval Length	12/31/2039*
Guideline Type	Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML	6627004000D520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML	6627004000D540	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.5ML	6627004000E520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a Gastroenterologist

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Trial and failure or contraindication to at least a short course (2-4 weeks) of oral corticosteroids

- **3.1.2** High-risk individual as evidence by ONE of the following:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High C-reactive protein (CRP) and erythrocyte sedimentation rate (ESR)
 - Steroid-requiring disease
 - History of hospitalization
 - C. difficile infection
 - CMV infection

OR

*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.11 (3 every 28 days) for 30 days
Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Simponi	
Diagnosis	Ulcerative Colitis (UC)
Approval Length	12 Month(s)*
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 50 MG/0.5ML	6627004000D520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 100 MG/ML	6627004000D540	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 50 MG/0.5ML	6627004000E520	Brand
SIMPONI	GOLIMUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 100 MG/ML	6627004000E540	Brand

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a Gastroenterologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Trial and failure or contraindication to at least a short course (2-4 weeks) of oral corticosteroids

AND

- **3.1.2** High-risk individual as evidence by ONE of the following:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High C-reactive protein (CRP) and erythrocyte sedimentation rate (ESR)
 - Steroid-requiring disease
 - History of hospitalization
 - C. difficile infection
 - CMV infection

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve at GPI 14 with a MDD of 0.11 (3 every 28 days) for

30 days Second PA: Approve at GPI 10 for 12 months
Second 1 A. Approve at Of 1 10 for 12 months

2. Revision History

Date	Notes
4/15/2024	Update COT language

Skyrizi (risankizumab)
(2) The contract of the contra

Prior Authorization Guideline

Guideline ID	GL-145256
Guideline Name	Skyrizi (risankizumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/1/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Skyrizi				
Diagnosis		Plaque Psoriasis		
Approval Length		12/31/2039*		
Guideline	Туре	Prior Authorization - All plans except IL and MN Plans		
Product Name	Generic Name GPI Brand/Gene		Brand/Generic	
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN 5250406070E210 Brand CARTRIDGE 180 MG/1.2ML		Brand	
SKYRIZI	YRIZI RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN 5250406070E220 Brand CARTRIDGE 360 MG/2.4ML		Brand	
SKYRIZI PEN	3020007070D020		Brand	

SKYRIZI RISANKIZUMAB-RZAA 150 MG/ML	SOLN PREFILLED SYRINGE	9025057070E540	Brand
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1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- **3** ONE of the following:
- **3.1** Both of the following:
- **3.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g., nails, scalp, genitals, or intertriginous areas

AND

3.1.2 Trial and failure, contraindication or intolerance to topical treatment (e.g., topical corticosteroids, calcipotriene, retinoids, calcineurin inhibitors, tazarotene)

OR

Natas	*For your stands to the years. Foton 2.DAs as follows with the course stand
Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
	30 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Skyrizi	
Diagnosis	Plaque Psoriasis
Approval Length	12 Month(s)*
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 180 MG/1.2ML	5250406070E210	Brand
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 360 MG/2.4ML	5250406070E220	Brand
SKYRIZI PEN	RISANKIZUMAB-RZAA SOLN AUTO-INJECTOR 150 MG/ML	9025057070D520	Brand
SKYRIZI	RISANKIZUMAB-RZAA SOLN PREFILLED SYRINGE 150 MG/ML	9025057070E540	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

- **3** ONE of the following:
 - **3.1** Both of the following:
 - **3.1.1** One of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than or equal to 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g., nails, scalp, genitals, or intertriginous areas

3.1.2 Trial and failure, contraindication or intolerance to topical treatment (e.g., topical corticosteroids, calcipotriene, retinoids, calcineurin inhibitors, tazarotene)

OR

3.2 Continuation of prior therapy with risankizumab, verified by paid claims or medical records (e.g. chart notes)

*For new starts to therapy: Enter 2 PAs as follows with the same start
date:
First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
30 days
Second PA: Approve at GPI 10 for 12 months

Product Name: Skyrizi	
Diagnosis	Psoriatic Arthritis (PsA)
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 180 MG/1.2ML	5250406070E210	Brand
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 360 MG/2.4ML	5250406070E220	Brand
SKYRIZI PEN	RISANKIZUMAB-RZAA SOLN AUTO-INJECTOR 150 MG/ML	9025057070D520	Brand
SKYRIZI	RISANKIZUMAB-RZAA SOLN PREFILLED SYRINGE 150 MG/ML	9025057070E540	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis

2 - Prescribed by or in consultation with a Dermatologist or Rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - Actively inflamed joints
 - Axial disease
 - Active skin, nail, or scalp psoriasis involvement
 - Dactylitis
 - Enthesitis

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
110163	date:
	First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
	30 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Skyrizi	
Diagnosis	Psoriatic Arthritis (PsA)
Approval Length	12 Month(s)*
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 180 MG/1.2ML	5250406070E210	Brand
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 360 MG/2.4ML	5250406070E220	Brand
SKYRIZI PEN	RISANKIZUMAB-RZAA SOLN AUTO-INJECTOR 150 MG/ML	9025057070D520	Brand
SKYRIZI	RISANKIZUMAB-RZAA SOLN PREFILLED SYRINGE 150 MG/ML	9025057070E540	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis

AND

2 - Prescribed by or in consultation with a Dermatologist or Rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - Actively inflamed joints
 - Axial disease
 - Active skin, nail, or scalp psoriasis involvement
 - Dactylitis
 - Enthesitis

OR

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
	30 days
	Second PA: Approve at GPI 10 for 12 months

Product Name: Skyrizi		
Diagnosis	Crohn's Disease	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 180 MG/1.2ML	5250406070E210	Brand
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 360 MG/2.4ML	5250406070E220	Brand
SKYRIZI PEN	RISANKIZUMAB-RZAA SOLN AUTO-INJECTOR 150 MG/ML	9025057070D520	Brand
SKYRIZI	RISANKIZUMAB-RZAA SOLN PREFILLED SYRINGE 150 MG/ML	9025057070E540	Brand

1 - Diagnosis of moderate to severely active Crohn's disease

AND

2 - Member is greater than 18 years of age

AND

3 - Prescribed by or in consultation with a Gastroenterologist

- **4** ONE of the following:
- **4.1** Documentation that the patient has been established on therapy on the medical benefit and one of the following traits:
 - **4.1.1** Member is a High-risk individual with ONE of the following traits:
 - Age less than 30 at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease

• Extraintestinal manifestations of inflammation (i.e. uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy, etc.)]

OR

4.1.2 Both of the following:

4.1.2.1 Member is a Low-risk individual

AND

4.1.2.2 ONE of the following:

- Intolerance or contraindication to 1 conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with 1 conventional therapy
- Demonstrated steroid dependence
- Conventional therapy is clinically inappropriate based on location of disease

OR

Product Name: Skyrizi		
Diagnosis	Crohn's Disease	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 180 MG/1.2ML	5250406070E210	Brand
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 360 MG/2.4ML	5250406070E220	Brand
SKYRIZI PEN	RISANKIZUMAB-RZAA SOLN AUTO-INJECTOR 150 MG/ML	9025057070D520	Brand

SKYRIZI	RISANKIZUMAB-RZAA SOLN PREFILLED SYRINGE 150 MG/ML	9025057070E540	Brand
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1 - Diagnosis of moderate to severely active Crohn's disease

AND

2 - Member is greater than 18 years of age

AND

3 - Prescribed by or in consultation with a Gastroenterologist

AND

- 4 ONE of the following:
- **4.1** Documentation that the patient has been established on therapy on the medical benefit and one of the following traits:
 - **4.1.1** Member is a High-risk individual with ONE of the following traits:
 - Age less than 30 at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease
 - Extraintestinal manifestations of inflammation (i.e. uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy, etc.)]

OR

- **4.1.2** Both of the following:
- 4.1.2.1 Member is a Low-risk individual

4.1.2.2 ONE of the following:

- Intolerance or contraindication to 1 conventional therapy (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with 1 conventional therapy
- Demonstrated steroid dependence
- Conventional therapy is clinically inappropriate based on location of disease

OR

4.2 Continuation of prior therapy with risankizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Skyrizi	
Diagnosis	Crohn's Disease
Approval Length	12 month(s)
Guideline Type	Quantity Limit - All Plans

Product Name	Generic Name	GPI	Brand/Generic
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 180 MG/1.2ML	5250406070E210	Brand
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 360 MG/2.4ML	5250406070E220	Brand
SKYRIZI PEN	RISANKIZUMAB-RZAA SOLN AUTO-INJECTOR 150 MG/ML	9025057070D520	Brand
SKYRIZI	RISANKIZUMAB-RZAA SOLN PREFILLED SYRINGE 150 MG/ML	9025057070E540	Brand

Approval Criteria

- **1** All of the following:
- **1.1** Trial and failure of a two-month trial of every 12 week therapy after completion of 3 doses of IV infusion for the induction dosing regimen

1.2 Provision of published literature supporting efficacy and safety of dosing regimen

AND

1.3 Based on subtherapeutic drug concentrations and absence (or low levels) of drug antibodies (when clinical lab available).

OR

2 - Continuation of previous therapy with ustekinumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

Product Name: Skyrizi		
Diagnosis	Plaque Psoriasis, Psoriatic Arthritis (PsA)	
Approval Length	12 month(s)	
Guideline Type	Quantity Limit - All Plans	

Product Name	Generic Name	GPI	Brand/Generic
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 180 MG/1.2ML	5250406070E210	Brand
SKYRIZI	RISANKIZUMAB-RZAA SUBCUTANEOUS SOLN CARTRIDGE 360 MG/2.4ML	5250406070E220	Brand
SKYRIZI PEN	RISANKIZUMAB-RZAA SOLN AUTO-INJECTOR 150 MG/ML	9025057070D520	Brand
SKYRIZI	RISANKIZUMAB-RZAA SOLN PREFILLED SYRINGE 150 MG/ML	9025057070E540	Brand

Approval Criteria

- 1 Both of the following:
- **1.1** Trial and failure of an adherent 3-month trial of standard maintenance dosing (every 12 weeks) with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)

1.2 Provision of published literature supporting efficacy and safety of dosing

OR

2 - Continuation of previous therapy with ustekinumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

2. Revision History

Date	Notes
4/17/2024	Update COT language

Soliqua (Insulin Glargine/Lixisenatide			
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Prior Authorization Guideline

Guideline ID	GL-129739
Guideline Name	Soliqua (Insulin Glargine/Lixisenatide)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Soliqua	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SOLIQUA 100/33	INSULIN GLARGINE-LIXISENATIDE SOL PEN-INJ 100- 33 UNIT-MCG/ML	2799100235D220	Brand

Approval Criteria

1 - Diagnosis of insulin dependent type 2 diabetes mellitus with a total basal insulin dose of less than 60 units/day

2 - Prescribed by, or in consultation with, an Endocrinologist or Certified Diabetic Educator

AND

3 - Trial and failure after an adequate trial, intolerance, or contraindication to use of ALL formulary basal insulins (e.g., Semglee-yfgn, insulin degludec) and ALL formulary glucagon-like peptide (GLP) 1 agonists (e.g., Trulicity) in combination

AND

4 - Prescriber supplies published literature to support the requested combination product will produce different clinical results than using the formulary basal insulins and GLP-1 agonists as separate products

Product Name: Soliqua	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
SOLIQUA 100/33	INSULIN GLARGINE-LIXISENATIDE SOL PEN-INJ 100- 33 UNIT-MCG/ML	2799100235D220	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Soliqua	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
SOLIQUA 100/33	INSULIN GLARGINE-LIXISENATIDE SOL PEN-INJ 100- 33 UNIT-MCG/ML	2799100235D220	Brand

1 - Diagnosis of insulin dependent type 2 diabetes mellitus with a total basal insulin dose of less than 60 units/day

AND

2 - Prescribed by, or in consultation with, an Endocrinologist or Certified Diabetic Educator

AND

3 - Trial and failure after an adequate trial, intolerance, or contraindication to use of ALL formulary basal insulins (e.g., Semglee-yfgn, insulin degludec) and ALL formulary glucagon-like peptide (GLP) 1 agonists (e.g., Trulicity) in combination

AND

4 - Prescriber supplies published literature to support the requested combination product will produce different clinical results than using the formulary basal insulins and GLP-1 agonists as separate products

2. Revision History

Date	Notes
10/25/2023	New Program

•	Solosec (secnidazole)		
	(2) The blackings contributing on The first the section most, waster a state with section price the contribute states		

Prior Authorization Guideline

Guideline ID	GL-132774
Guideline Name	Solosec (secnidazole)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Solosec	
Diagnosis	Bacterial vaginosis
Approval Length	12 month (s) with a fill count = 1
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SOLOSEC	SECNIDAZOLE GRANULES PACKET 2 GM	14000080003020	Brand

Approval Criteria

1 - Diagnosis of bacterial vaginosis

2 - Trial and failure, contraindication, or intolerance to metronidazole (oral or vaginal gel) and clindamycin

Product Name: Solosec	
Diagnosis	Bacterial vaginosis
Approval Length	One time fill
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
SOLOSE	SECNIDAZOLE GRANULES PACKET 2 GM	14000080003020	Brand

Approval Criteria

1 - Diagnosis of bacterial vaginosis

AND

2 - Trial and failure, contraindication, or intolerance to metronidazole (oral or vaginal gel) and clindamycin

Product Name: Solosec	
Diagnosis	trichomoniasis
Approval Length	12 month (s) with a fill count = 1
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SOLOSEC	SECNIDAZOLE GRANULES PACKET 2 GM	14000080003020	Brand

Approval Criteria

1 - Diagnosis of trichomoniasis

AND

- **2** Trial and failure, contraindication, or intolerance to a seven day course of one of the following:
 - oral metronidazole
 - tinidazole

Product Name: Solosec	
Diagnosis	trichomoniasis
Approval Length	One time fill
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
SOLOSEC	SECNIDAZOLE GRANULES PACKET 2 GM	14000080003020	Brand

Approval Criteria

1 - Diagnosis of trichomoniasis

AND

- **2** Trial and failure, contraindication, or intolerance to a seven day course of one of the following:
 - oral metronidazole
 - tinidazole

2. Revision History

Date	Notes

10/31/2023	New Program

Somatropin		
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Prior Authorization Guideline

Guideline ID	GL-145255	
Guideline Name	Somatropin	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	4/1/2012
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Omnitrope					
Diagnosis		Pediatric [less than 18 years of age])			
Approval Length		until age 18			
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans		ns			
Product Name	Generic Name		GPI	Brand/Generic	
OMNITROPE	SOMATROPIN SOLUTION CARTRIDGE 5 MG/1.5ML		3010002000E210	Brand	
OMNITROPE	SOMATROPIN SOLUTION CARTRIDGE 10 MG/1.5ML		3010002000E213	Brand	
OMNITROPE	SOMATROPIN FOR INJ 5.8 MG		30100020002123	Brand	

Approval Criteria
1 - One of the following:
1.1 ALL of the following:
1.1.1 Submission of medical records (e.g., chart notes) documenting radiological evidence of open epiphyses with date completed
AND
1.1.2 Submission of medical records (e.g., chart notes) documenting that the child's growth velocity value is subnormal (age specific growth rate less than the 25th percentile)
AND
1.1.3 Submission of medical records (e.g., chart notes) documenting that the child has delayed bone age (e.g., provide date of completion and the value of bone age)
AND
1.1.4 Submission of medical records (e.g., chart notes) documenting (e.g., provide date and value of test) that the child has subnormal GH response to at least one provocative stimulation test (less than 10 ng/mL)
AND
1.1.5 Member is less than 18 years of age
OR
1.2 Both of the following:
1.2.1 Member is less than 18 years of age

1.2.2 Diagnosis of Turner syndrome

AND

2 - Prescribed by or in consultation with an endocrinologist

Product Name: Omnitrope		
Diagnosis	Pediatric [less than 18 years of age])	
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
OMNITROPE	SOMATROPIN SOLUTION CARTRIDGE 5 MG/1.5ML	3010002000E210	Brand
OMNITROPE	SOMATROPIN SOLUTION CARTRIDGE 10 MG/1.5ML	3010002000E213	Brand
OMNITROPE	SOMATROPIN FOR INJ 5.8 MG	30100020002123	Brand

Approval Criteria

- 1 One of the following:
- **1.1** ALL of the following:
- **1.1.1** Submission of medical records (e.g., chart notes) documenting radiological evidence of open epiphyses with date completed

AND

1.1.2 Submission of medical records (e.g., chart notes) documenting that the child's growth velocity value is subnormal (age specific growth rate less than the 25th percentile)

AND

1.1.3 Submission of medical records (e.g., chart notes) documenting that the child has delayed bone age (e.g., provide date of completion and the value of bone age)

AND

1.1.4 Submission of medical records (e.g., chart notes) documenting (e.g., provide date and value of test) that the Child has subnormal GH response to at least one provocative stimulation test (less than 10 ng/mL)

AND

1.1.5 Member is less than 18 years of age

OR

- **1.2** Both of the following:
- 1.2.1 Member is less than 18 years of age

AND

1.2.2 Diagnosis of Turner syndrome

AND

2 - Prescribed by or in consultation with an endocrinologist

Product Name: Omnitrope		
Diagnosis Pediatric [less than 18 years of age])		
Approval Length	12 month(s)	
Therapy Stage Reauthorization		

Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
OMNITROPE	SOMATROPIN SOLUTION CARTRIDGE 5 MG/1.5ML		3010002000E210	Brand
OMNITROPE	SOMATROF	PIN SOLUTION CARTRIDGE 10 MG/1.5ML	3010002000E213	Brand
OMNITROPE	SOMATROF	PIN FOR INJ 5.8 MG	30100020002123	Brand

1 - Submission of medical records (e.g. chart notes) from the past 12 months documenting the member is under the age of 18 and continuing therapy with the requested drug

AND

2 - Prescribed by or in consultation with an endocrinologist

Product Name: Omnitrope		
Diagnosis Adult [18 years of age or older])		
Approval Length 12 month(s)		
Guideline Type Prior Authorization - ALL Plans		

Product Name	Generic Name	GPI	Brand/Generic
OMNITROPE	SOMATROPIN SOLUTION CARTRIDGE 5 MG/1.5ML	3010002000E210	Brand
OMNITROPE	SOMATROPIN SOLUTION CARTRIDGE 10 MG/1.5ML	3010002000E213	Brand
OMNITROPE	SOMATROPIN FOR INJ 5.8 MG	30100020002123	Brand

Approval Criteria

1 - Member is 18 years of age or older

AND

2 - Prescribed by or in consultation with an endocrinologist

AND

3 - One of the following:

3.1 Both of the following:

- Member has growth hormone deficiency as a child
- Submission of medical records (e.g. chart notes) of continued low IGF-1 levels or evidence of GH deficiency as noted by stimulation testing

OR

3.2 Both of the following:

- Submission of medical records (e.g. chart notes) of an abnormal structure of the hypothalamus or pituitary gland on MRI as a result of injury, tumor, infection or
- Evidence of GH deficiency as noted by stimulation testing or a diagnosis of panhypopituitarism

OR

3.3 Both of the following:

- Member has a previous 12-month authorization after the age of 18 years on file or a historical 12-month authorization after the age of 18 on file
- Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is benefitting from drug treatment (i.e., decreased fatigue, increased exercise endurance, age normalized IGF-1 levels, improvements in cholesterol panel, BMD, or body composition) including dates/values if applicable

Product Name: Serostim				
Approval L	ength	1 month(s)		
Therapy St	age	Initial Authorization		
Guideline 1	Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans			ns
Product Generic Name Name		GPI	Brand/Generic	

SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 4 MG	30100020102118	Brand
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 5 MG	30100020102121	Brand
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 6 MG	30100020102125	Brand

1 - Diagnosis of AIDS wasting or cachexia

AND

2 - Member continues on antiviral therapy

Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
	Thew to plan, reduction antona applies

Product Name: Serostim		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 4 MG	30100020102118	Brand
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 5 MG	30100020102121	Brand
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 6 MG	30100020102125	Brand

Approval Criteria

1 - Diagnosis of AIDS wasting or cachexia

AND 2 - Member continues on antiviral therapy		
Notes	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies	

Product Name: Serostim		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type Prior Authorization - ALL Plans		

Product Name	Generic Name	GPI	Brand/Generic
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 4 MG	30100020102118	Brand
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 5 MG	30100020102121	Brand
SEROSTIM	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 6 MG	30100020102125	Brand

1 - Submission of medical records (e.g., chart notes) documenting that member is benefitting from therapy (i.e., weight gain, increased muscle mass)

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
new to plan, reauthorization criteria applies

Product Name: Zorbtive	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - ALL Plans

Product Name	Generic Name	GPI	Brand/Generic
ZORBTIVE	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 8.8 MG	30100020102132	Brand

1 - Diagnosis of Short Bowel Syndrome

AND

2 - Member is on a special diet

Notes	*Member new to the plan (as evidenced by coverage effective date of
	less than or equal to 90 days) who initiated therapy using a manufactu
	rer-sponsored free drug program, provider samples, and/or vouchers
	will go through initial criteria, otherwise for continuation of therapy for
	new to plan, reauthorization criteria applies

Product Name: Zorbtive	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization - ALL Plans	

Product Name	Generic Name	GPI	Brand/Generic
ZORBTIVE	SOMATROPIN (NON-REFRIGERATED) FOR SUBCUTANEOUS INJ 8.8 MG	30100020102132	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months that the member is benefitting from therapy (i.e., improvements in necessary intravenous feeding requirements such as calories required, or volumes infused) including dates/values

Notes	*Member new to the plan (as evidenced by coverage effective date of
	less than or equal to 90 days) who initiated therapy using a manufactu
	rer-sponsored free drug program, provider samples, and/or vouchers
	will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

2. Revision History

Date	Notes
4/15/2024	Update COT language

Somavert (Pegvisomant)					
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Prior Authorization Guideline

Guideline ID	GL-144935
Guideline Name	Somavert (Pegvisomant)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	1/20/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Somavert				
Approval Length		12 month(s)		
Guideline Type		Prior Authorization-IL and MN Plans Only		
Product Name	Generic Name		GPI	Brand/Generic
SOMAVERT	PEGVISOMANT FOR INJ 10 MG (AS PROTEIN)		30180060002120	Brand
SOMAVERT	PEGVISOMANT FOR INJ 15 MG (AS PROTEIN)		30180060002130	Brand
SOMAVERT	PEGVISOMANT FOR INJ 20 MG (AS PROTEIN)		30180060002140	Brand
SOMAVERT	PEGVISOMANT FOR INJ 25 MG (AS PROTEIN)		30180060002150	Brand
SOMAVERT	PEGVISOMANT FOR INJ 30 MG (AS PROTEIN)		30180060002160	Brand

