



# 2026 PRIOR AUTHORIZATION CRITERIA

UCare Connect + Medicare (SNBC) (HMO D-SNP)
UCare's Minnesota Senior Health Options (MSHO) (HMO D-SNP)

UCare requires your provider to get prior authorization for certain drugs. This means that you'll need to get approval from us before you fill your prescriptions. If you don't get approval, UCare may not cover the drug.

UCare's MSHO and UCare Connect + Medicare (HMO D-SNP) are health plans that contract with both Medicare and the Minnesota Medical Assistance (Medicaid) program to provide benefits of both programs to enrollees. Enrollment in UCare's MSHO and UCare Connect + Medicare depends on contract renewal.

Effective: 01/01/2026

H5937\_5248\_072022\_C\_8 H2456\_5248\_072022 accepted

# Toll free 1-800-203-7225, TTY 1-800-688-2534

Attention. If you need free help interpreting this document, call the above number.

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ملاحظة: إذا أردت مساعدة مجانية لترجمة هذه الوثيقة، اتصل على الرقم أعلاه.

သတိ။ ဤစာရက်စာတမ်းအားအခမဲ့ဘာသာပြန်ပေးခြင်း အကူအညီလိုအပ်ပါက၊ အထက်ပါဖုန်းနံပါတ်ကိုခေါ် ဆိုပါ။

កំណត់សំគាល់ ។ បើអ្នកត្រូវការជំនួយក្នុងការបកប្រែឯកសារនេះដោយឥតគិតថ្លៃ សូមហៅទូរសព្ទតាមលេខខាងលើ ។

請注意,如果您需要免費協助傳譯這份文件,請撥打上面的電話號碼。

Attention. Si vous avez besoin d'une aide gratuite pour interpréter le présent document, veuillez appeler au numéro ci-dessus.

Thov ua twb zoo nyeem. Yog hais tias koj xav tau kev pab txhais lus rau tsab ntaub ntawv no pub dawb, ces hu rau tus najnpawb xov tooj saum toj no.

ပဉ်သူဉ်ပဉ်သးဘဉ်တက္၊ ဖဲနမ္၊်လိဉ်ဘဉ်တ၊မၤစားကလိုလာတ၊ကကျိုးထံဝဲဒဉ်လံဉ် တီလံဉ်မီတခါအံၤန္ဉာ,ကိုးဘဉ် လီတဲစိနိုါဂံ၊လာထးအံၤန္ဉ်တက္၊

알려드립니다. 이 문서에 대한 이해를 돕기 위해 무료로 제공되는 도움을 받으시려면 위의 전화번호로 연락하십시오.

ໂປຣດຊາບ. ຖ້າຫາກ ທ່ານຕ້ອງການການຊ່ວຍເຫຼືອໃນການແປເອກະສານນີ້ຟຣີ, ຈົ່ງ ໂທຣໄປທີ່ໝາຍເລກຂ້າງເທີງນີ້.

Hubachiisa. Dokumentiin kun tola akka siif hiikamu gargaarsa hoo feete, lakkoobsa gubbatti kenname bilbili.

Внимание: если вам нужна бесплатная помощь в устном переводе данного документа, позвоните по указанному выше телефону.

Digniin. Haddii aad u baahantahay caawimaad lacag-la'aan ah ee tarjumaadda (afcelinta) qoraalkan, lambarka kore wac.

Atención. Si desea recibir asistencia gratuita para interpretar este documento, llame al número indicado arriba.

Chú ý. Nếu quý vị cần được giúp đỡ dịch tài liệu này miễn phí, xin gọi số bên trên.