1 - Diagnosis of acromegaly

AND

2 - Prescribed by, or in consultation with, an Endocrinologist

SOMAVERT PEGVISOMANT FOR INJ 20 MG (AS PROTEIN)

AND

- 3 One of the following:
 - **3.1** All of the following:
 - 3.1.1 Inadequate response to, or not a candidate for, surgical correction

AND

3.1.2 Trial and failure, contraindication, or intolerance to somatostatin therapy

OR

3.2 Continuation of prior therapy with pegvisomant, verified by paid claims or medical records (e.g. chart notes)

Product Name: Somavert				
Approval Length		12/31/2039		
Guideline Type		Prior Authorization-All plans except IL and MN		
Product Name	Generic Name		GPI	Brand/Generic
SOMAVERT	PEGVISOMANT FOR INJ 10 MG (AS PROTEIN)		30180060002120	Brand
SOMAVERT	PEGVISOMANT FOR INJ 15 MG (AS PROTEIN)		30180060002130	Brand

Brand

30180060002140

SOMAVERT	PEGVISOMANT FOR INJ 25 MG (AS PROTEIN)	30180060002150	Brand
SOMAVERT	PEGVISOMANT FOR INJ 30 MG (AS PROTEIN)	30180060002160	Brand

1 - Diagnosis of acromegaly

AND

2 - Prescribed by, or in consultation with, an Endocrinologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Inadequate response to, or not a candidate for, surgical correction

AND

3.1.2 Trial and failure, contraindication, or intolerance to somatostatin therapy

OR

3.2 Continuation of prior therapy with pegvisomant, verified by paid claims or medical records (e.g. chart notes)

2. Revision History

Date	Notes
3/27/2024	Guideline Update

Standalone Personal Continuous Glucose Monitors (CG					

Prior Authorization Guideline

Guideline ID	GL-143341
Guideline Name	Standalone Personal Continuous Glucose Monitors (CGM)
Formulary	Quartz

Guideline Note:

Effective Date:	2/23/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

1. Criteria

Product Name: Freestyle Libre 2, Freestyle Libre 3				
Approval Length		12/31/2039		
Guideline Type		Prior Authorization - All Plans except IL and MN Plans		
Product Name	Ge	eneric Name	GPI	Brand/Generic
FREESTYLE LIBRE 2/READER/FLASH GLUCOSE MONITORING SYSTEM *CONTINUOUS BLOOD GLUCOSE SYSTEM RECEIVER*** *CONTINUOUS BLOOD GLUCOSE SYSTEM *CONTINUOUS BLOOD GLUCOSE SYSTEM		97202012026200	Brand	

FREESTYLE LIBRE 2/SENSOR/FLASH GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM SENSOR***	97202012046300	Brand
FREESTYLE LIBRE 3/SENSOR/GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM SENSOR***	97202012046300	Brand

1 - Trial and failure or intolerance to a Dexcom product

Notes	*If patent meets criteria approve all CGM components at NDC list "CG MABBOTT"
	Persons with insurance coverage of a formulary CGM may upgrade to the newer formulary model upon request (e.g. authorization for Freest yle Libre 2 and requesting Freestyle Libre 3)

Product Name: Freestyle Libre 2, Freestyle Libre 3		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
FREESTYLE LIBRE 2/READER/FLASH GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM RECEIVER***	97202012026200	Brand
FREESTYLE LIBRE 2/SENSOR/FLASH GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM SENSOR***	97202012046300	Brand
FREESTYLE LIBRE 3/SENSOR/GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM SENSOR***	97202012046300	Brand

Approval Criteria

1 - Trial and failure or intolerance to a Dexcom product

Notes	*If patent meets criteria please approve all CGM components at NDC I ist "CGMABBOTT"
	Persons with insurance coverage of a formulary CGM may upgrade to the newer formulary model upon request (e.g. authorization for Freest yle Libre 2 and requesting Freestyle Libre 3)

Product Name: Freestyle Libre 2, Freestyle Libre 3	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
FREESTYLE LIBRE 2/READER/FLASH GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM RECEIVER***	97202012026200	Brand
FREESTYLE LIBRE 2/SENSOR/FLASH GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM SENSOR***	97202012046300	Brand
FREESTYLE LIBRE 3/SENSOR/GLUCOSE MONITORING SYSTEM	*CONTINUOUS BLOOD GLUCOSE SYSTEM SENSOR***	97202012046300	Brand

1 - Submission of medical records (e.g., chart notes) documenting regular use of the device (average of at least 5 days per week)

Notes	*If patent meets criteria please approve all CGM components at NDC I
	ist "CGMABBOTT"
	Persons with insurance coverage of a formulary CGM may upgrade to
	the newer formulary model upon request (e.g. authorization for Freest
	yle Libre 2 and requesting Freestyle Libre 3)

2. Revision History

Date	Notes

2/23/2024	Remove Dexcom from criteria, removal of most requirements for Fre estyle libre

State Mandate Reference Docum	nent
(a) bibliographic bibliog, bibliog, the most desired a rate of the bibliographic provide articles.	

Prior Authorization Guideline

Guideline ID	GL-137462
Guideline Name	State Mandate Reference Document
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Guideline Type Administrative				
Product Name	Gener	ic Name	GPI	Brand/Generic
Arkansas				
California				
Connecticut				
Georgia				
Indiana				
Kentucky				
Maryland				
New York				
West Virginia				

State		
Mandate		
Colorado		
Delaware		
lowa		
Illinois		
Louisiana		
Maine		
Minnesota		
New Mexico		
North Dakota		
Oklahoma		
Pennsylvania		
South Dakota		
Texas		
Virginia		
Wisconsin		
Florida		
Massachusetts		

- **1** The following mandates apply to Illinois:
- **1.1** Effective 1/1/2018, step therapy requirements are deemed met if the provider submits medical records confirming the patient is currently stabilized on the requested medication for the medical condition under consideration.

OR

1.2 Effective 1/1/2019, any clinical criteria component involving a trial/failure requirement are deemed met if the prescription drug is used to treat the patient's stage four advanced metastatic cancer and treatment is consistent with the U.S. Food and Drug Administration-approved indication or the National Comprehensive Cancer Network Drugs & Biologics Compendium indication for the treatment of stage four advanced metastatic cancer.

OR

1.3 Effective 6/9/2023, all clinical criteria are deemed met for intravenous immunoglobulin (IVIg) therapy when the medication is being used for a diagnosis of pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) or pediatric acute onset neuropsychiatric syndrome (PANS).

OR

2 - For lowa, (effective 1/1/2018), when the provider confirms a patient has previously received either a documented step one prescription drug or submits medical records documenting another prescription drug was received that has the same mechanism of action as the documented step one prescription drug, and the prescription drug was discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event, the patient will not be required to try any other alternatives with the same mechanism of action. Where documented step one prescription drugs are deemed met due to this process, all documented step one prescription drugs with the same mechanism of action will count towards the number of alternatives to be tried/failed. If step through other prescription drugs with a different mechanism of action is still required, the patient must meet the additional criteria. Step therapy requirements are also deemed met if the provider submits medical records confirming that the patient is currently stabilized on the requested medication for the medical condition under consideration. Note: Samples and drugs obtained through coupon cards may not count as sufficient experience with the prescribed medication to be considered stable on the medication.

OR

3 - For Minnesota, (effective 1/1/2020), any clinical criteria component involving a trial/failure requirement are deemed met if the prescription drug is used to treat the patient's stage four advanced metastatic cancer, or an associated condition, and treatment is consistent with the U.S. Food and Drug Administration-approved indication or the National Comprehensive Cancer Network Drugs & Biologics Compendium indication for the treatment of stage four advanced metastatic cancer.

OR

4 - For Wisconsin, (effective 11/1/2019), any clinical criteria component involving a trial/failure requirement are deemed met when the provider confirms a patient has previously received either a documented step one prescription drug or submits medical records documenting another prescription drug was received that has the same mechanism of action as the

documented step one prescription drug, and the prescription drug was discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event, the patient will not be required to try any other alternatives within the same pharmacological class or with the same mechanism of action. Where documented step one prescription drugs are deemed met due to this process, all documented step one prescription drugs with the same mechanism of action will count towards the number of alternatives to be tried/failed. If step through other prescription drugs with a different mechanism of action is still required, the patient must meet the additional criteria. Any clinical criteria component involving a trial/failure requirement are also deemed met if the provider submits medical records confirming that the patient is currently stabilized on the requested medication for the medical condition under consideration, or if submitted justification and clinical documentation support that the required step one prescription drug is expected to be ineffective.

2. Background

Benefit/Coverage/Program Information

Background:

This document serves as a reference for changes requested to pharmacy utilization management programs based on state mandates. This includes but is not limited to step therapy, prior authorization regulations, supply limits, first line trial duration limitations, and pain therapy/end of life regulations.

Additional Clinical Rules:

Applicable clinical programs will apply.

3. Revision History

Date	Notes
12/7/2023	Updated to only include applicable states: MN, IL, IA, WI

Stelara (Ustekinumab)
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Prior Authorization Guideline

Guideline ID	GL-144939
Guideline Name	Stelara (Ustekinumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/20/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Stelara SC					
Diagnosis		Plaque Psoriasis			
Approval Length		12/31/2039*			
Guideline Type Prior Authorization - All Plans except IL and MN Plans		3			
Product Name	Generic Na	me	GPI	Brand/Generic	
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML		9025058500E520	Brand	
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML		9025058500E540	Brand	
STELARA	USTEKINUMAB INJ 45 MG/0.5ML		90250585002020	Brand	

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** ONE of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of ≥ 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g., nails, scalp, genitals, or intertriginous areas

AND

3.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

OR

3.2 Continuation of prior therapy with ustekinumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start
	date:
	First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
	30 days
	Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Stelara SC		
Diagnosis	Plaque Psoriasis	
Approval Length	12 months*	
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** ONE of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of ≥ 3%
 - Debilitating palmer/plantar psoriasis or other vulnerable areas that are difficult to treat (e.g., nails, scalp, genitals, or intertriginous areas

AND

3.1.2 Trial and failure, contraindication, or intolerance to topical treatment (e.g. topical corticosteroids, calcipotriene, retinoids)

	OR
3.2 Continuat records (e.g. c	ion of prior therapy with ustekinumab, verified by paid claims or medical hart notes)
Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for 30 days

Second PA: Approve at GPI 10 through 12/31/2039

Product Name: Stelara SC		
Diagnosis	Psoriatic Arthritis (PsA)	
Approval Length	12/31/2039*	
Guideline Type	Prior Authorization - All Plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

3 - One of the following:

- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

3.2 Continuation of prior therapy with ustekinumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same date: First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days	
	30 days Second PA: Approve at GPI 10 through 12/31/2039	

Product Name: Stelara SC		
Diagnosis	Psoriatic Arthritis (PsA)	
Approval Length	12 months*	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

OR

3.2 Continuation of prior therapy with ustekinumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
	30 days Second PA: Approve at GPI 10 for 12 months

Product Name: Stelara SC	
Diagnosis Moderate to Severely Active Crohn's Disease (CD)	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

1 - Diagnosis of moderate to severely active Crohn's Disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** Documentation that the patient has been established on therapy on the medical benefit and one of the following:
 - **3.1.1** Patient is considered high-risk based on at least ONE of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease
 - Extraintestinal manifestations of inflammation (i.e. uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy, etc.)

OR

- **3.1.2** Both of the following:
- **3.1.2.1** Patient is considered low-risk

AND

- **3.1.2.2** At least ONE of the following:
- Intolerance/contraindication to 1 conventional therapy (ex. azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with 1 conventional therapy

- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

OR

3.2 Continuation of prior therapy with ustekinumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Stelara SC	
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active Crohn's Disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** Documentation that the patient has been established on therapy on the medical benefit and one of the following:
 - **3.1.1** Patient is considered high-risk based on at least ONE of the following characteristics:

- Age less than 30 years at diagnosis
- Extensive anatomic involvement
- Perianal and/or severe rectal disease
- Deep ulcers
- Prior surgical resection
- Stricturing and/or penetrating behavior
- Fistulizing disease
- Extraintestinal manifestations of inflammation (i.e. uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy, etc.)

OR

- **3.1.2** Both of the following:
- **3.1.2.1** Patient is considered low-risk

AND

3.1.2.2 At least ONE of the following:

- Intolerance/contraindication to 1 conventional therapy (ex. azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with 1 conventional therapy
- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

OR

3.2 Continuation of prior therapy with ustekinumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Stelara SC				
Diagnosis		Moderate to Severely Active Ulcerative Colitis (UC)		
Approval Ler	ngth	12 month(s)		
Guideline Ty	/pe	Prior Authorization - IL and MN Plans		
Product G Name	Generic Na	me	GPI	Brand/Generic

STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** Patient is considered high-risk based on ONE of the following characteristics:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High CRP and ESR
 - Steroid-requiring disease
 - History of hospitalization
 - C. difficile infection
 - CMV infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

AND

3.1.3 Prescriber attests patient has been established on therapy with ustekinumab for ulcerative colitis through the medical benefit

OR

3.2 Continuation of prior therapy with ustekinumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Stelara SC	
Diagnosis Moderate to Severely Active Ulcerative Colitis (UC)	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** Patient is considered high-risk based on ONE of the following characteristics:

- Extensive colitis
- Deep ulcers
- Age less than 40 years
- High CRP and ESR
- Steroid-requiring disease
- History of hospitalization
- C. difficile infection
- CMV infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

AND

3.1.3 Prescriber attests patient has been established on therapy with ustekinumab for ulcerative colitis through the medical benefit

OR

3.2 Continuation of prior therapy with ustekinumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Stelara SC	
Approval Length	12 month(s)
Guideline Type	Quantity Limit

Product Name	Generic Name	GPI	Brand/Generic
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 45 MG/0.5ML	9025058500E520	Brand
STELARA	USTEKINUMAB SOLN PREFILLED SYRINGE 90 MG/ML	9025058500E540	Brand
STELARA	USTEKINUMAB INJ 45 MG/0.5ML	90250585002020	Brand

Approval Criteria

1 - ALL of the following:
1.1 For members with diagnoses of Ulcerative Colitis (UC) or Crohn's Disease (CD) requesting reduced interval or increased dose (dose other than 90mg, interval less than every 8 weeks), ALL of the following:
1.1.1 Failure of a two-month trial of every 8-week dosing regimen after completion of induction dosing regimen
AND
1.1.2 Based on subtherapeutic drug concentrations and absence (or low levels) of drug antibodies
AND
1.1.3 Provision of published literature supporting dose increase and/or frequency
OR
1.2 ALL of the following:
1.2.1 For members with diagnoses of Psoriatic Arthritis (PsA) or Plaque Psoriasis (PP)
AND
1.2.2 Failure of an adherent 3-month trial of standard maintenance dosing with concomitant methotrexate (unless contraindicated or not appropriate for indication being requested)
AND
1.2.3 Provision of published literature supporting efficacy and safety of dosing regimen

OR

2 - Continuation of previous therapy with ustekinumab with a quantity exception, confirmed by paid claims or medical records (e.g. chart notes).

2. Revision History

Date	Notes
4/15/2024	Updated guideline

Strensiq (asfotase alfa)				
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Prior Authorization Guideline

Guideline ID	GL-133238
Guideline Name	Strensiq (asfotase alfa)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Strensiq	
Approval Length	12 month(s)
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 18 MG/0.45ML	30905610002020	Brand
STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 28 MG/0.7ML	30905610002030	Brand
STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 40 MG/ML	30905610002040	Brand
STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 80 MG/0.8ML	30905610002050	Brand

- **1** Diagnosis of perinatal, infantile, or juvenile-onset hypophosphatasia (HPP) with submission of medical records (e.g., chart notes) of one of the following symptom onset by age 6 months:
 - **1.1** Both of the following:
 - Serum alkaline phosphatase (ALP) levels below the age/gender-adjusted normal range
 - Elevated tissue non-specific alkaline phosphatase (TNSALP) substrate (e.g. serum pyridoxal 5'-phosphate (PLP) level, serum or urine phosphoethanolamine (PEA) level, or urinary inorganic pyrophosphate level)

OR

1.2 Documentation of TNSALP gene mutation by ALPL genomic DNA testing

AND

2 - Prescribed by or in consultation with an endocrinologist or other specialist in the treatment of inborn errors of metabolism

AND

3 - Submission of medical records (e.g., chart notes) documenting radiographic evidence supporting the diagnosis (e.g. infantile rickets, craniosynotosis, non-traumatic fractures, osteoporosis or low bone mineral content for age, etc.)

Product Name: Strensiq	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 18 MG/0.45ML	30905610002020	Brand
STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 28 MG/0.7ML	30905610002030	Brand

STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 40 MG/ML	30905610002040	Brand
STRENSIQ	ASFOTASE ALFA SUBCUTANEOUS INJ 80 MG/0.8ML	30905610002050	Brand

1 - Submission of medical records (e.g., chart notes) within the past 12 months documenting objective improvements in skeletal quality and labs from baseline such as improvement in respiratory status, improved growth, improved radiographic findings, or decrease in TNSALP substrate levels

2. Revision History

Date	Notes
9/24/2023	New Program

Sunosi (solriamfetol)			
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Prior Authorization Guideline

Guideline ID	GL-144951	
Guideline Name	Sunosi (solriamfetol)	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	9/18/2019
P&T Revision Date:	7/18/2023

1. Criteria

Approval Criteria

Product Name: Sunosi					
Approval Length		12/31/2039			
Guideline Type		Prior Authorization - All plans except IL and MN Plans			
Product Name	Generic Name		GPI	Brand/Generic	
SUNOSI	SOLRIAMFE	TOL HCL TAB 75 MG (BASE EQUIV)	61370070200320	Brand	
SUNOSI	SOLRIAMFE	TOL HCL TAB 150 MG (BASE EQUIV)	61370070200340	Brand	

1 - One of the following:
1.1 Diagnosis of narcolepsy
OR
1.2 Diagnosis of excessive daytime sleepiness in narcolepsy
OR
1.3 All of the following:
 Diagnosis of obstructive sleep apnea (OSA) Current or prior treatment of the underlying obstruction (e.g. continuous positive airway pressure [CPAP], mandibular advancement device or surgical intervention, etc.) If using CPAP, it will be used concomitantly with solriamfetol
AND
2 - Prescribed by or in consultation with a Sleep Specialist, Neurologist or Psychiatrist
AND
3 - Member is 18 years of age or older
AND
4 - One of the following:
4.1 Inadequate clinical response after a 3-month trial and failure, contraindication or intolerance to one other first line alternative (e.g., modafinil, armodafinil)
OR
4.2 Continuation of prior therapy with solfiamfetol, verified by paid claims, medical records (e.g. chart notes), or provider attestation.

Product Name: Sunosi		
Approval Length 12 month(s)		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
SUNOSI	SOLRIAMFETOL HCL TAB 75 MG (BASE EQUIV)	61370070200320	Brand
SUNOSI	SOLRIAMFETOL HCL TAB 150 MG (BASE EQUIV)	61370070200340	Brand

- **1** One of the following:
- **1.1** Diagnosis of narcolepsy

OR

1.2 Diagnosis of excessive daytime sleepiness in narcolepsy

OR

- **1.3** All of the following:
 - Diagnosis of obstructive sleep apnea (OSA)
 - Current or prior treatment of the underlying obstruction (e.g. continuous positive airway pressure [CPAP], mandibular advancement device or surgical intervention, etc.)
 - If using CPAP, it will be used concomitantly with solriamfetol

AND

2 - Prescribed by or in consultation with a Sleep Specialist, Neurologist or Psychiatrist

AND

3 - Member is 18 years of age or older

- 4 One of the following:
- **4.1** Inadequate clinical response after a 3-month trial and failure, contraindication or intolerance to one other first line alternative (e.g., modafinil, armodafinil)

OR

4.2 Continuation of prior therapy with solfiamfetol, verified by paid claims, medical records (e.g. chart notes), or provider attestation.

Date	Notes
3/27/2024	Updated Guideline

5	Sympazan (Clobazam)				
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Guideline ID	GL-129121
Guideline Name	Sympazan (Clobazam)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Sympazan	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
SYMPAZAN	CLOBAZAM ORAL FILM 5 MG	72100007008205	Brand
SYMPAZAN	CLOBAZAM ORAL FILM 10 MG	72100007008210	Brand
SYMPAZAN	CLOBAZAM ORAL FILM 20 MG	72100007008220	Brand

Approval Criteria

1 - Person with a diagnosis of Lennox-Gastaut syndrome with continued seizure activity despite adequate trial and failure, contraindication or intolerance to at least two preferred antiepileptic drugs (e.g., levetiracetam, lamotrigine)

AND

2 - A trial of generic clobazam (tablets and solution) was not tolerated due to physical inability to swallow

Product Name: Sympazan	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
SYMPAZAN	CLOBAZAM ORAL FILM 5 MG	72100007008205	Brand
SYMPAZAN	CLOBAZAM ORAL FILM 10 MG	72100007008210	Brand
SYMPAZAN	CLOBAZAM ORAL FILM 20 MG	72100007008220	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Sympazan		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
SYMPAZAN	CLOBAZAM ORAL FILM 5 MG	72100007008205	Brand
SYMPAZAN	CLOBAZAM ORAL FILM 10 MG	72100007008210	Brand
SYMPAZAN	CLOBAZAM ORAL FILM 20 MG	72100007008220	Brand

1 - Member with a diagnosis of Lennox-Gastaut syndrome with continued seizure activity despite adequate trial and failure, contraindication or intolerance to at least two preferred antiepileptic drugs (e.g., levetiracetam, lamotrigine)

AND

2 - A trial of generic clobazam (tablets and solution) was not tolerated due to physical inability to swallow

Date	Notes
9/11/2023	New program

Systemic Lupus Erythematosus (SLE) Treatmer			

Guideline ID	GL-144878	
Guideline Name	Systemic Lupus Erythematosus (SLE) Treatments	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	8/16/2017
P&T Revision Date:	7/18/2023

1. Criteria

Approval Criteria

Product Name: Benlysta SC				
Approval Length		12 month(s)*		
Guideline Type Prior Authorization				
Product Name	Generic Name GPI Brand/Gene		Brand/Generic	
BENLYSTA	BELIMUMAB SUBCUTANEOUS SOLUTION AUTO- INJECTOR 200 MG/ML		9942201500D520	Brand
BENLYSTA	BELIMUMAB SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 200 MG/ML		9942201500E520	Brand

- **1** All of the following:
 - Diagnosis of Systemic Lupus Erythematosus (SLE) with or without lupus nephritis
 - Member does not have severe central nervous system lupus

2 - Prescribed by or in consultation with a rheumatologist or other specialist in the treatment of SLE

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to ALL of the following:
 - Hydroxychloroquine
 - Nonsteroidal anti-inflammatories (NSAIDs) (e.g., ibuprofen, naproxen)
 - A steroid-sparing immunosuppressive (e.g., azathioprine, methotrexate)
 - A short course of oral steroids

OR

3.2 Continuation of prior therapy with belimumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.29 (8 every 28 days) for
	30 days
	Second PA: Approve at GPI 10 for 12 months

Date	Notes
3/27/2024	Guideline Update.

ladalafil	for Benign Pr	ostate Hyperplas
The detail image-current techniques. The file may take a more more	of waters, a delete, builty death to probe for according and value.	

Guideline ID	GL-144565
Guideline Name	Tadalafil for Benign Prostate Hyperplasia
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Product Name: Generic Tadalafil 2.5mg, 5mg		
Approval Length 12 month(s)		
Therapy Stage	Therapy Stage Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
TADALAFIL	TADALAFIL TAB 2.5 MG	40304080000302	Generic
TADALAFIL	TADALAFIL TAB 5 MG	40304080000305	Generic

Approval Criteria

1 - Diagnosis of benign prostatic hyperplasia (BPH)

Product Name: Generic Tadalafil 2.5mg, 5mg	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
TADALAFIL	TADALAFIL TAB 2.5 MG	40304080000302	Generic
TADALAFIL	TADALAFIL TAB 5 MG	40304080000305	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Generic Tadalafil 2.5mg, 5mg	
Approval Length	12/31/2039
Guideline Type Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
TADALAFIL	TADALAFIL TAB 2.5 MG	40304080000302	Generic
TADALAFIL	TADALAFIL TAB 5 MG	40304080000305	Generic

Approval Criteria

1 - Diagnosis of benign prostatic hyperplasia (BPH)

Date	Notes
3/18/2024	Updated product name

Tavalisse (Fostamatinib)		
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Guideline ID	GL-128905
Guideline Name	Tavalisse (Fostamatinib)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Tavalisse		
Approval Length 12/31/2039		
Guideline Type Prior authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
TAVALISSE	FOSTAMATINIB DISODIUM TAB 100 MG (BASE EQUIVALENT)	85756040100310	Brand
TAVALISSE	FOSTAMATINIB DISODIUM TAB 150 MG (BASE EQUIVALENT)	85756040100320	Brand

Approval Criteria

1 - Diagnosis of chronic immune thrombocytopenia (ITP)

2 - Member's platelet count < 50,000/mL

AND

3 - Trial and failure of 2 prior ITP therapies (e.g. corticosteroids, rituximab, azathioprine, danazol, splenectomy, or eltrombopag)

AND

4 - Prescribed by, or in consultation with hematology

Product Name: Tavalisse		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TAVALISSE	FOSTAMATINIB DISODIUM TAB 100 MG (BASE EQUIVALENT)	85756040100310	Brand
TAVALISSE	FOSTAMATINIB DISODIUM TAB 150 MG (BASE EQUIVALENT)	85756040100320	Brand

Approval Criteria

1 - Diagnosis of chronic immune thrombocytopenia (ITP)

AND

2 - Member's platelet count < 50,000/mL

3 - Trial and failure of 2 prior ITP therapies (e.g. corticosteroids, rituximab, azathioprine, danazol, splenectomy, or eltrombopag)

AND

4 - Prescribed by, or in consultation with hematology

Product Name: Tavalisse		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization for IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TAVALISSE	FOSTAMATINIB DISODIUM TAB 100 MG (BASE EQUIVALENT)	85756040100310	Brand
TAVALISSE	FOSTAMATINIB DISODIUM TAB 150 MG (BASE EQUIVALENT)	85756040100320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
9/7/2023	New Program

1	Tegse	di (inc	terse	n)		
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Guideline ID	GL-131604
Guideline Name	Tegsedi (inotersen)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies.

1. Criteria

Product Na	Product Name: Tegsedi			
Diagnosis		Neuropathy due to hereditary transthyretin (hATTR) amyloidosis		
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization		
Product Generic Name Name		GPI	Brand/Generic	

TEGSEDI INOTERSEN SOD SUBCUTANEOUS MG/1.5ML (BASE EQ)	PREF SYR 284 6270104010E520	Brand
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1 - Diagnosis of neuropathy due to hereditary transthyretin (hATTR) amyloidosis with documentation of TTR gene mutation and biopsy proven amyloid deposits

AND

2 - Prescribed by, or in consultation with, a Neurologist, Cardiologist, or other expert in hereditary transthyretin-mediated amyloidosis (hATTR)

AND

3 - Member is 18 years of age or older

AND

4 - Drug is not being used in combination with another TTR-lowering agent (e.g., patisiran, vutrisiran)

AND

5 - Drug is not being used in combination with a TTR-stabilizing agent (e.g., diflunisal, tafamidis, tafamidis meglumine)

Product Name: Tegsedi				
Diagnosis Continuation of Coverage if New to Plan				
Approval Length		12 month(s)		
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization		
Product Generic Name Name		me	GPI	Brand/Generic

TEGSEDI INOTERSEN SOD SUBCUTANEOUS MG/1.5ML (BASE EQ)	PREF SYR 284 6270104010E520	Brand
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1 - Diagnosis of neuropathy due to hereditary transthyretin (hATTR) amyloidosis with documentation of TTR gene mutation and biopsy proven amyloid deposits

AND

2 - Prescribed by, or in consultation with, a Neurologist, Cardiologist, or other expert in hereditary transthyretin-mediated amyloidosis (hATTR)

AND

3 - Member is 18 years of age or older

AND

4 - Drug is not being used in combination with another TTR-lowering agent (e.g., patisiran, vutrisiran)

AND

5 - Drug is not being used in combination with a TTR-stabilizing agent (e.g., diflunisal, tafamidis, tafamidis meglumine)

AND

6 - The prescriber must provide clinical documentation of the member's initial response to therapy (e.g. clinical manifestation stability/improvement)

Product Name: Tegsed	i
Diagnosis	Neuropathy due to hereditary transthyretin (hATTR) amyloidosis

Approval Length	12/31/2039
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - All Plans except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TEGSEDI	INOTERSEN SOD SUBCUTANEOUS PREF SYR 284 MG/1.5ML (BASE EQ)	6270104010E520	Brand

1 - Submission of medical records (e.g., chart notes) documenting response to therapy or documentation of clinical stability for the previous 12 months

Product Name: Tegsedi	
Diagnosis	Neuropathy due to hereditary transthyretin (hATTR) amyloidosis
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TEGSEDI	INOTERSEN SOD SUBCUTANEOUS PREF SYR 284 MG/1.5ML (BASE EQ)	6270104010E520	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting response to therapy or documentation of clinical stability for the previous 12 months

Date	Notes
8/24/2023	2024 New Implementation

T	Testosterone		
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Guideline ID	GL-144887
Guideline Name	Testosterone
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	9/20/2017
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Preferred Testosterone Products: generic testosterone 1% gel, generic testosterone 1.62% gel, generic testosterone cypionate, generic testosterone enanthate				
Approval Length		12 month(s)		
Guideline Type Prior Authorization - IL and MN Plans				
Product Name	Gene	ric Name	GPI	Brand/Generic
TESTOSTERONE	TESTO (1.62%	STERONE TD GEL 20.25 MG/ACT)	23100030004050	Generic
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%) 231000		23100030004050	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 50 MG/5GM (1%) 23100030004030 Generic		Generic	
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 12.5 MG/ACT (1%) 23100030004040 Generic			

TESTOSTERONE	TESTOSTERONE TD GEL 25 MG/2.5GM (1%)	23100030004025	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 40.5 MG/2.5GM (1.62%)	23100030004047	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 20.25 MG/1.25GM (1.62%)	23100030004044	Generic
TESTOSTERONE ENANTHATE	TESTOSTERONE ENANTHATE IM INJ IN OIL 200 MG/ML	23100030202010	Generic
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 100 MG/ML	23100030102010	Generic
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 200 MG/ML	23100030102015	Generic

- **1** One of the following:
- 1.1 Diagnosis of gender dysphoria, transsexualism, or other gender identity diagnosis

OR

- **1.2** All of the following:
- **1.2.1** Submission of medical records (e.g., chart notes) demonstrating androgen deficiency** in one of the following diagnoses:
 - Primary or secondary hypogonadism
 - Mixed hypogonadism

AND

1.2.2 Submission of medical records (e.g. chart notes) documenting two morning testosterone levels (drawn between 7:00-10:00 a.m. or within 3 hours of waking for shift workers) below the lower limit of normal

AND

1.2.3 Submission of medical records (e.g., chart notes) documenting symptoms due to low testosterone other than decreased libido and/or other sexual dysfunction

\sim	

1.3 Continuation of prior therapy with testosterone, verified by paid claims or medical records (e.g. chart notes)

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s v co ore vo **,4 ve of by tes	Continuation of therapy/coverage criteria will not be applied to person who were not previously approved for overage but whose therapy was initiated using a manufacturer-sponsed free drug program, provider samples, and/or ouchers. Androgen deficiency is defined as a fasting, morning testosterone letel (drawn between 7 and 10 AM or within 3 hours waking for shift workers) below the lower limit of normal as defined the laboratory reference range. A single low stosterone is not diagnostic for androgen deficiency and must be contirmed with a second fasting, morning stosterone level.

Product Name: Non-Preferred Testosterone Products: Brand Androderm, generic testosterone topical solution	
Approval Length	12 month(s)
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
ANDRODERM	TESTOSTERONE TD PATCH 24HR 2 MG/24HR	23100030008503	Brand
ANDRODERM	TESTOSTERONE TD PATCH 24HR 4 MG/24HR	23100030008510	Brand

Approval Criteria

- 1 One of the following:
- 1.1 Diagnosis of gender dysphoria, transsexualism, or other gender identity diagnosis

OR

1.2 All of the following:

- **1.2.1** Submission of medical records (e.g., chart notes) demonstrating androgen deficiency** in one of the following diagnoses:
 - Primary or secondary hypogonadism
 - Mixed hypogonadism

1.2.2 Submission of medical records (e.g. chart notes) documenting two morning testosterone levels (drawn between 7:00 – 10:00 a.m. or within 3 hours of waking for shift workers) below the lower limit of normal

AND

1.2.3 Submission of medical records (e.g., chart notes) documenting symptoms due to low testosterone other than decreased libido and/or other sexual dysfunction

OR

1.3 Continuation of prior therapy with testosterone, verified by paid claims or medical records (e.g. chart notes)

AND

2 - Trial and failure, contraindication, or intolerance to at least one preferred testosterone option (with the same route of administration if available)

Notes	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage but whose therapy was initiated using a manufacturer-spons ored free drug program, provider samples, and/or vouchers. **Androgen deficiency is defined as a fasting, morning testosterone le vel (drawn between 7 and 10 AM or within 3 hours of waking for shift workers) below the lower limit of normal as defined by the laboratory reference range. A single low testosterone is not diagnostic for androgen deficiency and must be confirmed with a second fasting, morning testosterone level.

Product Name: Preferred Testosterone Products: generic testosterone 1% gel, generic testosterone 1.62% gel, generic testosterone cypionate, generic testosterone enanthate	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TESTOSTERONE	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Generic
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 20.25 MG/ACT (1.62%)	23100030004050	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 50 MG/5GM (1%)	23100030004030	Generic
TESTOSTERONE PUMP	TESTOSTERONE TD GEL 12.5 MG/ACT (1%)	23100030004040	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 25 MG/2.5GM (1%)	23100030004025	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 40.5 MG/2.5GM (1.62%)	23100030004047	Generic
TESTOSTERONE	TESTOSTERONE TD GEL 20.25 MG/1.25GM (1.62%)	23100030004044	Generic
TESTOSTERONE ENANTHATE	TESTOSTERONE ENANTHATE IM INJ IN OIL 200 MG/ML	23100030202010	Generic
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 100 MG/ML	23100030102010	Generic
TESTOSTERONE CYPIONATE	TESTOSTERONE CYPIONATE IM INJ IN OIL 200 MG/ML	23100030102015	Generic

- 1 One of the following:
- 1.1 Diagnosis of gender dysphoria, transsexualism, or other gender identity diagnosis

OR

- **1.2** All of the following:
- **1.2.1** Submission of medical records (e.g., chart notes) demonstrating androgen deficiency** in one of the following diagnoses:
 - Primary or secondary hypogonadism
 - Mixed hypogonadism

1.2.2 Submission of medical records (e.g. chart notes) documenting two morning testosterone levels (drawn between 7:00 – 10:00 a.m. or within 3 hours of waking for shift workers) below the lower limit of normal

AND

1.2.3 Submission of medical records (e.g., chart notes) documenting symptoms due to low testosterone other than decreased libido and/or other sexual dysfunction

OR

1.3 Continuation of prior therapy with testosterone, verified by paid claims or medical records (e.g. chart notes)

(e.g. chart notes)	
Notes	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage but whose therapy was initiated using a manufacturer-spons ored free drug program, provider samples, and/or vouchers. **Androgen deficiency is defined as a fasting, morning testosterone le vel (drawn between 7 and 10 AM or within 3 hours of waking for shift workers) below the lower limit of normal as defined by the laboratory reference range. A single low testosterone is not diagnostic for androgen deficiency and must be confirmed with a second fasting, morning testosterone level.

Product Name: Non-Preferred Testosterone Products: Brand Androderm, generic testosterone topical solution	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ANDRODERM	TESTOSTERONE TD PATCH 24HR 2 MG/24HR	23100030008503	Brand
ANDRODERM	TESTOSTERONE TD PATCH 24HR 4 MG/24HR	23100030008510	Brand

- **1** One of the following:
- 1.1 Diagnosis of gender dysphoria, transsexualism, or other gender identity diagnosis

OR

- **1.2** All of the following:
- **1.2.1** Submission of medical records (e.g., chart notes) demonstrating androgen deficiency** in one of the following diagnoses:
 - Primary or secondary hypogonadism
 - Mixed hypogonadism

AND

1.2.2 Submission of medical records (e.g. chart notes) documenting two morning testosterone levels (drawn between 7:00 – 10:00 a.m. or within 3 hours of waking for shift workers) below the lower limit of normal

AND

1.2.3 Submission of medical records (e.g., chart notes) documenting symptoms due to low testosterone other than decreased libido and/or other sexual dysfunction

OR

1.3 Continuation of prior therapy with testosterone, verified by paid claims or medical records (e.g. chart notes)

AND

2 - Trial and failure, contraindication, or intolerance to at least one preferred testosterone option (with the same route of administration if available)

Notes	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage but whose therapy was initiated using a manufacturer-spons ored free drug program, provider samples, and/or vouchers. **Androgen deficiency is defined as a fasting, morning testosterone le vel (drawn between 7 and 10 AM or within 3 hours of waking for shift workers) below the lower limit of normal as defined by the laboratory reference range. A single low testosterone is not diagnostic for androgen deficiency and must be co

Date	Notes
3/27/2024	Guideline Update.

Tezspire (tezepelumab)			
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Guideline ID	GL-144895
Guideline Name	Tezspire (tezepelumab)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	4/19/2023
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Tezspire					
Diagnosis		Eosinophilic Asthma			
Approval Length 12 month(s)					
Guideline Type Prior Authorization - IL and MN Plans					
Product Name	Generic Name		GPI	Brand/Generic	
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN AUTO- INJ 210 MG/1.91ML		4460807525D520	Brand	
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN PREF SYR 210 MG/1.91ML		4460807525E520	Brand	

Approval Criteria		
1 - Diagnosis of eosinophilic asthma		
AND		
AND		
2 - Prescribed by or in consultation with one of the following:		
AllergistImmunologistPulmonologist		
AND		
3 - Member is 12 years of age or older		
AND		
4 - One of the following:		
4.1 All of the following:		
4.1.1 All of the following:		
4.1.1.1 Submission of medical records (e.g., chart notes) documenting a blood eosinophil count of ≥ 150 cells/mm3		
AND		
AND		
4.1.1.2 All other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease have been ruled out		
AND		
4.1.1.3 One of the following:		

4.1.1.3.1 Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent $\ddagger \ge 3$ -month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

- **4.1.1.3.2** One of the following:
- **4.1.1.3.2.1** Member has an intolerance to medium to high dose inhaled corticosteroids (ICS) in combination with long-acting bronchodilator or leukotriene modifier

OR

- **4.1.1.3.2.2** Member has one or more of the following comorbid conditions that increase long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after evaluation by endocrine consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

4.1.1.4 Trial and failure, contraindication, or intolerance to at least two preferred self-administered biologic therapies for eosinophilic asthma (i.e. dupilumab, benralizumab, mepolizumab)

OR

4.2 Continuation of prior therapy with tezepelumab, verified by paid claims or medical records (e.g. chart notes)

Notes	‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.
	NOTE: II-5 inhibitor drugs in combination with omalizumab will be con sidered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms . Tezepelumab, in combination with other biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Name: Tezspire		
Diagnosis Eosinophilic Asthma		
Approval Length 12/31/2039		
Guideline Type Prior Authorization – All Plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN AUTO-INJ 210 MG/1.91ML	4460807525D520	Brand
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN PREF SYR 210 MG/1.91ML	4460807525E520	Brand

1 - Diagnosis of eosinophilic asthma

AND

- **2** Prescribed by or in consultation with one of the following:

 - Allergist Immunologist Pulmonologist

AND
3 - Member is 12 years of age or older
AND
4 - One of the following:
4.1 All of the following:
4.1.1 All of the following:
4.1.1.1 Submission of medical records (e.g., chart notes) documenting a blood eosinophil count of ≥ 150 cells/mm3
AND
4.1.1.2 All other causes of eosinophilia such as hypereosinophilic syndromes, neoplastic disease, or parasitic disease have been ruled out
AND
4.1.1.3 One of the following:
4.1.1.3.1 Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent ‡ ≥ 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier
OR
4.1.1.3.2 One of the following:
4.1.1.3.2.1 Member has an intolerance to medium to high dose inhaled corticosteroids (ICS) in combination with long-acting bronchodilator or leukotriene modifier

OR

- **4.1.1.3.2.2** Member has one or more of the following comorbid conditions that increase long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after evaluation by endocrine consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

4.1.1.4 Trial and failure, contraindication, or intolerance to at least two preferred self-administered biologic therapies for eosinophilic asthma (i.e. dupilumab, benralizumab, mepolizumab)

OR

4.2 Continuation of prior therapy with tezepelumab, verified by paid claims or medical records (e.g. chart notes)

Notes

‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.

NOTE: II-5 inhibitor drugs in combination with omalizumab will be con sidered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms . Tezepelumab, in combination with other biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).

*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Name: Tezspire

Diagnosis	Allergic Asthma
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN AUTO-INJ 210 MG/1.91ML	4460807525D520	Brand
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN PREF SYR 210 MG/1.91ML	4460807525E520	Brand

1 - Diagnosis of moderate-to-severe persistent allergic asthma as defined by Global Initiative for Asthma (GINA) Global Strategy for Asthma Management and Prevention Guidelines (Step 5)

AND

- **2** Prescribed by or in consultation with one of the following:
 - Allergist
 - Immunologist
 - Pulmonologist

AND

3 - Member is 12 years of age or older

AND

- 4 One of the following:
- **4.1** All of the following:
- **4.1.1** Serum IgE level ≥30 international units/mL

AND

4.1.2 Positive skin tests or in vitro reactivity to common aeroallergens (e.g. dust mites, pet dander, cockroaches, etc.)