# **Civil Rights Notice**

**Discrimination is against the law. UCare** does not discriminate on the basis of any of the following:

- race
- color
- national origin
- creed
- religion
- sexual orientation
- public assistance status

- age
- disability (including physical or mental impairment)
- sex (including sex stereotypes and genderidentity)
- marital status

- political beliefs
- medical condition
- health status
- receipt of health care services
- claims experience
- medical history
- genetic information

You have the right to file a discrimination complaint if you believe you were treated in a discriminatory way by UCare. You can file a complaint and ask for help filing a complaint in person or by mail, phone,

**UCare** 

Attn: Appeals and Grievances

PO Box 52

Minneapolis, MN 55440-0052 Toll Free: 1-800-203-7225 TTY: 1-800-688-2534

Fax: 612-884-2021 Email: cag@ucare.org

fax, or email at:

Auxiliary Aids and Services: UCare provides auxiliary aids and services, like qualified interpreters or information in accessible formats, free of charge and in a timely manner to ensure an equal opportunity to participate in our health care programs. Contact UCare at 612-676-3200 (voice) or 1-800-203-7225 (voice), 612-676-6810 (TTY), or 1-800-688-2534 (TTY).

Language Assistance Services: UCare provides translated documents and spoken language interpreting, free of charge and in a timely manner, when language assistance services are necessary to ensure limited English speakers have meaningful access to our information and services. Contact UCare at 612-676-3200 (voice) or 1-800-203-7225 (voice), 612-676-6810 (TTY), or 1-800-688-2534 (TTY).

# **Civil Rights Complaints**

You have the right to file a discrimination complaint if you believe you were treated in a discriminatory way by UCare. You may also contact any of the following agencies directly to file a discrimination complaint.

### U.S. Department of Health and Human Services Office for Civil Rights (OCR)

You have the right to file a complaint with the OCR, a federal agency, if you believe you have been discriminated against because of any of the following:

race

age

religion (in some cases)

color

disability

national origin

sex

Contact the OCR directly to file a complaint:

Office for Civil Rights

U.S. Department of Health and Human Services

Midwest Region

233 N. Michigan Avenue, Suite 240

Chicago, IL 60601

Customer Response Center: Toll-free: 800-368-1019

TDD Toll-free: 800-537-7697 Email: ocrmail@hhs.gov

### Minnesota Department of Human Rights (MDHR)

In Minnesota, you have the right to file a complaint with the MDHR if you have been discriminated against because of any of the following:

race

creed

• sex

national origin

sexual orientation

religion

color

marital status

 public assistance status

disability

Contact the MDHR directly to file a complaint:

Minnesota Department of Human Rights

540 Fairview Avenue North, Suite 201

St. Paul, MN 55104

651-539-1100 (voice)

800-657-3704 (toll-free)

711 or 800-627-3529 (MN Relay)

651-296-9042 (fax)

Info.MD4R@state.mn.us (email)

### Minnesota Department of Human Services (DHS)

You have the right to file a complaint with DHS if you believe you have been discriminated against in our health care programs because of any of the following:

- race
- color
- national origin
- religion (in some cases)
- age
- disability (including physical or mental impairment)
- sex (including sex stereotypes and gender identity)

Complaints must be in writing and filed within 180 days of the date you discovered the alleged discrimination. The complaint must contain your name and address and describe the discrimination you are complaining about. We will review it and notify you in writing about whether we have authority to investigate. If we do, we will investigate the complaint.

DHS will notify you in writing of the investigation's outcome. You have the right to appeal if you disagree with the decision. To appeal, you must send a written request to have DHS review the investigation outcome. Be brief and state why you disagree with the decision. Include additional information you think is important.

If you file a complaint in this way, the people who work for the agency named in the complaint cannot retaliate against you. This means they cannot punish you in any way for filing a complaint. Filing a complaint in this way does not stop you from seeking out other legal or administrative actions.