AND

- **4.1.3** One of the following:
- **4.1.3.1** Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent ‡ ≥ 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

- **4.1.3.2** One of the following:
- **4.1.3.2.1** Member has an intolerance to medium to high dose inhaled corticosteroids (ICS) in combination with long-acting bronchodilator or leukotriene modifier

OR

- **4.1.3.2.2** Member has one or more of the following comorbid conditions that increase long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after evaluation by endocrine consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

4.1.4 Trial and failure, contraindication, or intolerance to at least one preferred self-administered biologic therapy for allergic asthma (i.e. omalizumab)

OR

4.2 Continuation of prior therapy with tezepelumab, verified by paid claims or medical

records (e.g. cha	art notes)
Notes	‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.
	NOTE: II-5 inhibitor drugs in combination with omalizumab will be considered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms. Tezepelumab, in combination with other biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Name: Tezspire		
Diagnosis Allergic Asthma		
Approval Length 12/31/2039		
Guideline Type Prior Authorization – All plans except IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN AUTO-INJ 210 MG/1.91ML	4460807525D520	Brand
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN PREF SYR 210 MG/1.91ML	4460807525E520	Brand

Approval Criteria

1 - Diagnosis of moderate-to-severe persistent allergic asthma as defined by Global Initiative for Asthma (GINA) Global Strategy for Asthma Management and Prevention Guidelines (Step

AND

- 2 Prescribed by or in consultation with one of the following:
 - Allergist

ImmunologistPulmonologist
AND
3 - Member is 12 years of age or older
AND
4 - One of the following:
4.1 All of the following:
4.1.1 Serum IgE level ≥30 international units/mL
AND
4.1.2 Positive skin tests or in vitro reactivity to common aeroallergens (e.g. dust mites, pet dander, cockroaches, etc.)
AND
4.1.3 One of the following:
4.1.3.1 Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent ‡ ≥ 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier
OR
4.1.3.2 One of the following:
4.1.3.2.1 Member has an intolerance to medium to high dose inhaled corticosteroids (ICS) in combination with long-acting bronchodilator or leukotriene modifier
OR

- **4.1.3.2.2** Member has one or more of the following comorbid conditions that increase long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after evaluation by endocrine consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

4.1.4 Trial and failure, contraindication, or intolerance to at least one preferred self-administered biologic therapy for allergic asthma (i.e. omalizumab)

OR

4.2 Continuation of prior therapy with tezepelumab, verified by paid claims or medical records (e.g. chart notes)

Notes	‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.
	NOTE: II-5 inhibitor drugs in combination with omalizumab will be con sidered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms . Tezepelumab, in combination with other biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Name: Tezspire		
Diagnosis	Severe Asthma	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization – All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN AUTO-INJ 210 MG/1.91ML	4460807525D520	Brand
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN PREF SYR 210 MG/1.91ML	4460807525E520	Brand

1 - Diagnosis of severe asthma as defined by GINA guidelines

AND

- 2 Prescribed by or in consultation with one of the following:
 - Allergist
 - Immunologist
 - Pulmonologist

AND

3 - Member is 12 years of age or older

AND

- 4 One of the following:
 - **4.1** All of the following:
 - **4.1.1** One of the following:
 - History of ≥ 2 asthma exacerbations requiring systemic corticosteroids within the past 12 months
 - One asthma exacerbation requiring hospitalization in the past 12 months

AND

4.1.2 Asthma is non-eosinophilic (example: blood eosinophil counts of less than 150 cells/µL

AND

4.1.3 Asthma is non-allergic (example: Serum IgE level less than 30 international units/mL or negative skin tests to common aeroallergens)

AND

- **4.1.4** One of the following:
- **4.1.4.1** Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent ‡ ≥ 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

- **4.1.4.2** One of the following:
- **4.1.4.2.1** Member has an intolerance to medium to high dose inhaled corticosteroids (ICS) in combination with long-acting bronchodilator or leukotriene modifier

OR

- **4.1.4.2.2** Member has one or more of the following comorbid conditions that increase long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after evaluation by endocrine consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

4.1.5 For oral corticosteroid dependent asthma (requiring daily oral steroids): trial and failure, contraindication, or intolerance to at least one preferred self-administered biologic therapy for corticosteroid dependent asthma (i.e. dupilumab)

OR

4.2 Continuation of prior therapy with tezepelumab, verified by paid claims or medical records (e.g. chart notes)

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Notes	‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.
	NOTE: II-5 inhibitor drugs in combination with omalizumab will be considered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms. Tezepelumab, in combination with other biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).
	*Continuation of therapy/coverage criteria will not be applied to person s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

Product Name: Tezspire	
Diagnosis	Severe Asthma
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN AUTO-INJ 210 MG/1.91ML	4460807525D520	Brand
TEZSPIRE	TEZEPELUMAB-EKKO SUBCUTANEOUS SOLN PREF SYR 210 MG/1.91ML	4460807525E520	Brand

Approval Criteria

1 - Diagnosis of severe asthma as defined by GINA guidelines

AND
2 - Prescribed by or in consultation with one of the following:
AllergistImmunologistPulmonologist
AND
3 - Member is 12 years of age or older
AND
4 - One of the following:
4.1 All of the following:
4.1.1 One of the following:
 History of ≥ 2 asthma exacerbations requiring systemic corticosteroids within the past 12 months One asthma exacerbation requiring hospitalization in the past 12 months
AND
4.1.2 Asthma is non-eosinophilic (example: blood eosinophil counts of less than 150 cells/μL
AND
4.1.3 Asthma is non-allergic (example: Serum IgE level less than 30 international units/mL or negative skin tests to common aeroallergens)
AND

4.1.4 One of the following:

4.1.4.1 Symptoms are not well controlled or poorly controlled (refer to Table 1 in background section) despite an adherent ‡ ≥ 3-month trial of medium to high-dose inhaled corticosteroids in combination with a long-acting bronchodilator, long-acting muscarinic antagonist, or leukotriene modifier

OR

4.1.4.2 One of the following:

4.1.4.2.1 Member has an intolerance to medium to high dose inhaled corticosteroids (ICS) in combination with long-acting bronchodilator or leukotriene modifier

OR

- **4.1.4.2.2** Member has one or more of the following comorbid conditions that increase long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients > 40 years of age
 - Glaucoma
 - Recurrent thrush
 - Dysphonia
 - Growth inhibition, after evaluation by endocrine consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

4.1.5 For oral corticosteroid dependent asthma (requiring daily oral steroids): trial and failure, contraindication, or intolerance to at least one preferred self-administered biologic therapy for corticosteroid dependent asthma (i.e. dupilumab)

OR

4.2 Continuation of prior therapy with tezepelumab, verified by paid claims or medical records (e.g. chart notes)

	‡Adherent treatment is defined as a medication possession ratio (MP R) ≥ 70% based on the previous 120 days of prescription claims.

NOTE: II-5 inhibitor drugs in combination with omalizumab will be con sidered on a case-by-case basis if each individual agent with combination high dose ICS/LABA did not control symptoms . Tezepelumab, in combination with other biologics, has not been studied and coverage is not allowed except in extenuating circumstances (applies to both eosinophilic or non-eosinophilic asthma populations).

*Continuation of therapy/coverage criteria will not be applied to person

*Continuation of therapy/coverage criteria will not be applied to persor s who were not previously approved for coverage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers.

2. Background

Benefit/Coverage/Program Information

Table 1. Outcome Measure values for uncontrolled asthma

Measure	Not Well Controlled	Very Poorly Controlled
Baseline symptoms (outside of exacerbation)	> 2 days/week	Throughout the day
Nighttime awakening	1-3 times/week	≥ 4 times/week
Interference with normal activity	Some limitation	Extremely limited
Short acting beta agonist use for symptom control	> 2 days/week	Several times per day
FEV1	60-80% predicted or personal best	< 60% predicted or personal best
Asthma exacerbations requiring oral steroids ≥ 2 times in the past year	Yes	Yes

Asthma Control Test	16-19	≤ 15	
(ACT)			

Date	Notes
4/9/2024	Guideline Update.

Thrombopoietin Receptor Agonists			
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Guideline ID	GL-144923
Guideline Name	Thrombopoietin Receptor Agonists
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	8/21/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Doptele	Product Name: Doptelet, Promacta	
Diagnosis	Thrombocytopenia in Patients with Chronic Immune Thrombocytopenia (ITP)	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
DOPTELET	AVATROMBOPAG MALEATE TAB 20 MG (BASE EQUIV)	82405010200320	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 12.5 MG (BASE EQUIV)	82405030100310	Brand

PROMACTA	ELTROMBOPAG OLAMINE TAB 25 MG (BASE EQUIV)	82405030100320	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 50 MG (BASE EQUIV)	82405030100330	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 75 MG (BASE EQUIV)	82405030100340	Brand
PROMACTA	ELTROMBOPAG OLAMINE POWDER PACK FOR SUSP 25 MG (BASE EQUIV)	82405030103020	Brand
PROMACTA	ELTROMBOPAG OLAMINE POWDER PACK FOR SUSP 12.5 MG (BASE EQ)	82405030103030	Brand

1 - Diagnosis of chronic ITP with a platelet count less than 50,000/mcL

AND

2 - Prescribed by or in consultation with a hematologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to at least TWO prior ITP therapies (e.g., corticosteroids, rituximab, azathioprine, danazol, or splenectomy)

OR

3.2 Continuation of prior therapy with the requested drug, verified by paid claims or medical records (e.g. chart notes)

Product Name: Doptelet				
Diagnosis		Thrombocytopenia in Patients with Chronic Liver Disease (CLD)		
Approval L	ength	12 month(s) with a fill count of one		
Guideline 7	Guideline Type Prior Authorization - IL and MN Plans Only			
Product Generic Name Name		GPI	Brand/Generic	

DOPTELET AVATROMBOPAG MALEATE TAB 20 MG (BASE EQUIV)	82405010200320	Brand
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1 - Diagnosis of liver cirrhosis with a platelet count less than 50,000/mcL

AND

2 - Prescribed by or in consultation with a hematologist or gastroenterologist

AND

3 - Member is scheduled for a procedure with a moderate to high bleeding risk within the next 14 days

Product Name: Doptelet		
Diagnosis Thrombocytopenia in Patients with Chronic Liver Disease (CLD)		
Approval Length 5 Day(s)		
Guideline Type Prior Authorization - All plans except IL and MN		

П	Product Name	Generic Name	GPI	Brand/Generic
	DOPTELET	AVATROMBOPAG MALEATE TAB 20 MG (BASE EQUIV)	82405010200320	Brand

Approval Criteria

1 - Diagnosis of liver cirrhosis with a platelet count less than 50,000/mcL

AND

2 - Prescribed by or in consultation with a hematologist or gastroenterologist

AND

3 - Member is scheduled for a procedure with a moderate to high bleeding risk within the next 14 days

Product Name: Mulpleta		
Diagnosis	Thrombocytopenia in Patients with Chronic Liver Disease (CLD)	
Approval Length 12 month(s) with a fill count of one		
Guideline Type Prior Authorization - IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
MULPLETA	LUSUTROMBOPAG TAB 3 MG	82405045000320	Brand

Approval Criteria

1 - Diagnosis of liver cirrhosis with a platelet count less than 50,000/mcL

AND

2 - Prescribed by or in consultation with a hematologist or gastroenterologist

AND

3 - Member is scheduled for a procedure with a moderate to high bleeding risk within the next 14 days

Product Na	Product Name: Mulpleta			
Diagnosis		Thrombocytopenia in Patients with Chronic Liver Disease (CLD)		
Approval L	ength	1 Time(s)		
Guideline 7	Guideline Type Prior Authorization - All plans except IL and MN			
Product Generic Name		GPI	Brand/Generic	

MULPLETA LUSUTROMBOPAG TAB 3 MG	82405045000320	Brand

1 - Diagnosis of liver cirrhosis with a platelet count less than 50,000/mcL

AND

2 - Prescribed by or in consultation with a hematologist or gastroenterologist

AND

3 - Member is scheduled for a procedure with a moderate to high bleeding risk within the next14 days

Product Name: Promacta	
Diagnosis Chronic Hepatitis C-Associated Thrombocytopenia	
Approval Length 12 month(s)	
Guideline Type Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
PROMACTA	ELTROMBOPAG OLAMINE TAB 12.5 MG (BASE EQUIV)	82405030100310	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 25 MG (BASE EQUIV)	82405030100320	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 50 MG (BASE EQUIV)	82405030100330	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 75 MG (BASE EQUIV)	82405030100340	Brand
PROMACTA	ELTROMBOPAG OLAMINE POWDER PACK FOR SUSP 25 MG (BASE EQUIV)	82405030103020	Brand
PROMACTA	ELTROMBOPAG OLAMINE POWDER PACK FOR SUSP 12.5 MG (BASE EQ)	82405030103030	Brand

Approval Criteria

1 - Diagnosis of chronic hepatitis C virus (HCV) undergoing treatment with pegylated interferon/ribavirin

AND

2 - Prescribed by or in consultation with a hematologist, gastroenterologist, or infectious disease specialist

AND

- 3 One of the following:
- **3.1** Platelet count is less than 75,000/mcL

OR

3.2 Continuation of prior therapy with eltrombopag, verified by paid claims or medical records (e.g. chart notes)

Product Name: Promacta		
Diagnosis	Aplastic Anemia	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
PROMACTA	ELTROMBOPAG OLAMINE TAB 12.5 MG (BASE EQUIV)	82405030100310	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 25 MG (BASE EQUIV)	82405030100320	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 50 MG (BASE EQUIV)	82405030100330	Brand
PROMACTA	ELTROMBOPAG OLAMINE TAB 75 MG (BASE EQUIV)	82405030100340	Brand
PROMACTA	ELTROMBOPAG OLAMINE POWDER PACK FOR SUSP 25 MG (BASE EQUIV)	82405030103020	Brand
PROMACTA	ELTROMBOPAG OLAMINE POWDER PACK FOR SUSP 12.5 MG (BASE EQ)	82405030103030	Brand

Approval Criteria

1 - Diagnosis of severe aplastic anemia

AND

2 - Prescribed by or in consultation with a hematologist

AND

- 3 One of the following:
- **3.1** Trial and failure, contraindication, or intolerance to at least one immunosuppressive therapy (e.g., glucocorticoids, cyclosporine)

OR

3.2 Continuation of prior therapy with eltrombopag, verified by paid claims or medical records (e.g. chart notes)

Date	Notes
3/27/2024	Guideline Update.

Т	Tiglutik (riluzole)			
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Guideline ID	GL-131424
Guideline Name	Tiglutik (riluzole)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Tiglutik		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
TIGLUTIK	RILUZOLE SUSP 50 MG/10ML	74503070001820	Brand

Approval Criteria

1 - Diagnosis of amyotrophic lateral sclerosis (ALS)

AND

2 - A trial of generic riluzole tablets was not tolerated due to an inability to swallow solid dosage forms

Product Name: Tiglutik		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
TIGLUTIK	RILUZOLE SUSP 50 MG/10ML	74503070001820	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Tiglutik		
Approval Length 12/31/2039		
Guideline Type	Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
TIGLUTIK	RILUZOLE SUSP 50 MG/10ML	74503070001820	Brand

Approval Criteria

1 - Diagnosis of amyotrophic lateral sclerosis (ALS)

AND

2 - A trial of generic riluzole tablets was not tolerated due to an inability to swallow solid dosage forms

Date	Notes
10/10/2023	New program

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Guideline ID	GL-136666	
Guideline Name	Tobacco Cessation Therapy	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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Note:

Effective 2/1/2023 these restrictions and quantity limits do not apply to persons with IL plans

1. Criteria

Product Name: NICOTROL INHALER, NICOTROL NS					
Approval Length		12 month(s)			
Therapy Stage		Initial Authorization			
Guideline Type		Prior Authorization – MN plans			
Product Name	Generic Name		GPI	Brand/Generic	
NICOTROL INHALER	NICOTINE INHALER SYSTEM 10 MG (4 MG DELIVERED)		62100005002410	Brand	
NICOTROL NS	NICOTINE NASAL SPRAY 10 MG/ML (0.5 MG/SPRAY) 62100005002020 Brand		Brand		

- 1 Both of the following:
- **1.1** Person requires a smoking cessation product that is designed to alleviate acute cravings and/or replace behavioral activities of smoking

AND

1.2 Trial and failure, contraindication, or intolerance to nicotine gum or nicotine lozenges

OR

2 - Member with stage four metastatic cancer and smoking cessation therapy is supportive care related to their cancer diagnosis

Product Name: NICOTROL INHALER, NICOTROL NS		
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization – MN plans	

Product Name	Generic Name	GPI	Brand/Generic
NICOTROL INHALER	NICOTINE INHALER SYSTEM 10 MG (4 MG DELIVERED)	62100005002410	Brand
NICOTROL NS	NICOTINE NASAL SPRAY 10 MG/ML (0.5 MG/SPRAY)	62100005002020	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: NICOTROL INHALER, NICOTROL NS	
Approval Length	12/31/2039

Guideline Type		Prior Authorization – ALL plans except MN		
Product Name	Generic Name		GPI	Brand/Generic
NICOTROL INHALER	NICOTINE INHALER SYSTEM 10 MG (4 MG DELIVERED)		62100005002410	Brand
NICOTROL NS	NICOTINE NASAL SPRAY 10 MG/ML (0.5 MG/SPRAY)		62100005002020	Brand

1 - Person requires a smoking cessation product that is designed to alleviate acute cravings and/or replace behavioral activities of smoking

AND

2 - Trial and failure, contraindication, or intolerance to nicotine gum or nicotine lozenges

Date	Notes
11/21/2023	Criteria updated

Tobramycin for Inhalation
The final contract was to be the first to be to be to the contract and a self-or the contract and a self-or to contract an

Guideline ID	GL-144931	
Guideline Name	obramycin for Inhalation	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/16/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: generic tobramycin inhalation solution				
Approval Length		12 month(s)		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
TOBRAMYCIN	TOBRAMY	'CIN NEBU SOLN 300 MG/5ML	07000070002520	Generic

Approval Criteria

1 - One of the following:

- **1.1** All of the following:
- **1.1.1** Diagnosis of cystic fibrosis

AND

1.1.2 Submission of medical records (e.g., chart notes) documenting a current culture positive for, or history of recurrent Pseudomonas aeruginosa lung infections

OR

1.2 Continuation of prior therapy with inhaled tobramycin, verified by paid claims or medical records (e.g. chart notes)

Product Name: generic tobramycin inhalation solution	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TOBRAMYCIN	TOBRAMYCIN NEBU SOLN 300 MG/5ML	07000070002520	Generic

Approval Criteria

- 1 One of the following:
- **1.1** All of the following:
- **1.1.1** Diagnosis of cystic fibrosis

AND

1.1.2 Submission of medical records (e.g., chart notes) documenting a current culture positive for, or history of recurrent Pseudomonas aeruginosa lung infections

OR

1.2 Continuation of prior therapy with inhaled tobramycin, verified by paid claims or medical records (e.g. chart notes)

Date	Notes
3/27/2024	Guideline Update.

Tremfya (guselkumab)
The bear interpretate the depart. The their terminal content of the depart prime the court the arrangement to depart in the terminal court of the depart of

Guideline ID	GL-144979	
Guideline Name	Tremfya (guselkumab)	
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/16/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Tremfya				
Diagnosis		Moderate to Severe Plaque Psoriasis		
Approval Length		12 month(s)*		
Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
TREMFYA	GUSELKUMA	NB SOLN PEN-INJECTOR 100 MG/ML	9025054200D220	Brand
TREMFYA	GUSELKUMAB SOLN PREFILLED SYRINGE 100 MG/ML		9025054200E520	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Patient has one of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than 3%
 - Debilitating palmar or plantar psoriasis or other vulnerable areas that are difficult to treat such as nails, hairy/scalp areas, genitals, or intertriginous areas

AND

3.1.2 Submission of medical records (e.g., chart notes) documenting trial and failure (minimum 4 week trial duration), or contraindication to topical therapy (e.g., topical corticosteroids, calcipotriene, retinoids, calcineurin inhibitor, tazarotene)

OR

3.2 Continuation of prior therapy with guselkumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
	30 days Second PA: Approve at GPI 10 for 12 months

Product Name: Tremfya	
Diagnosis	Moderate to Severe Plaque Psoriasis

Approval Length	12/31/2039*	
Guideline Type	Prior Authorization – All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
TREMFYA	GUSELKUMAB SOLN PEN-INJECTOR 100 MG/ML	9025054200D220	Brand
TREMFYA	GUSELKUMAB SOLN PREFILLED SYRINGE 100 MG/ML	9025054200E520	Brand

1 - Diagnosis of moderate to severe plaque psoriasis

AND

2 - Prescribed by or in consultation with a dermatologist

AND

- 3 One of the following:
- **3.1** Both of the following:
- **3.1.1** Patient has one of the following:
 - Significant functional disability
 - Body surface area (BSA) involvement of greater than 3%
 - Debilitating palmar or plantar psoriasis or other vulnerable areas that are difficult to treat such as nails, hairy/scalp areas, genitals, or intertriginous areas

AND

3.1.2 Submission of medical records (e.g., chart notes) documenting trial and failure (minimum 4 week trial duration), or contraindication to topical therapy (e.g., topical corticosteroids, calcipotriene, retinoids, calcineurin inhibitor, tazarotene)

OR

3.2 Continuation of prior therapy with guselkumab, verified by paid claims or medical records (e.g. chart notes)	
Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date: First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for 30 days Second PA: Approve at GPI 10 to 12/31/2039

Product Name: Tremfya	
Diagnosis Psoriatic Arthritis (PsA)	
Approval Length	12 month(s)*
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
TREMFYA	GUSELKUMAB SOLN PEN-INJECTOR 100 MG/ML	9025054200D220	Brand
TREMFYA	GUSELKUMAB SOLN PREFILLED SYRINGE 100 MG/ML	9025054200E520	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- 3 One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis

OR

3.2 Continuation of prior therapy with guselkumab, verified by paid claims or medical records (e.g. chart notes)

Notes

*For new starts to therapy: Enter 2 PAs as follows with the same start date:
First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for 30 days

Product Name: Tremfya		
Diagnosis Psoriatic Arthritis (PsA)		
Approval Length	12/31/2039*	
Guideline Type Prior Authorization – All Plans Except IL and MN Plans		

Second PA: Approve at GPI 10 for 12 months

Product Name	Generic Name	GPI	Brand/Generic
TREMFYA	GUSELKUMAB SOLN PEN-INJECTOR 100 MG/ML	9025054200D220	Brand
TREMFYA	GUSELKUMAB SOLN PREFILLED SYRINGE 100 MG/ML	9025054200E520	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with a dermatologist or rheumatologist

AND

- **3** One of the following:
- **3.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:

- actively inflamed joints axial disease
- active skin, nail, or scalp psoriasis involvement
- enthesitis

OR

3.2 Continuation of prior therapy with guselkumab, verified by paid claims or medical records (e.g. chart notes)

Notes	*For new starts to therapy: Enter 2 PAs as follows with the same start date:
	First PA: Approve at GPI 14 with a MDD of 0.04 (1 every 28 days) for
	30 days Second PA: Approve at GPI 10 to 12/31/2039

Date	Notes
4/9/2024	Guideline Update.

T	Tresiba (insulin degludec)		
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Guideline ID	GL-129810	
Guideline Name	Tresiba (insulin degludec)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Brand Insulin Degludec U100	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
INSULIN DEGLUDEC FLEXTOUCH	INSULIN DEGLUDEC SOLN PEN-INJECTOR 100 UNIT/ML	2710400700D210	Brand
INSULIN DEGLUDEC	INSULIN DEGLUDEC INJ 100 UNIT/ML	27104007002020	Brand

Approval Criteria

1 - Diagnosis of diabetes mellitus

AND

- **2** Prescribed by or in consultation with one of the following:
 - Endocrinologist
 - Diabetes specialist

AND

- 3 One of the following:
- **3.1** Member cannot meet their glycemic goals despite adequate trials of insulin glargine including:
 - Dose escalation, unless dose increases cannot be tolerated due to nocturnal hypoglycemia or at least one severe low blood sugar event (requiring assistance from another) or would not be appropriate given the person's self-monitoring blood glucose profile
 - Splitting the dose
 - Submission of medical records (e.g., chart notes) documenting use of a specific adherence intervention deployed by a health care provider

OR

3.2 Member is intolerant to insulin glargine

Product Name: Brand Insulin Degludec U200	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
INSULIN DEGLUDEC FLEXTOUCH	INSULIN DEGLUDEC SOLN PEN-INJECTOR 200 UNIT/ML	2710400700D220	Brand

1 - Diagnosis of diabetes mellitus

AND

- **2** Prescribed by or in consultation with one of the following:
 - Endocrinologist
 - Diabetes specialist

AND

- **3** One of the following:
- **3.1** Member cannot meet their glycemic goals despite adequate trials of insulin glargine including:
 - Dose escalation, unless dose increases cannot be tolerated due to nocturnal hypoglycemia or at least one severe low blood sugar event (requiring assistance from another) or would not be appropriate given the person's self-monitoring blood glucose profile
 - Splitting the dose
 - Submission of medical records (e.g., chart notes) documenting use of a specific adherence intervention deployed by a health care provider

OR

3.2 Member is intolerant to insulin glargine

AND

4 - Member's daily basal insulin dose is greater than 100 units

Product Name: Brand Insulin Degludec U100 and U200	
Approval Length	12 month(s)

Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
INSULIN DEGLUDEC FLEXTOUCH	INSULIN DEGLUDEC SOLN PEN-INJECTOR 100 UNIT/ML	2710400700D210	Brand
INSULIN DEGLUDEC FLEXTOUCH	INSULIN DEGLUDEC SOLN PEN-INJECTOR 200 UNIT/ML	2710400700D220	Brand
INSULIN DEGLUDEC	INSULIN DEGLUDEC INJ 100 UNIT/ML	27104007002020	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Brand Insulin Degludec U100	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
INSULIN DEGLUDEC FLEXTOUCH	INSULIN DEGLUDEC SOLN PEN-INJECTOR 100 UNIT/ML	2710400700D210	Brand
INSULIN DEGLUDEC	INSULIN DEGLUDEC INJ 100 UNIT/ML	27104007002020	Brand

Approval Criteria

1 - Diagnosis of diabetes mellitus

AND

- 2 Prescribed by or in consultation with one of the following:
 - Endocrinologist

Diabetes specialist

AND

- **3** One of the following:
- **3.1** Member cannot meet their glycemic goals despite adequate trials of insulin glargine including:
 - Dose escalation, unless dose increases cannot be tolerated due to nocturnal hypoglycemia or at least one severe low blood sugar event (requiring assistance from another) or would not be appropriate given the person's self-monitoring blood glucose profile
 - Splitting the dose
 - Submission of medical records (e.g., chart notes) documenting use of a specific adherence intervention deployed by a health care provider

OR

3.2 Member is intolerant to insulin glargine

Product Name: Brand Insulin Degludec U200	
Approval Length	12/31/2039
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
INSULIN DEGLUDEC FLEXTOUCH	INSULIN DEGLUDEC SOLN PEN-INJECTOR 200 UNIT/ML	2710400700D220	Brand

Approval Criteria

1 - Diagnosis of diabetes mellitus

AND

2 - Prescribed by or in consultation with one of the following:

- Endocrinologist
- Diabetes specialist

AND

- 3 One of the following:
- **3.1** Member cannot meet their glycemic goals despite adequate trials of insulin glargine including:
 - Dose escalation, unless dose increases cannot be tolerated due to nocturnal hypoglycemia or at least one severe low blood sugar event (requiring assistance from another) or would not be appropriate given the person's self-monitoring blood glucose profile
 - Splitting the dose
 - Submission of medical records (e.g., chart notes) documenting use of a specific adherence intervention deployed by a health care provider

OR

3.2 Member is intolerant to insulin glargine

AND

4 - Member's daily basal insulin dose is greater than 100 units

Date	Notes
10/12/2023	2024 New Implementation

	Tudorza Pressair					
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Guideline ID	GL-127804
Guideline Name	Tudorza Pressair
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Tudorza Pressair				
Approval Length	12 month(s)			
Therapy Stage	Initial Authorization			
Guideline Type	Step Therapy - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
	ACLIDINIUM BROMIDE AEROSOL POWD BREATH ACTIVATED 400 MCG/ACT	44100007108020	Brand

Approval Criteria

1 - Trial and failure, intolerance, or contraindication to both an inhaled tiotropium bromide product and an inhaled umeclidinium product

Product Name: Tudorza Pressair	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Step Therapy - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
	ACLIDINIUM BROMIDE AEROSOL POWD BREATH ACTIVATED 400 MCG/ACT	44100007108020	Brand

1 - Submission of medical notes (e.g. chart notes) from the past 12 months that member is continuing therapy with the requested drug

Product Name: Tudorza Pressair	
Approval Length 12/31/2039	
Guideline Type Step Therapy - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
	ACLIDINIUM BROMIDE AEROSOL POWD BREATH ACTIVATED 400 MCG/ACT	44100007108020	Brand

Approval Criteria

1 - Trial and failure, intolerance, or contraindication to both an inhaled tiotropium bromide product and an inhaled umeclidinium product

Date	Notes
8/21/2023	New Program

Vaccine	<u> </u>			
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Guideline ID	GL-136474	
Guideline Name	Vaccines	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Recombivax-HB, Engerix-B, Arexvy, Abrysvo, Boostrix, Prehevbrio, Twinrix, Prevnar 13, Prevnar 20, Vaxneuvance, Pneumovax, Adacel, Boostrix. Tdvax, Tenivac, Shingrix	
Approval Length	12 month(s)
Guideline Type	Administrative

Product Name	Generic Name	GPI	Brand/Generic
RECOMBIVAX HB	HEPATITIS B VACCINE (RECOMBINANT) SUSP 5 MCG/0.5ML	17100010201815	Brand
RECOMBIVAX HB	HEPATITIS B VACCINE (RECOMBINANT) SUSP 10 MCG/ML	17100010201820	Brand
ENGERIX-B	HEPATITIS B VACCINE (RECOMBINANT) SUSP PREF SYR 20 MCG/ML	1710001020E630	Brand
HEPLISAV-B	HEPATITIS B VACCINE RECOMB ADJUVANTED PREF SYR 20 MCG/0.5ML	1710001030E520	Brand

AREXVY	RSVPREF3 VACCINE RECOMB ADJUVANTED FOR IM SUSP 120 MCG/0.5ML	17100072101920	Brand
ABRYSVO	RSV PRE-FUSION F A&B VAC RECOMB FOR IM SOLN 120 MCG/0.5ML	17100072202120	Brand
BOOSTRIX	TET-DIPH-ACELL PERTUSS AD PREF SYR 5-2.5- 18.5 LF-MCG/0.5ML	1899000322E620	Brand
PREHEVBRIO	HEPATITIS B VACCINE 3-ANTIGEN (RECOMBINANT) SUSP 10 MCG/ML	17100010401820	Brand
ENGERIX-B	HEPATITIS B VACCINE (RECOMBINANT) SUSP 20 MCG/ML	17100010201830	Brand
ENGERIX-B	HEPATITIS B VACCINE (RECOMBINANT) SUSP PREF SYR 10 MCG/0.5ML	1710001020E625	Brand
RECOMBIVAX HB	HEPATITIS B VACCINE (RECOMBINANT) SUSP 40 MCG/ML	17100010201840	Brand
RECOMBIVAX HB	HEPATITIS B VACCINE (RECOMBINANT) SUSP PREF SYR 5 MCG/0.5ML	1710001020E610	Brand
RECOMBIVAX HB	HEPATITIS B VACCINE (RECOMBINANT) SUSP PREF SYR 10 MCG/ML	1710001020E620	Brand
TWINRIX	HEP A-HEP B VACCINE SUSP PREF SYR 720-20 ELU-MCG/ML	1710990205E620	Brand
PREVNAR 13	PNEUMOCOCCAL 13-VALENT CONJUGATE VACCINE INJ	17200065301800	Brand
VAXNEUVANCE	PNEUMOCOCCAL 15-VALENT CONJUGATE VACCINE SUS PREF SYR 0.5 ML	1720006535E620	Brand
PREVNAR 20	PNEUMOCOCCAL 20-VALENT CONJUGATE VACCINE SUS PREF SYR 0.5 ML	1720006540E620	Brand
ADACEL	TET TOX-DIPH-ACELL PERTUSS AD INJ 5-2-15.5 LF-LF-MCG/0.5ML	18990003221815	Brand
PNEUMOVAX 23	PNEUMOCOCCAL VACCINE POLYVALENT INJ 25 MCG/0.5ML	17200065002205	Brand
PNEUMOVAX 23/1 DOSE	PNEUMOCOCCAL VACCINE POLYVALENT INJ 25 MCG/0.5ML	17200065002205	Brand
TDVAX	TETANUS-DIPHTHERIA TOXOIDS (TD) INJ 2-2 LF/0.5ML	18990002201805	Brand
BOOSTRIX	TET TOX-DIPH-ACELL PERTUSS AD INJ 5-2.5- 18.5 LF-LF-MCG/0.5ML	18990003221820	Brand
TENIVAC	TETANUS-DIPHTHERIA TOXOIDS (TD) INJ 5-2 LFU	18990002202210	Brand
SHINGRIX	ZOSTER VAC RECOMBINANT ADJUVANTED FOR IM INJ 50 MCG/0.5ML	17100095401920	Brand

1 - Member is 18 years or older*

AND

- 2 One of the following:
- **2.1** The requested vaccination will be used for a Food and Drug Administration (FDA) approved indication

OR

2.2 The requested vaccination will be used in accordance with Advisory Committee on Immunization Practices (ACIP) recommendation

Notes	*Vaccines listed above are considered excluded for persons under the
	age of 18 years. They are covered under the medical benefit

Date	Notes
12/5/2023	New Program

Valtoc	o (diazep	am)	
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Guideline ID	GL-129092	
Guideline Name	Valtoco (diazepam)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Valtoco				
Approval L	oval Length 12/31/2039			
Guideline ⁻	Гуре	Prior Authorization - All plans excep	ot IL and MN Plans	
Product Name	Generic Na	eric Name GPI Brand/Generi		Brand/Generic
VALTOCO 15 MG DOSE	DIAZEPAM NASAL SPRAY THER PACK 2 X 7.5 MG/0.1ML (15 MG DOSE)		7210003000C440	Brand
VALTOCO 20 MG DOSE	DIAZEPAM NASAL SPRAY THER PACK 2 X 10 MG/0.1ML (20 MG DOSE)		7210003000C450	Brand
VALTOCO 5 MG DOSE	DIAZEPAM N	ASAL SPRAY 5 MG/0.1 ML	72100030000920	Brand

VALTOCO 10 MG DOSE	DIAZEPAM NASAL SPRAY 10 MG/0.1 ML	72100030000930	Brand
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1 - Diagnosis of a seizure disorder (epilepsy)

AND

2 - Prescribed by or in consultation with a neurologist or other specialist with experience in the management of epilepsy

AND

3 - Member is between the ages of 6 and 12 years old

AND

4 - Submission of medical records (e.g., chart notes) that support history of frequent episodes of acute seizure activity

Product Name: Valtoco		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
VALTOCO 15 MG DOSE	DIAZEPAM NASAL SPRAY THER PACK 2 X 7.5 MG/0.1ML (15 MG DOSE)	7210003000C440	Brand
VALTOCO 20 MG DOSE	DIAZEPAM NASAL SPRAY THER PACK 2 X 10 MG/0.1ML (20 MG DOSE)	7210003000C450	Brand
VALTOCO 5 MG DOSE	DIAZEPAM NASAL SPRAY 5 MG/0.1 ML	72100030000920	Brand

VALTOCO DIAZEPAM NASAL SPRAY 10 MG/0.1 ML 10 MG DOSE	72100030000930	Brand
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1 - Diagnosis of a seizure disorder (epilepsy)

AND

2 - Prescribed by or in consultation with a neurologist or other specialist with experience in the management of epilepsy

AND

3 - Member is between the ages of 6 and 12 years old

AND

4 - Submission of medical records (e.g., chart notes) that support history of frequent episodes of acute seizure activity

Product Name: Valtoco		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
VALTOCO 15 MG DOSE	DIAZEPAM NASAL SPRAY THER PACK 2 X 7.5 MG/0.1ML (15 MG DOSE)	7210003000C440	Brand
VALTOCO 20 MG DOSE	DIAZEPAM NASAL SPRAY THER PACK 2 X 10 MG/0.1ML (20 MG DOSE)	7210003000C450	Brand
VALTOCO 5 MG DOSE	DIAZEPAM NASAL SPRAY 5 MG/0.1 ML	72100030000920	Brand

VALTOCO DIAZEPAM NASAL SPRAY 10 N 10 MG DOSE	G/0.1 ML 721000300009	Brand
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1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
9/7/2023	2024 New Implementation

Vascepa (Icosa	Vascepa (Icosapent Ethyl)				
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Guideline ID	GL-129625	
Guideline Name	Vascepa (Icosapent Ethyl)	
Formulary • Quartz		

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Icosapent Ethyl			
Approval Length 12 month(s)			
Therapy Stage	Therapy Stage Initial Authorization		
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
ICOSAPENT ETHYL	ICOSAPENT ETHYL CAP 0.5 GM	39500035100110	Generic
ICOSAPENT ETHYL	ICOSAPENT ETHYL CAP 1 GM	39500035100120	Generic

Approval Criteria

1 - Diagnosis of established cardiovascular disease* OR diabetes mellitus with ≥ 2 additional risk factors for cardiovascular disease**

AND

2 - Prescribed by, or in consultation with, a Cardiologist, Endocrinologist, or other lipid specialist

AND

3 - Triglycerides ≥ 150 mg/dL

AND

4 - Using as an adjunct to maximally tolerated statin therapy

OR

5 - Clinical documentation to support statin intolerance***

Notes

*ASCVD refers to the following conditions: coronary heart disease such as myocardial infarction, angina, coronary artery stenosis >50%; cerebrovascular disease such as transient ischemic at tack, ischemic stroke, or carotid artery stenosis > 50%; peripheral artery disease such as claudication; and aortic athero sclerotic disease such as abdominal aortic aneurysm and descending thoracic aneurysm.

- **Additional risk factors may include: current smoker, family history of premature ASCVD, LDL cholesterol persistently ≥ 160 mg/dL, chronic kidney disease, metabolic syndrome, South Asian ancestry, or other guideline supported risk factors
- ***Statin intolerance is defined as the inability to tolerate at least 2 stat ins, with:
- ♣ one started at the lowest starting dose
- ♣ statin dose reduction was attempted to resolve symptoms or lab ab normalities (not discontinuation)
- * symptoms or lab abnormalities reversed with statin discontinuation but returned with re-challenge of statins

♣ symptoms or lab abnormalities are not due to established predispos
itions such as drug interactions, significant
changes in physical activity, or underlying muscle disease

Product Name: Generic Icosapent Ethyl			
Approval Length 12 month(s)			
Therapy Stage	Therapy Stage Reauthorization		
Guideline Type Prior Authorization - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
ICOSAPENT ETHYL	ICOSAPENT ETHYL CAP 0.5 GM	39500035100110	Generic
ICOSAPENT ETHYL	ICOSAPENT ETHYL CAP 1 GM	39500035100120	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

N	otes
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*ASCVD refers to the following conditions: coronary heart disease such as myocardial infarction, angina, coronary artery stenosis >50%; cerebrovascular disease such as transient ischemic at tack, ischemic stroke, or carotid artery stenosis > 50%; peripheral artery disease such as claudication; and aortic athero sclerotic disease such as abdominal aortic aneurysm and descending thoracic aneurysm.

- **Additional risk factors may include: current smoker, family history of premature ASCVD, LDL cholesterol persistently ≥ 160 mg/dL, chronic kidney disease, metabolic syndrome, South Asian ancestry, or other guideline supported risk factors
- ***Statin intolerance is defined as the inability to tolerate at least 2 stat ins, with:
- one started at the lowest starting dose
- * statin dose reduction was attempted to resolve symptoms or lab ab normalities (not discontinuation)
- * symptoms or lab abnormalities reversed with statin discontinuation but returned with re-challenge of statins
- symptoms or lab abnormalities are not due to established predispos itions such as drug interactions, significant changes in physical activity, or underlying muscle disease

Product Name: Generic Icosapent Ethyl		
Approval Length 12/31/2039		
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
ICOSAPENT ETHYL	ICOSAPENT ETHYL CAP 0.5 GM	39500035100110	Generic
ICOSAPENT ETHYL	ICOSAPENT ETHYL CAP 1 GM	39500035100120	Generic

1 - Diagnosis of established cardiovascular disease* OR diabetes mellitus with ≥ 2 additional risk factors for cardiovascular disease**

AND

2 - Prescribed by, or in consultation with, a Cardiologist, Endocrinologist, or other lipid specialist

AND

3 - Triglycerides ≥ 150 mg/dL

AND

4 - Using as an adjunct to maximally tolerated statin therapy

OR

5 - Clinical documentation to support statin intolerance***

Notes	*ASCVD refers to the following conditions: coronary heart disease such as myocardial infarction, angina, coronary artery
	stenosis >50%; cerebrovascular disease such as transient ischemic at tack, ischemic stroke, or carotid artery stenosis >
	50%; peripheral artery disease such as claudication; and aortic athero

sclerotic disease such as abdominal aortic aneurysm and descending thoracic aneurysm. **Additional risk factors may include: current smoker, family history of premature ASCVD, LDL cholesterol persistently ≥ 160 mg/dL, chronic kidney disease, metabolic syndrome, South Asian ancestry, or other guideline supported risk factors ***Statin intolerance is defined as the inability to tolerate at least 2 stat ins, with: ♣ one started at the lowest starting dose * statin dose reduction was attempted to resolve symptoms or lab ab normalities (not discontinuation) * symptoms or lab abnormalities reversed with statin discontinuation but returned with re-challenge of statins * symptoms or lab abnormalities are not due to established predispos itions such as drug interactions, significant changes in physical activity, or underlying muscle disease

Date	Notes
10/25/2023	New Program

Vemlidy (tenofovir alafenamide)				
(i) but saving over sideles. Note the least of loved a read and and defining provide actually				

Guideline ID	GL-131349
Guideline Name	Vemlidy (tenofovir alafenamide)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Vemlidy				
Approval Length 12/31/2039				
Guideline Type Prior Authorization - All plans except IL and MN Plans		1		
Product Name	Generic Name		GPI	Brand/Generic
VEMLIDY TENOFOVI		ALAFENAMIDE FUMARATE TAB 25 MG	12352083200320	Brand

1 - Diagnosis of chronic hepatitis B

AND

- 2 One of the following:
 - Member has failed entecavir
 - Submission of medical records (e.g., chart notes) documenting member has lamivudine resistance

AND

3 - Trial and failure, contraindication, or intolerance to tenofovir disoproxil fumarate

Product Name: Vemlidy	
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
VEMLIDY	TENOFOVIR ALAFENAMIDE FUMARATE TAB 25 MG	12352083200320	Brand

Approval Criteria

- **1** ALL of the following:
- 1.1 Diagnosis of chronic hepatitis B

AND

- **1.2** One of the following:
 - Member has failed entecavir
 - Submission of medical records (e.g., chart notes) documenting member has lamivudine resistance

AND

1.3 Trial and failure, contraindication, or intolerance to tenofovir disoproxil fumarate

OR

2 - (Minnesota plans only): Member has stage four metastatic cancer and the requested drug is being used to treat cancer-related hepatitis B infection

Product Name: Vemlidy	
Approval Length 12 month(s)	
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Ш	Product Name	Generic Name	GPI	Brand/Generic
	VEMLIDY	TENOFOVIR ALAFENAMIDE FUMARATE TAB 25 MG	12352083200320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
10/8/2023	2024 New Implementation

Verkazia (cyclosporine ophthalmic emulsion 0.19				

Guideline ID	GL-129065	
Guideline Name	Verkazia (cyclosporine ophthalmic emulsion 0.1%)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Verkazia	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization*
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
VERKAZIA	CYCLOSPORINE (OPHTH) EMULSION 0.1%	86720020001630	Brand

Approval Criteria

1 - Diagnosis of moderate to severe vernal keratoconjunctivitis (e.g. visual deficit and/or continuous symptoms)

AND

2 - Positive skin tests or in vitro reactivity to common aeroallergens (e.g. pollen, dust mites, pet dander, cockroaches, etc.)

AND

3 - Continued symptoms despite a two-week trial of an ophthalmic steroid

AND

- **4** Trial and failure with first-line treatments including all of the following:
- **4.1** Lifestyle changes (e.g. cold compresses, trigger avoidance, artificial tears)

AND

- **4.2** A three-week trial of one of the following, in combination with ophthalmic cyclosporine $(0.05\ \%\ or\ 0.09\%)$
 - Topical ophthalmic mast cell stabilizer (e.g., cromolyn sodium, lodoxamide tromethamine, nedocromil sodium)
 - Topical dual action mast cell stabilizer/antihistamine (e.g., olopatadine, azelastine hydrochloride, epinastine, ketotifen fumarate)

AND

5 - Prescribed by or in consultation with an ophthalmologist

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through
reauthorization criteria

Product Name: Verkazia

Approval Length	12 month(s)
Therapy Stage	Reauthorization*
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
VERKAZIA	CYCLOSPORINE (OPHTH) EMULSION 0.1%	86720020001630	Brand

1 - Diagnosis of moderate to severe vernal keratoconjunctivitis (e.g. visual deficit and/or continuous symptoms)

AND

2 - Positive skin tests or in vitro reactivity to common aeroallergens (e.g. pollen, dust mites, pet dander, cockroaches, etc.)