Contact **DHS** directly to file a discrimination complaint:

Civil Rights Coordinator
Minnesota Department of Human Services
Equal Opportunity and Access Division
P.O. Box 64997
St. Paul, MN 55164-0997
651-431-3040 (voice) or use your preferred relay service

### **ACNE AGENTS\_NVT\_2026**

# **MEDICATION(S)**

TRETINOIN CREAM (0.025 %, 0.05 %, 0.1 %)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**ACTIMMUNE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# ADBRY\_(UCARE)\_2026

### MEDICATION(S)

**ADBRY** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For atopic dermatitis (initial requests): Two of the following were ineffective or not tolerated: a) a medium to very high potency topical steroid, b) a topical calcineurin inhibitor OR c) an oral immunosuppressant. For atopic dermatitis (continuation requests): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

For atopic dermatitis: Prescribed by, or in consultation with, an allergist, immunologist, or dermatologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For atopic dermatitis (initial requests): Member has moderate to severe atopic dermatitis defined as: 1) One of the following: a) Body surface area involvement of 10 percent or more or b) Involvement of the face, head, neck, hands, feet, groin, or intertriginous areas and 2) At least two of the following: a) Intractable pruritus (itching), b) Cracking and oozing/bleeding of skin or c) Impaired activities of daily living. For atopic dermatitis (all requests): Will not be used in combination with another targeted immunomodulator product for the prescribed indication.

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

YES

Rending CMS Approval

### ADCIRCA\_NVT\_2026

# **MEDICATION(S)**

ALYQ, TADALAFIL (PAH)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

Autheterization.

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERF

N/A

### PREREQUISITE THERAPY REQUIRED

**ADEMPAS** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For all indications: Diagnosis confirmed by right heart catheterization. For pulmonary arterial hypertension: Both of the following were ineffective or not tolerated: one endothelial receptor antagonist (ambrisentan, bosentan or macitentan (Opsumit)) and one phosphodiesterase-5 inhibitor (sildenafil or tadalafil). For persistent/recurrent chronic thromboembolic pulmonary hypertension (WHO Group 4): Trial of other agents not required.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

N/A

# **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

EVEROLIMUS 10 MG TAB, EVEROLIMUS 2 MG TAB SOL, EVEROLIMUS 2.5 MG TAB, EVEROLIMUS 3 MG TAB SOL, EVEROLIMUS 5 MG TAB, EVEROLIMUS 5 MG TAB SOL, **EVEROLIMUS 7.5 MG TAB** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# CINS REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

# AIMOVIG\_(UCARE)\_2026

# **MEDICATION(S)**

**AIMOVIG** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For migraine prevention (initial requests): Member has 4 or more migraine days per month for the previous 3 months or longer. For migraine prevention (continuation requests): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**AKEEGA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ALECENSA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### ALPHA1\_NVT\_2026

### **MEDICATION(S)**

PROLASTIN-C 1000 MG/20ML SOLUTION

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Both of the following: 1) Diagnosis of congenital alpha1-antitrypsin deficiency is confirmed by both of the following: A) circulating baseline alpha1-antitrysin level is below the standard protective threshold (less than 11 micromol/L or less than 50 mg per deciliter by nephelometry) and B) high risk alpha1-antitrypsin deficiency genotype (SS, SZ, ZZ, or null/null) and 2) Prescriber attests that member does not have IgA deficiency with known anti-IgA antibody.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a pulmonologist

# **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

**ALUNBRIG** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ALYFTREK** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ARCALYST** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ARIKAYCE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, an infectious disease specialist or pulmonologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

### **ATTRUBY NVT 2026**

### MEDICATION(S)

**ATTRUBY** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For cardiomyopathy of transthyretin-mediated amyloidosis (ATTR-CM)(initial requests): Diagnosis confirmed by one of the following: A) Cardiac biopsy with positive Congo Red staining and ATTR confirmation by mass spectrometry or immunofluorescence staining or B) Non-invasive testing demonstrating all of the following: i) Serum kappa/lambda free light chain ratio 0.26 to 1.65, ii) Absence of monoclonal protein via serum protein immunofixation, iii) Absence of monoclonal protein via urine protein immunofixation and iv) Myocardial uptake of 99mTc-PYP demonstrated by a greater than 1.5 heart-to-contralateral ratio or grade 2 or greater visual evidence. For ATTR-CM (continuation requests): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

# PRESCRIBER RESTRICTION

For ATTR-CM: Prescribed by, or in consultation with, a cardiologist or other provider experienced in the treatment of ATTR-CM.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For ATTR-CM (all requests): Will not be used in combination with inotersen (Tegsedi), patisiran (Onpattro), vutrisiran (Amvuttra) or tafamidis (Vyndagel/Vyndamax).