AND

3 - Continued symptoms despite a two-week trial of an ophthalmic steroid

AND

- **4** Trial and failure with first-line treatments including all of the following:
 - **4.1** Lifestyle changes (e.g. cold compresses, trigger avoidance, artificial tears)

AND

- **4.2** A three-week trial of one of the following, in combination with ophthalmic cyclosporine (0.05 % or 0.09%)
 - Topical ophthalmic mast cell stabilizer (e.g., cromolyn sodium, lodoxamide tromethamine, nedocromil sodium)

• Topical Dual action mast cell stabilizer/antihistamine (e.g., olopatadine, azelastine hydrochloride, epinastine, ketotifen fumarate)

AND

5 - Prescribed by or in consultation with an ophthalmologist

AND

6 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months, the member has improved while on therapy

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) and were not previously approved for c overage whose therapy was initiated using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through
reauthorization criteria

Date	Notes
7/28/2023	2024 New Implementation

Verquvo (vericiguat)			
The birth regions in the last tension most, count, a state she birth to provide an own burst before			

Guideline ID	GL-141086
Guideline Name	Verquvo (vericiguat)
Formulary	Quartz

Guideline Note:

Effective Date:	2/3/2024
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1. Criteria

Product Name: Verquvo					
Approval Length	12/31/2039				
Guideline Type Prior Authorization - ALL Plans Except IL and MN Plans					
		ODI		1/0	

Product Name	Generic Name	GPI	Brand/Generic
VERQUVO	VERICIGUAT TAB 2.5 MG	40900085000321	Brand
VERQUVO	VERICIGUAT TAB 5 MG	40900085000330	Brand
VERQUVO	VERICIGUAT TAB 10 MG	40900085000340	Brand

Approval Criteria

1 - Diagnosis of symptomatic chronic heart failure (HF)

AND
2 - Ejection fraction less than 45%
AND
3 - Hospitalization related to HF in the past 6 months
AND
4 - One of the following:
4.1 Both of the following:
4.1.1 Trial and failure, contraindication or intolerance to one of the following:
 Angiotensin-converting enzyme inhibitor (e.g., enalapril, lisinopril, ramipril) Angiotensin II receptor blocker (e.g., candesartan, losartan, valsartan) Angiotensin receptor neprilysin inhibitors (e.g., sacubitril/valsartan)
AND
4.1.2 Trial and failure, contraindication or intolerance to beta-blocker (e.g., bisoprolol, carvedilol, metoprolol succinate)
OR
4.2 Trial and failure, contraindication or intolerance to an aldosterone antagonist (e.g., eplerenone, spironolactone)
AND
5 - Prescribed by or in consultation with a cardiologist

Product Name: Verquvo	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
VERQUVO	VERICIGUAT TAB 2.5 MG	40900085000321	Brand
VERQUVO	VERICIGUAT TAB 5 MG	40900085000330	Brand
VERQUVO	VERICIGUAT TAB 10 MG	40900085000340	Brand

1 - Diagnosis of symptomatic chronic heart failure (HF)

AND

2 - Ejection fraction less than 45%

AND

3 - Hospitalization related to HF in the past 6 months

AND

- 4 One of the following:
- **4.1** Both of the following:
- **4.1.1** Trial and failure, contraindication or intolerance to one of the following:
 - Angiotensin-converting enzyme inhibitor (e.g., enalapril, lisinopril, ramipril)
 - Angiotensin II receptor blocker (e.g., candesartan, losartan, valsartan)
 - Angiotensin receptor neprilysin inhibitors (e.g., sacubitril/valsartan)

AND

4.1.2 Trial and failure, contraindication or intolerance to beta-blocker (e.g., bisoprolol, carvedilol, metoprolol succinate)

OR

4.2 Trial and failure, contraindication or intolerance to an aldosterone antagonist (e.g., eplerenone, spironolactone)

AND

5 - Prescribed by or in consultation with a cardiologist

	*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
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Product Name: Verquvo		
Approval Length 12 month(s)		
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
VERQUVO	VERICIGUAT TAB 2.5 MG	40900085000321	Brand
VERQUVO	VERICIGUAT TAB 5 MG	40900085000330	Brand
VERQUVO	VERICIGUAT TAB 10 MG	40900085000340	Brand

Approval Criteria

1 - Diagnosis of symptomatic chronic heart failure (HF)

AND

2 - Ejection fraction less than 45%

AND
3 - Hospitalization related to HF in the past 6 months
AND
4 - One of the following:
4.1 BOTH of the following:
4.1.1 Trial and failure, contraindication or intolerance to one of the following:
 Angiotensin-converting enzyme inhibitor (e.g., enalapril, lisinopril, ramipril) Angiotensin II receptor blocker (e.g., candesartan, losartan, valsartan) Angiotensin receptor neprilysin inhibitors (e.g., sacubitril/valsartan)
AND
4.1.2 Trial and failure, contraindication or intolerance to beta-blocker (e.g., bisoprolol, carvedilol, metoprolol succinate)
OR
4.2 Trial and failure, contraindication or intolerance to an aldosterone antagonist (e.g., eplerenone, spironolactone)
AND
5 - Prescribed by or in consultation with a cardiologist
AND
6 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member's is stable or an improvement is seen while on therapy with the requested drug

*Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufactu rer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies
new to plan, reauthorization criteria applies

Date	Notes
2/3/2024	Update Program

V	iagra (silde	nafil)	
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Guideline ID	GL-144569
Guideline Name	Viagra (sildenafil)
Formulary	Quartz

Guideline Note:

Effective Date:	4/21/2024
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1. Criteria

Approval Length		12/31/2039		
Guideline Type		Quantity Limit - ALL Plans Except IL and MN Plans		
Product Name	Generic N	ame	GPI	Brand/Generic
SILDENAFIL CITRATE	SILDENAFIL	CITRATE TAB 25 MG	40304070100310	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 50 MG		40304070100320	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 100 MG		40304070100330	Generic

1 - Submission of medical records (e.g., chart notes) documenting the actual number of intercourse events within a 30-day period and the dose required to achieve an erection

AND

2 - Submission of medical records (e.g., chart notes) documenting that the quantity for treatment cannot be met with the commercially available dose forms within the quantity limit*

Notes	*QTY Limit: MAX 15 doses per 30 days
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Product Name: Generic sildenafil 25mg, 50mg, 100mg	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Quantity Limit - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 25 MG	40304070100310	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 50 MG	40304070100320	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 100 MG	40304070100330	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting the actual number of intercourse events within a 30-day period and the dose required to achieve an erection

AND

2 - Submission of medical records (e.g., chart notes) documenting that the quantity for treatment cannot be met with the commercially available dose forms within the quantity limit*

Notes	1*QTY Limit: MAX 15 doses per 30 days
LINDIAS	I "O I Y I IMIT: IVIA X 15 MOSES DEL 311 MAVS

Product Name: Generic	c sildenafil 25mg, 50mg, 100mg
Approval Length	12 month(s)

Therapy Stage	Reauthorization
Guideline Type	Quantity Limit - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 25 MG	40304070100310	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 50 MG	40304070100320	Generic
SILDENAFIL CITRATE	SILDENAFIL CITRATE TAB 100 MG	40304070100330	Generic

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
3/18/2024	Updated product name

Viberzi (eluxadoline)
The best and the property of the best of the section would would be seen the property of the section to section be a section to the section of the section to the section t

Guideline ID	GL-129221
Guideline Name	Viberzi (eluxadoline)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Viberzi	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
VIBERZI	ELUXADOLINE TAB 75 MG	52558020000330	Brand
VIBERZI	ELUXADOLINE TAB 100 MG	52558020000340	Brand

Approval Criteria

1 - Both of the following:

1.1 Covered for persons with diarrhea predominant irritable bowel syndrome (IBS)

AND

1.2 Have failed, or been intolerant to, a one-month trial of conventional therapy (such as loperamide or diphenoxylate/atropine)

Product Name: Viberzi	
Approval Length 12 month(s)	
Therapy Stage Reauthorization Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
VIBERZI	ELUXADOLINE TAB 75 MG	52558020000330	Brand
VIBERZI	ELUXADOLINE TAB 100 MG	52558020000340	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Viberzi	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
VIBERZI	VIBERZI ELUXADOLINE TAB 75 MG		Brand
VIBERZI	ELUXADOLINE TAB 100 MG	52558020000340	Brand

Approval Criteria

1 - Both of the following:

1.1 Covered for persons with diarrhea predominant irritable bowel syndrome (IBS)

AND

1.2 Have failed, or been intolerant to, a one-month trial of conventional therapy (such as loperamide or diphenoxylate/atropine)

Date	Notes
10/6/2023	New Program

Vimpat (lacosamide)
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Guideline ID	GL-128134
Guideline Name	Vimpat (lacosamide)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Generic Lacosamide	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Step Therapy - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
LACOSAMIDE	LACOSAMIDE TAB 50 MG	72600036000320	Generic
LACOSAMIDE	LACOSAMIDE TAB 100 MG	72600036000330	Generic
LACOSAMIDE	LACOSAMIDE TAB 150 MG	72600036000340	Generic
LACOSAMIDE	LACOSAMIDE TAB 200 MG	72600036000350	Generic

- **1** Trial and failure, contraindication, or intolerance to at least TWO preferred anticonvulsants:
 - lamotrigine
 - levetiracetam
 - carbamazepine
 - valproate
 - oxcarbazepine
 - gabapentin
 - pregabalin
 - topiramate
 - phenytoin
 - zonisamide
 - primidone

Product Name: Generic Lacosamide	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Step Therapy - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
LACOSAMIDE	LACOSAMIDE TAB 50 MG	72600036000320	Generic
LACOSAMIDE	LACOSAMIDE TAB 100 MG	72600036000330	Generic
LACOSAMIDE	LACOSAMIDE TAB 150 MG	72600036000340	Generic
LACOSAMIDE	LACOSAMIDE TAB 200 MG	72600036000350	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Generic Lacosamide				
Approval Length		12/31/2039		
Guideline Type		Step Therapy - All other plans except IL and MN Plans		
Product Generic Name		Name	GPI	Brand/Generic

LACOSAMIDE	LACOSAMIDE TAB 50 MG	72600036000320	Generic
LACOSAMIDE	LACOSAMIDE TAB 100 MG	72600036000330	Generic
LACOSAMIDE	LACOSAMIDE TAB 150 MG	72600036000340	Generic
LACOSAMIDE	LACOSAMIDE TAB 200 MG	72600036000350	Generic

- **1** Trial and failure, contraindication, or intolerance to at least TWO preferred anticonvulsants:
 - lamotrigine
 - levetiracetam
 - carbamazepine
 - valproate
 - oxcarbazepine
 - gabapentin
 - pregabalin
 - topiramate
 - phenytoin
 - zonisamide
 - primidone

Date	Notes
8/25/2023	New Program

Vitamin D Analogs
(a) The ball of the grant brighty in The Box Service count or and a disk delta to probe the country of trades.

Guideline ID	GL-131955
Guideline Name	Vitamin D Analogs
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Brand: Rayaldee, Generic: Doxercalciferol, paricalcitol		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
RAYALDEE	CALCIFEDIOL CAP ER 30 MCG	30905025000230	Brand
PARICALCITOL	PARICALCITOL CAP 1 MCG	30905070000110	Generic
DOXERCALCIFEROL	DOXERCALCIFEROL CAP 0.5 MCG	30905040000105	Generic
DOXERCALCIFEROL	DOXERCALCIFEROL CAP 1 MCG	30905040000110	Generic
DOXERCALCIFEROL	DOXERCALCIFEROL CAP 2.5 MCG	30905040000120	Generic
PARICALCITOL	PARICALCITOL CAP 2 MCG	30905070000120	Generic
PARICALCITOL	PARICALCITOL CAP 4 MCG	30905070000140	Generic

1 - Dose-adjusted trial and failure (unable to achieve parathyroid hormone level goals), contraindication, or intolerance with calcitriol.

Product Name: Brand: Rayaldee, Generic: Doxercalciferol, paricalcitol				
Approval Length		12 month(s)		
Therapy Stage		Reauthorization		
Guideline Type		Prior Authorization-IL and MN Plans Only		
Product Name	Ge	neric Name	GPI	Brand/Generic
RAYALDEE	CAI	LCIFEDIOL CAP ER 30 MCG	30905025000230	Brand
PARICALCITOL	PAI	RICALCITOL CAP 1 MCG	30905070000110	Generic
DOXERCALCIFEROL	DO	XERCALCIFEROL CAP 0.5 MCG	30905040000105	Generic
DOXERCALCIFEROL	DO	XERCALCIFEROL CAP 1 MCG	30905040000110	Generic
DOXERCALCIFEROL	DO	XERCALCIFEROL CAP 2.5 MCG	30905040000120	Generic
PARICALCITOL	PAI	RICALCITOL CAP 2 MCG	30905070000120	Generic
		-	·	·

Approval Criteria

PARICALCITOL

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

PARICALCITOL CAP 4 MCG

Product Name: Brand: Rayaldee, Generic: Doxercalciferol, paricalcitol				
Approval Length		12/31/2039		
Guideline Type Prior Authorization-All plans except IL and MN				
Product Name	Ge	neric Name	GPI	Brand/Generic
RAYALDEE	CAI	CIFEDIOL CAP ER 30 MCG	30905025000230	Brand
PARICALCITOL	PAF	RICALCITOL CAP 1 MCG	30905070000110	Generic
DOXERCALCIFEROL	DO:	XERCALCIFEROL CAP 0.5 MCG	30905040000105	Generic
DOXERCALCIFEROL	DO	XERCALCIFEROL CAP 1 MCG	30905040000110	Generic

Generic

30905070000140

DOXERCALCIFEROL	DOXERCALCIFEROL CAP 2.5 MCG	30905040000120	Generic
PARICALCITOL	PARICALCITOL CAP 2 MCG	30905070000120	Generic
PARICALCITOL	PARICALCITOL CAP 4 MCG	30905070000140	Generic

1 - Dose-adjusted trial and failure (unable to achieve parathyroid hormone level goals), contraindication, or intolerance with calcitriol.

2. Revision History

Date	Notes
11/6/2023	New program

Vivjoa (Oteseconazole)
So the followage was to deplay to Treeling to be the strong or come or stand any best to be provided to come the strong of the s

Prior Authorization Guideline

Guideline ID	GL-131407
Guideline Name	Vivjoa (Oteseconazole)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Na	ame: Vivjoa			
Approval Length		12 month(s)		
Guideline Type		Prior Authorization-IL and MN Plans Only		
Product Name	Generic Na	me	GPI	Brand/Generic
VIVJOA	OTESECONA WEEKS)	ZOLE CAP THERAPY PACK 150 MG (12	1140805000B220	Brand

Approval Criteria

1 - Current diagnosis of vulvovaginal candidiasis with positive KOH test

2 - History of recurrent vulvovaginal candidiasis with ≥ 3 episodes of vulvovaginal candidiasis in the past 12 months

AND

3 - Person is not of reproductive potential (i.e. postmenopausal, permanent infertility, tubal ligation, etc.)

AND

4 - Trial and failure, contraindication, or intolerance of maintenance fluconazole therapy (at least 6 months)

Product Name: Vivjoa	
Approval Length	3 month(s)
Guideline Type	Prior Authorization-All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
VIVJOA	OTESECONAZOLE CAP THERAPY PACK 150 MG (12 WEEKS)	1140805000B220	Brand

Approval Criteria

1 - Current diagnosis of vulvovaginal candidiasis with positive KOH test

AND

2 - History of recurrent vulvovaginal candidiasis with ≥ 3 episodes of vulvovaginal candidiasis in the past 12 months

3 - Person is not of reproductive potential (i.e. postmenopausal, permanent infertility, tubal ligation, etc.)

AND

4 - Trial and failure, contraindication, or intolerance of maintenance fluconazole therapy (at least 6 months)

2. Revision History

Date	Notes
10/24/2023	New Program

Vowst (Fecal microbiota spores, live-brp)k)
(3) Characteristic Child to the majority and with the property of the contract	

Prior Authorization Guideline

Guideline ID	GL-143523	
Guideline Name	Vowst (Fecal microbiota spores, live-brpk)	
Formulary	Quartz	

Guideline Note:

Effective Date:	4/1/2024
P&T Approval Date:	7/18/2023
P&T Revision Date:	

1. Criteria

Product Name: Vowst				
Approval L	Length 12 month (s) with a fill count = 1			
Guideline	e Type Prior Authorization			
Product Generic Name		GPI	Brand/Generic	
VOWST	FECAL MICR	OBIOTA SPORES, LIVE-BRPK CAPS	52522020100120	Brand

Approval Criteria

1 - One of the following:

- **1.1** Both of the following:
- 1.1.1 Diagnosis of at least 2 recurrent* episodes of Clostridioides difficile (C diff) infection (≥3 C diff infection episodes)

1.1.2 C diff infection is refractory to standard antibiotic therapy (i.e., has received vancomycin or fidaxomicin therapy with previous episodes)

OR

1.2 Diagnosis of recurrent* C. diff episode after previous treatment with fecal microbiota therapy

AND

2 - Has a positive stool test for toxigenic C diff from a recent stool sample

AND

- **3** Prescribed by or in consultation with one of the following:
 - Infectious Disease specialist
 - Gastroenterologist

AND

4 - Member is 18 years or older

Notes	*Recurrent defined as recurrence of diarrhea and positive C diff test w	
	ithin 8 weeks after treatment of prior episode	

2. Revision History

Date	Notes
2/29/2024	New Program

Vyndaqel, Vyndamax (tafamidis)			
(2) The find the grown and displace for the find the local county count and death of public the constitutions.			

Prior Authorization Guideline

Guideline ID	GL-131932
Guideline Name	Vyndaqel, Vyndamax (tafamidis)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: (Vyndaqel, Vyndamax		
Approval Length 12 month(s)		
Therapy Stage Initial Authorization		
Guideline Type Prior Authorization		

Product Name	Generic Name	GPI	Brand/Generic
VYNDAQEL	TAFAMIDIS MEGLUMINE (CARDIAC) CAP 20 MG	40550080200120	Brand
VYNDAMAX	TAFAMIDIS CAP 61 MG	40550080000120	Brand

Approval Criteria

1 - Diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM)

AND

2 - Prescribed by, or in consultation with, Cardiology, transthyretin amyloidosis (ATTR) specialist, or medical geneticist

AND

3 - Age ≥ 18

AND

4 - New York Heart Association (NYHA) functional class I, II, or III heart failure

AND

5 - No previous history of heart transplantation or estimated glomerular filtration rate less than 25mL per minute per 1.73m2 of body-surface area

Product Name: (Vyndaqel, Vyndamax		
Approval Length 12 month(s)		
Therapy Stage Reauthorization		
Guideline Type Prior Authorization		

Product Name	Generic Name	GPI	Brand/Generic
VYNDAQEL	TAFAMIDIS MEGLUMINE (CARDIAC) CAP 20 MG	40550080200120	Brand
VYNDAMAX	TAFAMIDIS CAP 61 MG	40550080000120	Brand

Approval Criteria

1 - Diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM)
AND
2 - Prescribed by, or in consultation with, Cardiology, transthyretin amyloidosis (ATTR) specialist, or medical geneticist
AND
3 - Age ≥ 18
AND
4 - New York Heart Association (NYHA) functional class I, II, or III heart failure
AND
5 - No previous history of heart transplantation or estimated glomerular filtration rate less than 25mL per minute per 1.73m2 of body-surface area
AND
6 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.
AND
7 - Individual has not progressed to NYHA Class IV heart failure.
2. Revision History

Notes

Date

10/16/2023	New Program

Xcopri (cenobamate)				
The behavior and continued to the term of the continued o				

Prior Authorization Guideline

Guideline ID	GL-127849
Guideline Name	Xcopri (cenobamate)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Xcopri		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type Step Therapy - IL and MN Plans		

Product Name	Generic Name	GPI	Brand/Generic
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 12.5 MG & 14 X 25 MG	7212001000B720	Brand
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 50 MG & 14 X 100 MG	7212001000B725	Brand
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 150 MG & 14 X 200 MG	7212001000B730	Brand
XCOPRI	CENOBAMATE TAB PACK 100 MG & 150 MG TABS (250 MG DAILY DOSE)	7212001000B738	Brand

XCOPRI	CENOBAMATE TAB PACK 150 MG & 200 MG TABS (350 MG DAILY DOSE)	7212001000B740	Brand
XCOPRI	CENOBAMATE TAB 50 MG	72120010000320	Brand
XCOPRI	CENOBAMATE TAB 100 MG	72120010000325	Brand
XCOPRI	CENOBAMATE TAB 150 MG	72120010000330	Brand
XCOPRI	CENOBAMATE TAB 200 MG	72120010000335	Brand

- **1** Trial and failure of at least two preferred anticonvulsants:

 - lamotrigine levetiracetam
 - carbamazepine
 - valproate
 - oxcarbazepine

 - gabapentin pregabalin topiramate
 - phenytoin zonisamide

 - primidone

Product Name: Xcopri			
Approval Length	12 month(s)		
Therapy Stage	Therapy Stage Reauthorization		
Guideline Type Step Therapy - IL and MN Plans			

Product Name	Generic Name	GPI	Brand/Generic
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 12.5 MG & 14 X 25 MG	7212001000B720	Brand
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 50 MG & 14 X 100 MG	7212001000B725	Brand
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 150 MG & 14 X 200 MG	7212001000B730	Brand
XCOPRI	CENOBAMATE TAB PACK 100 MG & 150 MG TABS (250 MG DAILY DOSE)	7212001000B738	Brand
XCOPRI	CENOBAMATE TAB PACK 150 MG & 200 MG TABS (350 MG DAILY DOSE)	7212001000B740	Brand

XCOPRI	CENOBAMATE TAB 50 MG	72120010000320	Brand
XCOPRI	CENOBAMATE TAB 100 MG	72120010000325	Brand
XCOPRI	CENOBAMATE TAB 150 MG	72120010000330	Brand
XCOPRI	CENOBAMATE TAB 200 MG	72120010000335	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Product Name: Xcopri		
Approval Length	12/31/2039	
Guideline Type	Step Therapy - All plans except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic	
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 12.5 MG & 7212001000B720 Brand 14 X 25 MG			
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 50 MG & 7212001000B725 Brand 14 X 100 MG			
XCOPRI	CENOBAMATE TAB TITRATION PACK 14 X 150 MG & 14 X 200 MG	7212001000B730	Brand	
XCOPRI	CENOBAMATE TAB PACK 100 MG & 150 MG TABS (250 MG DAILY DOSE)	7212001000B738	Brand	
XCOPRI	CENOBAMATE TAB PACK 150 MG & 200 MG TABS (350 MG DAILY DOSE)	7212001000B740	Brand	
XCOPRI	CENOBAMATE TAB 50 MG	72120010000320	Brand	
XCOPRI	CENOBAMATE TAB 100 MG	72120010000325	Brand	
XCOPRI	CENOBAMATE TAB 150 MG	72120010000330	Brand	
XCOPRI	CENOBAMATE TAB 200 MG	72120010000335	Brand	

Approval Criteria

- 1 Trial and failure of at least two preferred anticonvulsants:
 - lamotrigine
 - levetiracetam
 - carbamazepine

- valproate oxcarbazepine gabapentin pregabalin topiramate phenytoin zonisamide primidone

2. Revision History

Date	Notes
8/25/2023	New Program

Xdemvy
(3) The Manufacture for things, or There is no based as a most of a minute and place to proper the controlled frame.

Prior Authorization Guideline

Guideline ID	GL-135582
Guideline Name	Xdemvy
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Xdemvy				
Approval Length 2 month(s)				
Guideline Type		Prior Authorization – All plans excep	ot IL and MN plans	•
Dreshust Conside Name		Duorad/Conoria		

Product Name	Generic Name	GPI	Brand/Generic
XDEMVY	LOTILANER OPHTH SOLN 0.25%	86106050002020	Brand

Approval Criteria

- **1** Diagnosis of demodex blepharitis with all of the following:
 - Presence of erythema of the upper eyelid margin

• Presence of mites upon examination of eyelashes by light microscopy OR presence of collarettes on slit lamp examination

AND

2 - Member is 18 years of age or older

Product Name: Xdemvy				
Approval Length 12 month(s)				
Therapy Stage		Initial Authorization		
Guideline Type		Prior Authorization- IL and MN plans	3	
Droduct	Canaria Nama		CDI	Prond/Conorio

Product Name	Generic Name	GPI	Brand/Generic
XDEMVY	LOTILANER OPHTH SOLN 0.25%	86106050002020	Brand

Approval Criteria

- **1** Diagnosis of demodex blepharitis with all of the following:
 - Presence of erythema of the upper eyelid margin
 - Presence of mites upon examination of eyelashes by light microscopy OR presence of collarettes on slit lamp examination

AND

2 - Member is 18 years of age or older

Product Name: Xdemvy				
Approval Length		12 month(s)		
Therapy Stage		Reauthorization		
Guideline Type		Prior Authorization- IL and MN plans		
Product Name	Generic Name		GPI	Brand/Generic
XDEMVY	LOTILANER OPHTH SOLN 0.25%		86106050002020	Brand

- 1 Diagnosis of demodex blepharitis with all of the following:
 - Presence of erythema of the upper eyelid margin
 - Presence of mites upon examination of eyelashes by light microscopy OR presence of collarettes on slit lamp examination

AND

2 - Member is 18 years of age or older

AND

- 3 One of the following:
- **3.1** At least 11 months has elapsed since previous treatment with lotilaner (Xdemvy)

OR

3.2 Person is established on therapy and has not completed the initial 6 week treatment course

2. Revision History

Date	Notes
11/6/2023	New Program

Xeljanz (tofacitinib)				
The hand and provide the control of				

Prior Authorization Guideline

Guideline ID	GL-144983
Guideline Name	Xeljanz (tofacitinib)
Formulary	Quartz

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	10/16/2013
P&T Revision Date:	10/16/2023

1. Criteria

Product Name: Xeljanz, Xeljanz ER				
Diagnosis		Moderate to Severely Active Psoriatic Arthritis (PsA)		
Approval Length 12/31/2039				
Guideline ⁻	Prior Authorization - All Plans Except IL and MN Plans		8	
Product Name	Generic Name GPI Brand/Ger		Brand/Generic	
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT) 66603		66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT) 66603065100330 Bra		Brand	
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT) Brand		Brand	

XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

1 - Diagnosis of moderate to severely active psoriatic arthritis

AND

- **2** Prescribed by or in consultation with one of the following:
 - dermatologist
 - rheumatologist

AND

- **3** One of the following:
- **3.1** Both of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xeljanz, Xeljanz ER	
Diagnosis	Moderate to Severely Active Psoriatic Arthritis (PsA)
Approval Length	12 month(s)
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active psoriatic arthritis

AND

- 2 Prescribed by or in consultation with one of the following:
 - dermatologist
 - rheumatologist

AND

- 3 One of the following:
 - **3.1** Both of the following:

- **3.1.1** Submission of medical records (e.g., chart notes) documenting at least ONE of the following:
 - actively inflamed joints
 - axial disease
 - active skin, nail, or scalp psoriasis involvement
 - dactylitis
 - enthesitis

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xeljanz, Xeljanz ER	
Diagnosis Moderate to Severely Active Rheumatoid Arthritis	
Approval Length	12/31/2039
Guideline Type Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Notes	**Absolute contraindications to methotrexate are pregnancy, nursing,
	alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm
	bocytopenia or significant anemia, or hypersensitivity to methotrexate

Product Name: Xeljanz, Xeljanz ER	
Diagnosis	Moderate to Severely Active Rheumatoid Arthritis
Approval Length	12 month(s)

Guideline Type		Prior Authorization - IL and MN Plans		
Product Name	Generic Name		GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)		66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)		66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)		66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)		66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)		66603065107550	Brand

1 - Diagnosis of moderate to severely active rheumatoid arthritis (RA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
 - **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Notes	**Absolute contraindications to methotrexate are pregnancy, nursing,
	alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm
	bocytopenia or significant anemia, or hypersensitivity to methotrexate

Product Name: Xeljanz, Xeljanz ER		
Diagnosis	Polyarticular Juvenile Idiopathic Arthritis (PJIA)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

Approval Criteria

1 - Diagnosis of polyarticular juvenile idiopathic arthritis (PJIA)

AND

2 - Prescribed by or in consultation with a rheumatologist

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Notes	**Absolute contraindications to methotrexate are pregnancy, nursing,
	alcoholism, alcoholic liver disease or other chronic liver disease, immu
	nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm
	bocytopenia or significant anemia, or hypersensitivity to methotrexate

Product Name: Xeljanz, Xeljanz ER		
Diagnosis Polyarticular Juvenile Idiopathic Arthritis (PJIA)		
Approval Length	12 month(s)	
Guideline Type Prior Authorization - IL and MN Plans		

П	Product Name	Generic Name	GPI	Brand/Generic
	XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand

XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

1 - Diagnosis of polyarticular juvenile idiopathic arthritis (PJIA)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Submission of medical records (e.g., chart notes) documenting a 3-month trial and failure, intolerance, or contraindication to ONE of the following:
 - methotrexate (MTX)**
 - leflunomide
 - hydroxychloroquine
 - sulfasalazine

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)		
Notes	**Absolute contraindications to methotrexate are pregnancy, nursing, alcoholism, alcoholic liver disease or other chronic liver disease, immu nodeficiency syndromes, bone marrow hyperplasia, leukopenia, throm bocytopenia or significant anemia, or hypersensitivity to methotrexate	

Product Name: Xeljanz, Xeljanz ER		
Diagnosis	Ankylosing Spondylitis (AS)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

1 - Diagnosis of ankylosing spondylitis (AS)

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

- **3** One of the following:
- **3.1** All of the following:

3.1.1 Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xeljanz, Xeljanz ER	
Diagnosis Ankylosing Spondylitis (AS)	
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

Approval Criteria

1 - Diagnosis of ankylosing spondylitis (AS)

2 - Prescribed by or in consultation with a rheumatologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Minimum duration of a 1-month trial and failure, contraindication, or intolerance of scheduled prescription doses of TWO different NSAIDs (e.g., naproxen, nabumetone, diclofenac)

AND

3.1.2 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xeljanz, Xeljanz ER			
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)		
Approval Length	12/31/2039		
Guideline Type Prior Authorization - All Plans Except IL and MN Plans			S
Product Generic Name GPI		GPI	Brand/Generic

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand

XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

1 - Diagnosis of moderate to severely active Crohn's disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
- **3.1** One of the following:
- **3.1.1** Member is considered high-risk based on at least one of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease
 - Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)

OR

- **3.1.2** Both of the following:
- **3.1.2.1** Member is considered low-risk

3.1.2.2 One of the following:

- Trial and failure, contraindication, or intolerance to two conventional therapies (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with one conventional therapy
- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

AND

3.1.3 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xeljanz, Xeljanz ER		
Diagnosis	Moderate to Severely Active Crohn's Disease (CD)	
Approval Length	12 month(s)	
Guideline Type	Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand

XELJANZ TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand
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1 - Diagnosis of moderate to severely active Crohn's disease (CD)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- **3** One of the following:
- **3.1** One of the following:
- **3.1.1** Member is considered high-risk based on at least one of the following characteristics:
 - Age less than 30 years at diagnosis
 - Extensive anatomic involvement
 - Perianal and/or severe rectal disease
 - Deep ulcers
 - Prior surgical resection
 - Stricturing and/or penetrating behavior
 - Fistulizing disease
 - Extraintestinal manifestations of inflammation (e.g., uveitis, erythema nodosum, pyoderma gangrenosum, spondyloarthropathy)

OR

- **3.1.2** Both of the following:
- **3.1.2.1** Member is considered low-risk

AND

3.1.2.2 One of the following:

- Trial and failure, contraindication, or intolerance to two conventional therapies (e.g., azathioprine, balsalazide, corticosteroids, mesalamine, mercaptopurine, methotrexate, sulfasalazine)
- Inadequate disease control or inability to achieve remission after an adequate trial of 3 months with one conventional therapy
- Demonstrated steroid dependence
- Conventional therapy clinically inappropriate based on location of disease

3.1.3 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xeljanz, Xeljanz ER		
Diagnosis	Moderate to Severely Active Ulcerative Colitis (UC)	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

2 - Prescribed by or in consultation with a gastroenterologist

AND

- **3** One of the following:
- **3.1** All of the following:
- **3.1.1** Member is considered high-risk based on at least one of the following characteristics:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High CRP and ESR
 - Steroid-requiring disease
 - History of hospitalization
 - C. difficile infection
 - CMV infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

AND

3.1.3 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xeljanz, Xeljanz ER

Diagnosis	Moderate to Severely Active Ulcerative Colitis (UC)
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
XELJANZ	TOFACITINIB CITRATE TAB 5 MG (BASE EQUIVALENT)	66603065100320	Brand
XELJANZ	TOFACITINIB CITRATE TAB 10 MG (BASE EQUIVALENT)	66603065100330	Brand
XELJANZ	TOFACITINIB CITRATE ORAL SOLN 1 MG/ML (BASE EQUIVALENT)	66603065102020	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 11 MG (BASE EQUIVALENT)	66603065107530	Brand
XELJANZ XR	TOFACITINIB CITRATE TAB ER 24HR 22 MG (BASE EQUIVALENT)	66603065107550	Brand

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

- 3 One of the following:
 - **3.1** All of the following:
 - **3.1.1** Member is considered high-risk based on at least one of the following characteristics:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High CRP and ESR
 - Steroid-requiring disease History of hospitalization

 - C. difficile infection

• CMV infection

AND

3.1.2 Trial and failure, contraindication, or intolerance to a short course (2 to 4 weeks) of oral corticosteroids

AND

3.1.3 Trial and failure, contraindication, or intolerance to at least one TNF agent (e.g. adalimumab, etanercept, etc.) as per package label

OR

3.2 Continuation of prior therapy with tofacitinib, verified by paid claims or medical records (e.g. chart notes)

2. Definitions

Definition	Description
Inadequate Disease Control of UC/CD:	Worsening of baseline symptoms (i.e. bowel frequency, presence of blood, abdominal pain or tenderness, fever, etc.), extraintestinal manifestations (i.e. fatigue, joint pain, skin rash, and ocular symptoms), laboratory assessment (i.e. Creactive protein (CRP), hemoglobin, ESR white blood count (WBC), albumin, platelets, fecal calprotectin, etc.) and/or recent endoscopy results demonstrating ongoing inflammation
Steroid Dependence:	Demonstrated steroid dependence (defined as equivalent to prednisone 10mg daily for >3 months) with the inability to taper or when tapering of dose leads to loss of symptom control
Inflammatory status: Signs/Symptoms/Labs/Endoscopy for diagnosis	-Bloody diarrhea, weight loss, tenesmus, urgency, abdominal pain, fever, joint swelling/redness, localized abdominal tenderness, anemia, cutaneous signs -CBC,

	CMP, CRP, ESR, stool cultures, C difficile assay, fecal calprotectin -endoscopy, colonoscopy, sigmoidoscopy
Ulcerative Colitis Disease Severity:	Based on the degree of presentation of the signs and symptoms and change in baseline inflammatory status Moderate disease - more than four stools per day with minimal signs of toxicity, anemia, abdominal pain, low grade fever Severe disease - more than six bloody stools per day, fever, tachycardia, anemia, elevated ESR or CRP
Crohn's Disease Classification:	Stricturing - narrowing of bowel that may cause bowel obstruction; Penetrating - fistulae may form between bowel and other structures; Inflammatory - nonstricturing, nonpenetrating - inflammation without strictures or fistula

3. Revision History

Date	Notes
4/9/2024	Guideline Update.

Xenl	Xenleta (Lefamulin)				
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Prior Authorization Guideline

Guideline ID	GL-129632	
Guideline Name	Xenleta (Lefamulin)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Xenleta				
Approval Length *See Note				
Guideline Type Prior Authorization - IL and MN Plans				
Product Name			GPI	Brand/Generic
XENLETA	LEFAMULIN ACETATE TAB 600 MG		16240040100320	Brand

Approval Criteria

1 - Person has been receiving drug during hospitalization and needs to complete the course of therapy as an outpatient.

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- 2 Both of the following:
- **2.1** Outpatient treatment of bacterial resistant strains as ordered by or in consultation with an Infectious Disease Specialist

2.2 Report of susceptibilities documenting resistance to preferred alternatives

OR

3 - (Illinois plans only) – the requested FDA approved drug is being used for the long-term treatment of tick-borne disease.

Notes	Approval Length-12 months Fill Limit- 1 Fill
	The Entire TT III

Product Name: Xenleta		
Approval Length One fill		
Guideline Type Prior Authorization - All plans except IL and MN		

Product Name	Generic Name	GPI	Brand/Generic
XENLETA	LEFAMULIN ACETATE TAB 600 MG	16240040100320	Brand

Approval Criteria

1 - Person has been receiving drug during hospitalization and needs to complete the course of therapy as an outpatient.

OR

2 - Outpatient treatment of bacterial resistant strains as ordered by or in consultation with an Infectious Disease Specialist

3 - Report of susceptibilities documenting resistance to preferred alternatives

2. Revision History

Date	Notes
10/25/2023	New program

Xermelo (telotristat)		
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Prior Authorization Guideline

Guideline ID	GL-131938
Guideline Name	Xermelo (telotristat)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Xermelo		
Approval Length	12 month(s)	
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
XERMELO	TELOTRISTAT ETHYL TAB 250 MG (AS TELOTRISTAT ETIPRATE)	52570075100330	Brand

Approval Criteria

1 - Diagnosis of diarrhea secondary to carcinoid syndrome

2 - Age greater than or equal to 18 years

AND

3 - 3-month trial and failure (≥ 4 bowel movements per day) with a somatostatin analog such as octreotide, lanreotide, or pasireotide.

AND

4 - Used in combination with a somatostatin analog

Product Name: Xermelo		
12 month(s)		
Reauthorization		
Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
XERMELO	TELOTRISTAT ETHYL TAB 250 MG (AS TELOTRISTAT ETIPRATE)	52570075100330	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Xermelo				
Approval Length		12/31/2039		
Guideline Type		Prior Authorization-All plans except IL and MN		
Product Generic Na Name		me	GPI	Brand/Generic

_	LOTRISTAT ETHYL TAB 250 MG (AS TELOTRISTAT IPRATE)	52570075100330	Brand
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1 - Diagnosis of diarrhea secondary to carcinoid syndrome

AND

2 - Age greater than or equal to 18 years

AND

3 - 3-month trial and failure (≥ 4 bowel movements per day) with a somatostatin analog such as octreotide, lanreotide, or pasireotide.

AND

4 - Used in combination with a somatostatin analog

2. Revision History

Date	Notes
10/31/2023	New program

Xolair (Omalizumab)				
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Prior Authorization Guideline

Guideline ID	GL-145159
Guideline Name	Xolair (Omalizumab)
Formulary	Quartz

Guideline Note:

Effective Date:	4/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Na	Product Name: Xolair			
Diagnosis		Asthma		
Approval Length		12 month(s)		
Therapy Stage Init		Initial Authorization		
Guideline Type		Prior Authorization - IL and MN Plar	ns Only	
Product Generic Na Name		me	GPI	Brand/Generic

XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED 4460306000E520 Brail SYRINGE 150 MG/ML		Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

1 - Diagnosis of moderate-to-severe persistent allergic asthma as defined by Global Initiative for Asthma (GINA) Global Strategy for Asthma Management and Prevention Guidelines (Step 5)

AND

2 - Member is 6 years or older

AND

3 - Serum IgE level ≥ 30 international units/mL

AND

4 - Positive skin tests or in vitro reactivity to common aeroallergens (e.g. dust mites, pet dander, cockroaches)

AND

5 - Member is a non-smoker or smoking cessation therapy has been recommended

- 6 One of the following:
- **6.1** Member has not well controlled or poorly controlled asthma despite episodic use of systemic corticosteroids or at least 3 months of medium to high-dose inhaled corticosteroids (ICS) in combination with long acting beta2 agonist (LABA) or leukotriene modifiers

OR

- **6.2** Member has one of the following adverse effects from medium to high dose ICS or long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients greater than 40 years of age
 - Glaucoma
 - Recurrent Thrush
 - Dysphonia
 - Growth inhibition, after evaluation by Endocrine Consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

7 - Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

Product Name: Xolair		
Diagnosis Asthma		
Approval Length	12/31/2039	
Guideline Type	Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

1 - Diagnosis of moderate-to-severe persistent allergic asthma as defined by Global Initiative for Asthma (GINA) Global Strategy for Asthma Management and Prevention Guidelines (Step 5)

AND

2 - Member is 6 years or older

AND

3 - Serum IgE level ≥ 30 international units/mL

AND

4 - Positive skin tests or in vitro reactivity to common aeroallergens (e.g. dust mites, pet dander, cockroaches)

AND

5 - Member is a non-smoker or smoking cessation therapy has been recommended

AND

- 6 One of the following:
 - 6.1 Member has not well controlled or poorly controlled asthma despite episodic use of

systemic corticosteroids or at least 3 months of medium to high-dose inhaled corticosteroids (ICS) in combination with long acting beta2 agonist (LABA) or leukotriene modifiers

OR

- **6.2** Member has one of the following adverse effects from medium to high dose ICS or long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients greater than 40 years of age
 - Glaucoma
 - Recurrent Thrush
 - Dysphonia
 - Growth inhibition, after evaluation by Endocrine Consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

7 - Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

Due de et Name e Valain		
Product Name: Xolair		
Diagnosis	Urticaria	
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type	Prior Authorization - IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

1 - Diagnosis of chronic (at least 3 months), refractory urticaria

AND

- **2** Member has tried and failed both of the following:
 - Scheduled, high dose non-sedating antihistamines at least one short course of corticosteroids

AND

3 - Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

Product Name: Xolair	
Diagnosis Urticaria	
Approval Length 12/31/2039	
Guideline Type Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

1 - Diagnosis of chronic (at least 3 months), refractory urticaria

AND

- 2 Member has tried and failed both of the following:
 - Scheduled, high dose non-sedating antihistamines at least one short course of corticosteroids

AND

3 - Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

Product Name: Xolair	
Diagnosis	Immunotherapy
Approval Length	12 month(s)
Guideline Type Prior Authorization	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

Approval Criteria

1 - Prescribed by an allergist

2 - Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

Product Name: Xolair	
Diagnosis	Polyps
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type Prior Authorization - IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

Approval Criteria

1 - Diagnosis of chronic rhinosinusitis with nasal polyposis

AND

2 - All of the following:

- At least eight weeks of moderate to severe nasal congestion/blockage/obstruction OR diminished sense of smell or rhinorrhea
- Submission of medical records (e.g., chart notes) Documented nasal polyps by direct exam, endoscopy, or sinus CT scan (i.e., nasal polyp score five out of eight)

No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation
AND
3 - Prescribed by, or in consultation with, a specialist experienced in the treatment of nasal polyps (e.g., Otolaryngologist, Allergist)
AND
4 - Trial and failure, contraindication, or intolerance to one of the following:
 Greater than or equal to 2 nasal steroid sprays (i.e. failed two nasal sprays) IM injections for polyps with one previous nasal spray
AND
5 - Trial and failure, contraindication, or intolerance to one of the following:
 Oral corticosteroids for nasal polyps Prior surgery for nasal polyps greater than six months ago
AND
6 - Requested medication will be used in combination with a nasal corticosteroid medication
AND
7 - Requested medication will not used in combination with other biologic therapies (e.g. penralizumab, dupilumab, mepolizumab)
AND
3 - Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

Product Name: Xolair	
Diagnosis	Polyps
Approval Length	12/31/2039
Guideline Type Prior Authorization - All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

1 - Diagnosis of chronic rhinosinusitis with nasal polyposis

AND

- 2 All of the following:
 - At least eight weeks of moderate to severe nasal congestion/blockage/obstruction OR diminished sense of smell or rhinorrhea
 - Submission of medical records (e.g., chart notes) Documented nasal polyps by direct exam, endoscopy, or sinus CT scan (i.e., nasal polyp score five out of eight)
 - No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation

AND

3 - Prescribed by, or in consultation with, a specialist experienced in the treatment of nasal polyps (e.g., Otolaryngologist, Allergist)

- 4 Trial and failure, contraindication, or intolerance to one of the following:
 - Greater than or equal to 2 nasal steroid sprays (i.e. failed two nasal sprays)
 - IM injections for polyps with one previous nasal spray

AND

- **5** Trial and failure, contraindication, or intolerance to one of the following:
 - Oral corticosteroids for nasal polyps
 - Prior surgery for nasal polyps greater than six months ago

AND

6 - Requested medication will be used in combination with a nasal corticosteroid medication

AND

7 - Requested medication will not used in combination with other biologic therapies (e.g. benralizumab, dupilumab, mepolizumab)

AND

8 - Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

Product Name: Xolair		
Diagnosis	All Indications	
Approval Length	12 month(s)	
Therapy Stage	Reauthorization	
Guideline Type	uideline Type Prior Authorization - IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand

1 - Submission of medial records documenting a positive clinical response to therapy and improvement in disease state from previous 12 months

AND

- **2** Submission of medical records (e.g., chart notes) documenting from the previous 12 months in improvement to one of the following:
 - Decreased frequency of corticosteroid use to treat or prevent an exacerbation
 - Reductions in symptom exacerbation frequency or intensity
 - Decreased frequency of unscheduled clinic, urgent care or emergency department visits due to asthma
 - Increase in percent predicted FEV1 from pre-treatment baseline
 - Increase in percent predicted FEV1 from pre-treatment baseline
 - Reduction use of ICS, leukotriene or beta agonist therapy
 - Improvement in nasal polyposis score

2. Background

Benefit/Coverage/Program Information

Outcome Measure values for uncontrolled asthma

Measure	Not Well Controlled	Very Poorly Controlled
Baseline symptoms	> 2 days/week	Throughout the day
(outside of exacerbation)		
Nighttime awakening	1-3 times/week	≥ 4 times/week
Interference with normal activity	Some limitation	Extremely limited
Short acting beta agonist use for symptom control	> 2 days/week	Several times per day
FEV1	60-80% predicted or personal best	< 60% predicted or personal best
Asthma exacerbations requiring oral steroids ≥ 2 times in the past year	Yes	Yes
Asthma Control Test (ACT)	16-19	≤ 15

Xolair (Omalizumab)		
Security commission with an arrive and come of the left to produce commission of the		

Prior Authorization Guideline

Guideline ID	GL-144993 Xolair (Omalizumab)	
Guideline Name		
Formulary	Quartz	

Guideline Note:

Effective Date:	5/1/2024
P&T Approval Date:	1/16/2013
P&T Revision Date:	7/18/2023

1. Criteria

Product Name: Xolair				
Diagnosis Asthma				
Approval Length 12 month(s)				
Guideline ¹	Туре	Prior Authorization - IL and MN Plan	ns Only	
Product Name	Generic Name		GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML		4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML		4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML		4460306000D530	Brand

XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

1	- Diagnosi	is of moderate	-to-severe p	persistent	allergic ast	hma as o	defined by	Global Initia	ative
fc	or Asthma ((GINA) Global	Strategy fo	r Asthma	Manageme	ent and P	Prevention (Guidelines	(Step
5	5)								

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2 - Member is 6 years or older

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Serum IgE level ≥ 30 international units/mL

AND

3.1.2 Positive skin tests or in vitro reactivity to common aeroallergens (e.g. dust mites, pet dander, cockroaches)

AND

3.1.3 Member is a non-smoker or smoking cessation therapy has been recommended

AND

3.1.4 One of the following:

3.1.4.1 Member has not well controlled or poorly controlled asthma despite episodic use of systemic corticosteroids or at least 3 months of medium to high-dose inhaled corticosteroids (ICS) in combination with long acting beta2 agonist (LABA) or leukotriene modifiers

OR

- **3.1.4.2** Member has one of the following adverse effects from medium to high dose ICS or long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients greater than 40 years of age
 - Glaucoma
 - Recurrent Thrush
 - Dysphonia
 - Growth inhibition, after evaluation by Endocrine Consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