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

N/A

Pending CMS Approval

**AUGTYRO** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### AUSTEDO\_NVT\_2026

### **MEDICATION(S)**

AUSTEDO, AUSTEDO XR, AUSTEDO XR PATIENT TITRATION 12 & 18 & 24 & 30 MG TBER THPK

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### OFF LABEL USES

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For tardive dyskinesia (initial requests): A) One of the following: i) Member has failed to respond to a change in current antidopaminergic therapy OR ii) Member is unable to switch current antidopaminergic therapy OR iii) Member has symptoms of tardive dyskinesia and is not using antidopaminergic therapy AND B) Member has a functional disability due to tardive dyskinesia. For chorea associated with Huntington's disease (initial requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist or psychiatrist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

pending and proval

AVMAPKI FAKZYNJA CO-PACK

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**AYVAKIT** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**BALVERSA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

RUFINAMIDE

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Trial of at least one anti-epileptic medication was ineffective or not tolerated.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

### BENLYSTA\_NVT\_2026

### **MEDICATION(S)**

BENLYSTA 200 MG/ML SOLN A-INJ, BENLYSTA 200 MG/ML SOLN PRSYR

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For systemic lupus erythematosus initial requests: Two of the following were ineffective or not tolerated: a) hydroxychloroquine, b) methotrexate, c) azathioprine, d) mycophenolate or e) a corticosteroid. For all requests: Prescriber attests that member does not have severe active CNS lupus and member is not taking other biologics. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a rheumatology specialist, nephrologist, or dermatologist.

### COVERAGE DURATION

Approved for duration of 1 year.

### OTHER CRITERIA

For lupus erythematosus initial therapy: Diagnosis of active systemic lupus erythematosus is confirmed by one of the following: A) anti-double stranded DNA value greater than 30 IU/mL or B) low complement (C3/C4) or C) positive for anti-Smith antibodies. For systemic lupus erythematosus (all requests): Will not be given in combination with other biologics. For active lupus nephritis (all requests): Will not be used in combination with voclosporin (Lupkynis).

### PART B PREREQUISITE

### PREREQUISITE THERAPY REQUIRED

YES

pending chis Approval

**BESREMI** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For polycythemia vera: One of the following: A) Both of the following: i) High risk disease as defined by one of the following: a) Age 60 years or older or b) History of thrombosis and ii) Trial of hydroxyurea was ineffective, contraindicated, or not tolerated or B) Both of the following: i) Low risk as defined by both of the following: a) Age less than 60 years and b) No history of thrombosis.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**BOSULIF** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**BRAFTOVI** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**BRUKINSA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**CABOMETYX** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**CALQUENCE 100 MG TAB** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**CAPLYTA** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For schizophrenia: Two of the following were ineffective or not tolerated: a) aripiprazole, b) olanzapine, c) quetiapine, d) risperidone, e) ziprasidone, f) lurasidone, or g) asenapine. For bipolar depression: Two of the following were ineffective or not tolerated: a) lurasidone, b) quetiapine, or c) asenapine.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**CAPRELSA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**CARGLUMIC ACID** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**CAYSTON** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

TADALAFIL 2.5 MG TAB, TADALAFIL 5 MG TAB

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

### COBENFY\_NVT\_2026

### **MEDICATION(S)**

COBENFY, COBENFY STARTER PACK

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Two of the following were ineffective or not tolerated: a) aripiprazole, b) olanzapine, c) quetiapine, d) risperidone, e) ziprasidone, f) lurasidone, or g) asenapine.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

### COMETRIQ\_NVT\_2026

### **MEDICATION(S)**

COMETRIQ (100 MG DAILY DOSE), COMETRIQ (140 MG DAILY DOSE), COMETRIQ (60 MG DAILY DOSE)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