3.1.5 Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

OR

3.2 Continuation of prior therapy with omalizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xolair				
Diagnosis	Asthma			
Approval Length	12/31/2039			
Guideline Type	e Prior Authorization – All Plans Except IL and MN Plans Only			
Product Generic Name GPI Brand/G			Brand/Generic	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand

XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

1 - Diagnosis of moderate-to-severe persistent allergic asthma as defined by Global Initiative for Asthma (GINA) Global Strategy for Asthma Management and Prevention Guidelines (Step 5)

AND

2 - Member is 6 years or older

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Serum IgE level ≥ 30 international units/mL

AND

3.1.2 Positive skin tests or in vitro reactivity to common aeroallergens (e.g. dust mites, pet dander, cockroaches)

AND

3.1.3 Member is a non-smoker or smoking cessation therapy has been recommended

- **3.1.4** One of the following:
- **3.1.4.1** Member has not well controlled or poorly controlled asthma despite episodic use of systemic corticosteroids or at least 3 months of medium to high-dose inhaled corticosteroids (ICS) in combination with long acting beta2 agonist (LABA) or leukotriene modifiers

OR

- **3.1.4.2** Member has one of the following adverse effects from medium to high dose ICS or long-term risks of adverse effects from high dose ICS or oral corticosteroids:
 - Cataracts in patients greater than 40 years of age
 - Glaucoma
 - Recurrent Thrush
 - Dysphonia
 - Growth inhibition, after evaluation by Endocrine Consult
 - Diagnosis of osteoporosis, treatment resistant to FDA approved osteoporosis treatment

AND

3.1.5 Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

OR

3.2 Continuation of prior therapy with omalizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xolair			
Diagnosis	Urticaria		
Approval Length	12 month(s)		
Guideline Type	Prior Authorization - IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

1 - Diagnosis of chronic (at least 3 months), refractory urticaria

AND

- 2 One of the following:
- 2.1 All of the following:
- **2.1.1** Member has tried and failed both of the following:
 - Scheduled, high dose non-sedating antihistamines
 - At least one short course of corticosteroids

AND

2.1.2 Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

OR

2.2 Continuation of prior therapy with omalizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xolair		
Diagnosis	Urticaria	
Approval Length	12/31/2039	
Guideline Type	Prior Authorization – All Plans Except IL and MN Plans	

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

1 - Diagnosis of chronic (at least 3 months), refractory urticaria

AND

- 2 One of the following:
- 2.1 All of the following:
- **2.1.1** Member has tried and failed both of the following:
 - Scheduled, high dose non-sedating antihistamines
 - At least one short course of corticosteroids

AND

2.1.2 Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

OR

2.2 Continuation of prior therapy with omalizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xolair	
Diagnosis	Immunotherapy
Approval Length	12 month(s)
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

Approval Criteria

1 - Prescribed by an allergist

AND

- **2** One of the following:
- **2.1** Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

OR

2.2 Continuation of prior therapy with omalizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xolair	
Diagnosis	Polyps
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

Approval Criteria

1 - Diagnosis of chronic rhinosinusitis with nasal polyposis

AND

2 - Prescribed by, or in consultation with, a specialist experienced in the treatment of nasal polyps (e.g., Otolaryngologist, Allergist)

AND

3 - One of the following: **3.1** All of the following: **3.1.1** All of the following: At least eight weeks of moderate to severe nasal congestion/blockage/obstruction OR diminished sense of smell or rhinorrhea Submission of medical records (e.g., chart notes) • Documented nasal polyps by direct exam, endoscopy, or sinus CT scan (i.e., nasal polyp score five out of eight) No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation **AND 3.1.2** Trial and failure, contraindication, or intolerance to one of the following: Greater than or equal to 2 nasal steroid sprays (i.e. failed two nasal sprays) IM injections for polyps with one previous nasal spray AND **3.1.3** Trial and failure, contraindication, or intolerance to one of the following: Oral corticosteroids for nasal polyps Prior surgery for nasal polyps greater than six months ago

AND

3.1.4 Requested medication will be used in combination with a nasal corticosteroid medication

AND

3.1.5 Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

OR

3.2 Continuation of prior therapy with omalizumab, verified by paid claims or medical records (e.g. chart notes)

Product Name: Xolair	
Diagnosis	Polyps
Approval Length	12/31/2039
Guideline Type	Prior Authorization – All Plans Except IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 75 MG/0.5ML	4460306000D510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 150 MG/ML	4460306000D520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN AUTO- INJECTOR 300 MG/2ML	4460306000D530	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 75 MG/0.5ML	4460306000E510	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 150 MG/ML	4460306000E520	Brand
XOLAIR	OMALIZUMAB SUBCUTANEOUS SOLN PREFILLED SYRINGE 300 MG/2ML	4460306000E530	Brand

Approval Criteria

1 - Diagnosis of chronic rhinosinusitis with nasal polyposis

AND

2 - Prescribed by, or in consultation with, a specialist experienced in the treatment of nasal polyps (e.g., Otolaryngologist, Allergist)

AND

- 3 One of the following:
 - **3.1** All of the following:

3.1.1 All of the following:

- At least eight weeks of moderate to severe nasal congestion/blockage/obstruction OR diminished sense of smell or rhinorrhea
- Submission of medical records (e.g., chart notes) Documented nasal polyps by direct exam, endoscopy, or sinus CT scan (i.e., nasal polyp score five out of eight)
- No chronic or acute infection requiring systemic treatment within two weeks before therapy initiation

AND

- **3.1.2** Trial and failure, contraindication, or intolerance to one of the following:
 - Greater than or equal to 2 nasal steroid sprays (i.e. failed two nasal sprays)
 - IM injections for polyps with one previous nasal spray

AND

- **3.1.3** Trial and failure, contraindication, or intolerance to one of the following:
 - Oral corticosteroids for nasal polyps
 - Prior surgery for nasal polyps greater than six months ago

AND

3.1.4 Requested medication will be used in combination with a nasal corticosteroid medication

AND

3.1.5 Prescriber attestation that the initial six (6) months of therapy must be done in the clinic setting by a healthcare professional

OR

3.2 Continuation of prior therapy with omalizumab, verified by paid claims or medical records (e.g. chart notes)

2. Background

Benefit/Coverage/Program Information				
Outcome Measure values for uncontrolled asthma				
Measure	Not Well Controlled	Very Poorly Controlled		
Baseline symptoms (outside of exacerbation)	> 2 days/week	Throughout the day		
Nighttime awakening	1-3 times/week	≥ 4 times/week		
Interference with normal activity	Some limitation	Extremely limited		
Short acting beta agonist use for symptom control	> 2 days/week	Several times per day		
FEV1	60-80% predicted or personal best	< 60% predicted or personal best		
Asthma exacerbations requiring oral steroids ≥ 2 times in the past year	Yes	Yes		
Asthma Control Test (ACT)	16-19	≤ 15		

3. Revision History

Date	Notes
4/9/2024	Guideline Update.

Χ	Xuriden (Uridine triacetate)			
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Prior Authorization Guideline

Guideline ID	GL-131951
Guideline Name	Xuriden (Uridine triacetate)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Xuriden	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
XURIDEN	URIDINE TRIACETATE ORAL GRANULES PACKET 2 GM	30903875203020	Brand

Approval Criteria

1 - Diagnosis of hereditary orotic aciduria

Product Name: Xuriden	
Approval Length 12 month(s)	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization-IL and MN Plans Only	

Product Name	Generic Name	GPI	Brand/Generic
XURIDEN	URIDINE TRIACETATE ORAL GRANULES PACKET 2 GM	30903875203020	Brand

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug and that there has been a response to therapy (improvement in hematologic counts and urine orotic acid levels).

Product Name: Xuriden	
Approval Length 3 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
XURIDEN	URIDINE TRIACETATE ORAL GRANULES PACKET 2 GM	30903875203020	Brand

Approval Criteria

1 - Diagnosis of hereditary orotic aciduria

Product Name: Xuriden			
Approval Length		12/31/2039	
Therapy Stage Re		Reauthorization	
Guideline Type P		Prior Authorization-All plans except IL and MN	
Product Generic Name Name		GPI	Brand/Generic

XURIDEN URIDINE TRIACETATE ORAL GRANULES PACKE	ET 2 30903875203020	Brand
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1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug and that there has been a response to therapy (improvement in hematologic counts and urine orotic acid levels).

Date	Notes
10/31/2023	New program

,	Xyrem (sodium oxybate)				
	Substanting-most in dialogue. Took may be made using a made using factor to a more flower budge.				

Guideline ID	GL-131921
Guideline Name	Xyrem (sodium oxybate)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2023
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1. Criteria

Product Name: Generic Sodium oxybate		
Approval Length 12 month(s)		
Therapy Stage	Initial Authorization	
Guideline Type Prior Authorization-IL and MN Plans Only		

Product Name	Generic Name	GPI	Brand/Generic
SODIUM OXYBATE	SODIUM OXYBATE ORAL SOLUTION 500 MG/ML	62450060202020	Generic

Approval Criteria

1 - Diagnosis of cataplexy in narcolepsy or excessive daytime sleepiness in narcolepsy

2 - Trial and failure, contraindication, or intolerance to one preferred narcolepsy medication (e.g. modafinil, armodafinil)

Product Name: Generic Sodium oxybate	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization-IL and MN Plans Only

Product Name	Generic Name	GPI	Brand/Generic
SODIUM OXYBATE	SODIUM OXYBATE ORAL SOLUTION 500 MG/ML	62450060202020	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months documenting an improvement in sleepiness symptoms.

Product Name: Generic Sodium oxybate	
Approval Length	3 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization-All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
SODIUM OXYBATE	SODIUM OXYBATE ORAL SOLUTION 500 MG/ML	62450060202020	Generic

Approval Criteria

1 - Diagnosis of cataplexy in narcolepsy or excessive daytime sleepiness in narcolepsy

2 - Trial and failure, contraindication, or intolerance to one preferred narcolepsy medication (e.g. modafinil, armodafinil)

Product Name: Generic Sodium oxybate		
Approval Length	12/31/2039	
Therapy Stage	Reauthorization	
Guideline Type	Prior Authorization-All plans except IL and MN	

Product Name	Generic Name	GPI	Brand/Generic
SODIUM OXYBATE	SODIUM OXYBATE ORAL SOLUTION 500 MG/ML	62450060202020	Generic

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months documenting an improvement in sleepiness symptoms.

Date	Notes
10/31/2023	New program

Zeposia (Ozanimod)
(a) The little frequency to think you have been const, consist, a date that his joint in a consist and consists are consistent and consists and consi

Guideline ID	GL-143573
Guideline Name	Zeposia (Ozanimod)
Formulary	Quartz

Guideline Note:

Effective Date:	4/1/2024
P&T Approval Date:	1/17/2023
P&T Revision Date:	7/18/2023

Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Zeposia			
Diagnosis Ulcerative colitis (UC)			
Approval Length 12/31/2039			
Guideline Type	Prior Authorization - Applies to ALL	Prior Authorization - Applies to ALL plans except IL and MN	
Product Generic Name	Name	GPI	Brand/Generic

ZEPOSIA 7-DAY STARTER PACK	OZANIMOD CAP PACK 4 X 0.23 MG & 3 X 0.46 MG	6240705020B210	Brand
ZEPOSIA STARTER KIT	OZANIMOD CAP PACK 4 X 0.23 MG & 3 X 0.46 MG & 21 X 0.92 MG	6240705020B215	Brand
ZEPOSIA	OZANIMOD HCL CAP 0.92 MG	62407050200120	Brand

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with Gastroenterologist

AND

- 3 One of the following:
- **3.1** All of the following:
- **3.1.1** Member is considered high risk based on at least ONE of the following characteristics:
 - Extensive colitis
 - Deep ulcers
 - Age less than 40 years
 - High CRP and ESR
 - Steroid-requiring disease
 - History of hospitalization (due to UC)
 - C. difficile infection
 - CMV infection

AND

3.1.2 Trial and failure, intolerance, or contraindication to a short course (2-4 weeks) of oral corticosteroids

- **3.1.3** Trial and failure, intolerance or contraindication to TWO of the following:
 - adalimumab
 - upadacitinib
 - golimumab
 - ustekinumab
 - tofacitinib/ER

OR

3.2 Continuation of prior therapy with ozanimod, verified by paid claims or medical records (e.g. chart notes)

Product Name: Zeposia	
Diagnosis	Ulcerative colitis (UC)
Approval Length	12 month(s)
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ZEPOSIA 7-DAY STARTER PACK	OZANIMOD CAP PACK 4 X 0.23 MG & 3 X 0.46 MG	6240705020B210	Brand
ZEPOSIA STARTER KIT	OZANIMOD CAP PACK 4 X 0.23 MG & 3 X 0.46 MG & 21 X 0.92 MG	6240705020B215	Brand
ZEPOSIA	OZANIMOD HCL CAP 0.92 MG	62407050200120	Brand

Approval Criteria

1 - Diagnosis of moderate to severely active ulcerative colitis (UC)

AND

2 - Prescribed by or in consultation with Gastroenterologist
AND
3 - One of the following:
3.1 All of the following:
3.1.1 Member is considered high risk based on at least ONE of the following characteristics:
 Extensive colitis Deep ulcers Age less than 40 years High CRP and ESR Steroid-requiring disease History of hospitalization (due to UC) C. difficile infection CMV infection
AND
3.1.2 Trial and failure, intolerance, or contraindication to a short course (2-4 weeks) of oral corticosteroids
AND
3.1.3 Trial and failure, intolerance or contraindication to TWO of the following:
 adalimumab upadacitinib golimumab ustekinumab tofacitinib/ER
OR
3.2 Continuation of prior therapy with ozanimod, verified by paid claims or medical records (e.g. chart notes)

Product Name: Zeposia	
Diagnosis	Multiple Sclerosis
Approval Length	12 month(s)
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
ZEPOSIA 7-DAY STARTER PACK	OZANIMOD CAP PACK 4 X 0.23 MG & 3 X 0.46 MG	6240705020B210	Brand
ZEPOSIA STARTER KIT	OZANIMOD CAP PACK 4 X 0.23 MG & 3 X 0.46 MG & 21 X 0.92 MG	6240705020B215	Brand
ZEPOSIA	OZANIMOD HCL CAP 0.92 MG	62407050200120	Brand

1 - Diagnosis of a relapsing form of multiple sclerosis

AND

2 - Prescribed by or in consultation with a Neurologist

AND

- **3** One of the following:
- **3.1** Paid claims or submission of medical records (e.g. chart notes) show there have been adequate trials of ALL appropriate formulary therapeutic alternatives and there was (a) inadequate clinical response, (b) inappropriate clinical response, (c) intolerance or (d) allergy to those medications

OR

3.2 An exception to the formulary may be considered when ALL appropriate formulary therapeutic alternatives have not been tried and there is submission of medical record documentation (e.g. chart notes) demonstrating that ALL appropriate formulary therapeutic alternatives will be (a) ineffective, (b) less effective or (c) will result in adverse effects

Date	Notes
2/29/2024	New Program

Zokinvy (Lonafarnib)			
The State Progress of Subspace The Board and control, or State Sta			

Guideline ID	GL-129641
Guideline Name	Zokinvy (Lonafarnib)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Zokinvy	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
ZOKINVY	LONAFARNIB CAP 50 MG	99463045000120	Brand
ZOKINVY	LONAFARNIB CAP 75 MG	99463045000130	Brand

Approval Criteria

1 - Diagnosis of Hutchinson-Gilford progeria syndrome OR other FDA approved diagnosis

2 - Prescribed by, or in consultation with, a specialist in the treatment of progeria or related-syndromes

Product Name: Zokinvy	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization

Product Name	Generic Name	GPI	Brand/Generic
ZOKINVY	LONAFARNIB CAP 50 MG	99463045000120	Brand
ZOKINVY	LONAFARNIB CAP 75 MG	99463045000130	Brand

Approval Criteria

1 - The prescriber must provide clinical documentation from an office visit in the preceding 12 months that use of the drug has slowed the disease progression and function is improved relative to the expected natural course of the disease.

Date	Notes
10/6/2023	New Program

Zontivity (vorapaxar)		
(3) had been from your in deplayer. The first factor count, county a detect like little joins his countly actuales.		

Guideline ID	GL-132750
Guideline Name	Zontivity (vorapaxar)
Formulary	Quartz

Guideline Note:

Effective Date:	1/1/2024
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Note:

Member new to the plan (as evidenced by coverage effective date of less than or equal to 90 days) who initiated therapy using a manufacturer-sponsored free drug program, provider samples, and/or vouchers will go through initial criteria, otherwise for continuation of therapy for new to plan, reauthorization criteria applies

1. Criteria

Product Name: Zontivity				
Approval Length		12/31/2039		
Guideline Type		Prior Authorization - All plans except IL and MN Plans		
Product Name	Generic Na	me	GPI	Brand/Generic
ZONTIVITY	VORAPAXAR EQUIVALENT	SULFATE TAB 2.08 MG (BASE)	85155780300320	Brand

- **1** Diagnosis of one of the following:
 - Peripheral Arterial Disease (PAD)
 - History of myocardial infarction (MI)

AND

2 - Prescribed by or in consultation with a Cardiologist

AND

3 - Submission of medical records (e.g., chart notes) documenting increased risk of thrombotic cardiovascular events despite being on combination therapy with BOTH aspirin and P2Y12 therapy (e.g., clopidogrel, ticagrelor, or prasugrel)

Product Name: Zontivity	
Approval Length	12 month(s)
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ZONTIVITY	VORAPAXAR SULFATE TAB 2.08 MG (BASE EQUIVALENT)	85155780300320	Brand

Approval Criteria

- **1** Diagnosis of one of the following:
 - Peripheral Arterial Disease (PAD)
 - History of myocardial infarction (MI)

AND

2 - Prescribed by or in consultation with a Cardiologist

AND

3 - Submission of medical records (e.g., chart notes) documenting increased risk of thrombotic cardiovascular events despite being on combination therapy with BOTH aspirin and P2Y12 therapy (e.g., clopidogrel, ticagrelor, or prasugrel)

Product Name: Zontivity	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Prod Nam		Generic Name	GPI	Brand/Generic
ZON	TIVITY	VORAPAXAR SULFATE TAB 2.08 MG (BASE EQUIVALENT)	85155780300320	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
9/7/2023	2024 New Implementation

Zoryve (roflumilast cream)		
The birth regions in the last tension most, usual, a state shell had been been unash an indicated as		

Guideline ID GL-131913	
Guideline Name	Zoryve (roflumilast cream)
Formulary • Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

duct Name: Zoryve			
ength	12/31/2039		
Гуре	Prior Authorization – All plans except IL and MN plans		
Generic Name		GPI	Brand/Generic
ROFLUMILAST CREAM 0.3%		90250045003720	Brand
	ength ype Generic Na	ength 12/31/2039 Type Prior Authorization – All plans except Generic Name	ength 12/31/2039 Type Prior Authorization – All plans except IL and MN plans Generic Name GPI

Approval Criteria

1 - Diagnosis of psoriasis

2 - 12 years or older

AND

3 - Prescribed by, or in consultation with a dermatologist, or other specialist in the treatment psoriasis

AND

- 4 One of the following:
 - Trial and failure, contraindication, or intolerance to one preferred high or super-high potency topical corticosteroid (see the formulary at QuartzBenefits.com for a complete listing of options)
 - Person has facial or other sensitive area involvement and a trial and failure, contraindication, or intolerance to one preferred non steroid therapy (e.g., calcipotriene, retinoids)

Product Name: Zoryve	
Approval Length 12 month(s)	
Therapy Stage Initial Authorization	
Guideline Type Prior Authorization- II and MN plans	

Product Name	Generic Name	GPI	Brand/Generic
ZORYVE	ROFLUMILAST CREAM 0.3%	90250045003720	Brand

Approval Criteria

1 - Diagnosis of psoriasis

2 - 12 years or older

AND

3 - Prescribed by, or in consultation with a dermatologist, or other specialist in the treatment psoriasis

AND

- 4 One of the following:
 - Trial and failure, contraindication, or intolerance to one preferred high or super-high potency topical corticosteroid (see the formulary at QuartzBenefits.com for a complete listing of options)
 - Person has facial or other sensitive area involvement and a trial and failure, contraindication, or intolerance to one preferred non steroid therapy (e.g., calcipotriene, retinoids)

Product Name: Zoryve	
Approval Length 12/31/2039	
Therapy Stage Reauthorization	
Guideline Type Prior Authorization- II and MN plans	

Product Name	Generic Name	GPI	Brand/Generic
ZORYVE	ROFLUMILAST CREAM 0.3%	90250045003720	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug

Date	Notes
10/31/2023	New Program

4	Ztlido (Lidocaine Patch)						
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Guideline ID GL-129640		
Guideline Name	Ztlido (Lidocaine Patch)	
Formulary	Quartz	

Guideline Note:

Effective Date:	1/1/2024
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1. Criteria

Product Name: Ztlido	
Approval Length 12 month(s)	
Therapy Stage	Initial Authorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ZTLIDO	LIDOCAINE PATCH 1.8% (36 MG)	90850060005910	Brand

Approval Criteria

1 - Person with a diagnosis of post-herpetic neuralgia

2 - Trial and failure, contraindication or intolerance of an equivalent dose of generic lidocaine 5% transdermal patches

AND

3 - Trial and failure, contraindication or intolerance to at least one other preferred drug with evidence for reducing symptoms of post-herpetic neuralgia symptoms

AND

4 - The prescriber provides an evidence-based clinical rationale for why different results from those seen with generic 5% lidocaine patches would be expected

OR

5 - (Minnesota plans only) – the person has stage four metastatic cancer and the requested drug is being used to treat cancer-related pain

Product Name: Ztlido	
Approval Length	12 month(s)
Therapy Stage	Reauthorization
Guideline Type	Prior Authorization - IL and MN Plans

Product Name	Generic Name	GPI	Brand/Generic
ZTLIDO	LIDOCAINE PATCH 1.8% (36 MG)	90850060005910	Brand

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that within the past 12 months the member is continuing therapy with the requested drug.

Product Name: Ztlido	
Approval Length 12/31/2039	
Guideline Type	Prior Authorization - All plans except IL and MN

Product Name	Generic Name	GPI	Brand/Generic
ZTLIDO	LIDOCAINE PATCH 1.8% (36 MG)	90850060005910	Brand

1 - Person with a diagnosis of post-herpetic neuralgia

AND

2 - Trial and failure, contraindication or intolerance of an equivalent dose of generic lidocaine 5% transdermal patches

AND

3 - Trial and failure, contraindication or intolerance to at least one other preferred drug with evidence for reducing symptoms of post-herpetic neuralgia symptoms

AND

4 - The prescriber provides an evidence-based clinical rationale for why different results from those seen with generic 5% lidocaine patches would be expected

Date	Notes
10/6/2023	New Program