## CINIS REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

### CONTINUOUS GLUCOSE MONITORS\_UCARE 2026

### MEDICATION(S)

DEXCOM G6 RECEIVER, DEXCOM G6 SENSOR, DEXCOM G6 TRANSMITTER, DEXCOM G7 15 DAY SENSOR, DEXCOM G7 RECEIVER, DEXCOM G7 SENSOR, FREESTYLE LIBRE 14 DAY READER, FREESTYLE LIBRE 14 DAY SENSOR, FREESTYLE LIBRE 2 PLUS SENSOR, FREESTYLE LIBRE 2 READER, FREESTYLE LIBRE 3 PLUS SENSOR, FREESTYLE LIBRE 3 READER, FREESTYLE LIBRE READER

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 3 years.

### OTHER CRITERIA

For Diabetes Mellitus (Initial Requests) - Approve if the member has been treated with insulin in the past 180 days OR has a history of problematic hypoglycemia with documentation of at least one of the following: Recurrent level 2 hypoglycemic events (glucose less than 54mg/dL (3.0mmol/L) that persist despite multiple (2 or more) attempts to adjust medication(s) and/or modify the diabetes treatment plan, or, a history of one level 3 hypoglycemic event (glucose less than 54mg/dL (3.0mmol/L) characterized by altered mental and/or physical state requiring third-party assistance for treatment of hypoglycemia, AND the member (or the members caregiver) must have been properly trained on using the requested

continuous glucose monitor (CGM) as evidenced by the treating practitioner providing a prescription, AND the CGM is prescribed according to its Food and Drug Administration (FDA) indicated use, AND the prescriber has had an in-person visit or approved telehealth visit with the member within the past six months, prior to ordering the CGM, to evaluate their diabetes control. For Diabetes Mellitus (Continuation Requests) - Approve if the treating practitioner conducts an in-person or Medicareapproved telehealth visit with the member to document adherence to their CGM regimen and diabetes treatment plan every six months following the initial prescription of the CGM.

### PART B PREREQUISITE

N/A

Pending CMS Approval PREREQUISITE THERAPY REQUIRED

**COPIKTRA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### COSENTYX\_(UCARE)\_2026

### **MEDICATION(S)**

COSENTYX 150 MG/ML SOLN PRSYR, COSENTYX 75 MG/0.5ML SOLN PRSYR, COSENTYX (300 MG DOSE), COSENTYX SENSOREADY (300 MG), COSENTYX SENSOREADY PEN, COSENTYX UNOREADY

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For ankylosing spondylitis (all requests): Trial of other agents not required. For psoriatic arthritis (all requests): Trial of other agents not required. For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of at least 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For non-radiographic axial spondyloarthritis (initial requests): Trial of two non-steroidal anti-inflammatory drugs (NSAIDs) was ineffective or not tolerated. For hidradenitis suppurativa (initial requests): Member must have both of the following: a) At least 3 cysts and b) Trial of one oral antibiotic was ineffective or not tolerated. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For psoriatic arthritis, non-radiographic axial spondyloarthritis or ankylosing spondylitis: Prescribed by, or in consultation with, a rheumatology specialist. For plaque psoriasis and hidradenitis suppurativa: Prescribed by, or in consultation with, a dermatologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

**OTHER CRITERIA** 

N/A

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

YES

Pending CMS Approval

COTELLIC

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

CRESEMBA 186 MG CAP, CRESEMBA 74.5 MG CAP

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

For invasive aspergillosis: 3 months. For invasive mucormycosis: 6 months.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**CYSTADROPS** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### CYSTEAMINE\_(UCARE)\_2026

### **MEDICATION(S)**

**CYSTAGON** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

Concomitant use of Cystagon and Procysbi

### REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For nephrotic cystinosis: Prescribed by or in consultation with a nephrologist or a metabolic disease specialist (or specialist who focuses in the treatment of metabolic diseases)

### **COVERAGE DURATION**

Approved for duration of 1 year

### OTHER CRITERIA

For Nephrotic Cystinosis (initial requests): Approve if the prescriber attests the diagnosis was established by genetic testing confirming a mutation of the CTNS gene or the member has a white blood cell cystine concentration above the upper limit of the normal reference range for the reporting laboratory. For Nephrotic Cystinosis (continuation requests): Approve if the member has had a clinical benefit (e.g., decrease in white blood cell cystine levels from baseline) with the requested medication.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

Pending CMS Approval

### DARAPRIM\_(UCARE)\_2026

### **MEDICATION(S)**

**PYRIMETHAMINE 25 MG TAB** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

### DAURISMO\_NVT\_2026

### **MEDICATION(S)**

**DAURISMO** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**METYROSINE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

DIACOMIT

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Trial of at least one anti-epileptic medication was ineffective or not tolerated.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

### DRONABINOL\_NVT\_2026

### **MEDICATION(S)**

**DRONABINOL** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

## CINIS REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

Approval will be based off BvD coverage determination.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

### **DUPIXENT\_(UCARE)\_2026**

### MEDICATION(S)

**DUPIXENT** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### OFF LABEL USES

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For atopic dermatitis (initial requests): Two of the following were ineffective or not tolerated: a) A medium to very high potency topical steroid, b) A topical calcineurin inhibitor or c) An oral immunosuppressant (trial of other agents not required for patients under 2 years of age). For asthma (initial requests): History within the last year of at least one asthma exacerbation requiring one of the following, despite regular use of inhaled corticosteroids plus an additional controller(s): a) Treatment with systemic corticosteroids, b) Emergency department visit or c) Hospitalization. For nasal polyps (initial requests): Trial of a nasal corticosteroid was ineffective or not tolerated. For eosinophilic esophagitis (initial requests): Trial of topical corticosteroid was ineffective or not tolerated. For prurigo nodularis (initial requests): Trial of other agents not required. For chronic obstructive pulmonary disease (COPD)(initial requests): History, within the last year, of at least one severe or two moderate COPD exacerbations despite receiving optimized (triple therapy) maintenance therapy. For chronic spontaneous urticaria (initial requests): One of the following: a) Patient remains symptomatic despite H1 antihistamine treatment or b) Has intolerance or contraindication to H1 antihistamine treatment. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For atopic dermatitis, prurigo nodularis, or chronic spontaneous urticaria: Prescribed by, or in consultation with, an allergist, immunologist, or dermatologist. For asthma or COPD: Prescribed by, or in consultation with, an allergist, pulmonologist, or immunologist. For nasal polyps: Prescribed by, or in consultation with, an allergist, immunologist, or otolaryngologist. For eosinophilic esophagitis:

Prescribed by, or in consultation with, an allergist or gastroenterologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For atopic dermatitis (initial requests): Member has moderate to severe atopic dermatitis defined as: 1) One of the following: a) Body surface area involvement of 10 percent or more or b) Involvement of the face, head, neck, hands, feet, groin, or intertriginous areas and 2) At least two of the following: a) Intractable pruritus (itching), b) Cracking and oozing/bleeding of skin or c) Impaired activities of daily living. For asthma (initial requests): One of the following: 1) Eosinophilic phenotype with baseline blood eosinophil concentration greater than or equal to 150 cells/microliter or 2) Oral corticosteroiddependent asthma. For nasal polyps (initial requests): All of the following: a) Diagnosis of chronic rhinosinusitis with nasal polyposis, lasting at least 12 weeks, b) Bilateral nasal polyposis confirmed with sinus CT scan, and c) Moderate to severe symptoms of nasal congestion, blockage, or obstruction (such as loss of smell, rhinorrhea, or facial pain). For eosinophilic esophagitis (initial requests): Both of the following: 1) Endoscopic biopsy with at least 15 eosinophils per high-power field (hpf) and 2) Symptoms of esophageal dysfunction (e.g. dysphagia). For prurigo nodularis (initial requests): Both of the following apply: a) Diagnosis has persisted for at least 6 weeks and b) Nodules present at baseline. For COPD (initial requests): Eosinophilic phenotype with baseline blood eosinophil concentration greater than or equal to 300 cells/microliter. For all atopic dermatitis, asthma, prurigo nodularis, or COPD (all requests): Will not be used in combination with another targeted immunomodulator product for the prescribed indication.

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

### **EMGALITY\_(UCARE)\_2026**

### **MEDICATION(S)**

EMGALITY, EMGALITY (300 MG DOSE)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For migraine prevention (initial requests): Member has 4 or more migraine days per month for the previous 3 months or longer. For episodic cluster headache prophylaxis initial requests: Trial of verapamil was ineffective or not tolerated. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

### ENBREL\_(UCARE)\_2026

### MEDICATION(S)

ENBREL 25 MG/0.5ML SOLN PRSYR, ENBREL 25 MG/0.5ML SOLUTION, ENBREL 50 MG/ML SOLN PRSYR, ENBREL MINI, ENBREL SURECLICK

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For rheumatoid arthritis (initial requests): Trial of methotrexate at a dose of at least 20mg/week (or maximally tolerated dose) was ineffective or not tolerated. For polyarticular juvenile idiopathic arthritis (pJIA)(initial requests): Trial of methotrexate at a dose of at least 15 mg/week required (or maximally tolerated dose) was ineffective or not tolerated. For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of at least 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For ankylosing spondylitis (AS)(all requests): Trial of other agents not required. For psoriatic arthritis (all requests): Trial of other agents not required. For juvenile psoriatic arthritis: Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For rheumatoid arthritis, psoriatic arthritis, juvenile psoriatic arthritis, pJIA or ankylosing spondylitis: Prescribed by, or in consultation with, a rheumatology specialist. For plaque psoriasis: Prescribed by, or in consultation with, a dermatologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

**OTHER CRITERIA** 

N/A

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

YES

Pending CMS Approval

### ENDARI\_NVT\_2026

### **MEDICATION(S)**

L-GLUTAMINE 5 GM PACKET

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For initial requests: One of the following: 1) Both of the following: A) Trial of a maximally tolerated hydroxyurea dose was ineffective or not tolerated and B) Member has had at least 1 vaso-occlusive crisis in the prior 12 months, while on hydroxyurea (if applicable) or 2) Prescriber is a hematologist. For continuation requests: Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a hematologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

### **EPCLUSA\_NVT\_2026**

### **MEDICATION(S)**

SOFOSBUVIR-VELPATASVIR

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

All of the following: 1) Current HCV-RNA titer is provided, 2) One of the following: a) Member does not have cirrhosis or b) Member has compensated cirrhosis and one of the following: i) Does not have genotype 3 or ii) has genotype 3 but no NS5A resistance-associated substitution Y93H, or c) Member has decompensated cirrhosis and will receive weight-based ribavirin.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a gastroenterologist, hepatologist, infectious disease or transplant specialist.

### COVERAGE DURATION

Coverage duration of 12 to 24 weeks. Applied consistent with current AASLD-IDSA guidance.

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

### **EPIDIOLEX NVT\_2026**

### **MEDICATION(S)**

**EPIDIOLEX** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Trial of at least one anti-epileptic medication was ineffective or not tolerated.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**ERIVEDGE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### ERLEADA\_NVT\_2026

### **MEDICATION(S)**

**ERLEADA** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For metastatic castration-sensitive prostate cancer: Trial of abiraterone was ineffective or not tolerated. For nonmetastatic castration-resistant prostate cancer: Trial of other agents not required.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

### **ESBRIET NVT 2026**

### MEDICATION(S)

PIRFENIDONE 267 MG CAP, PIRFENIDONE 267 MG TAB, PIRFENIDONE 801 MG TAB

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For idiopathic pulmonary fibrosis (initial requests): Diagnosis confirmed by one of the following: A) Surgical lung biopsy or transbronchial lung cryobiopsy revealing histopathological pattern of unspecified interstitial pneumonia (UIP), B) High-resolution computed tomography (HRCT) indicates definite UIP pattern C) Both of the following: HRCT indicates possible UIP pattern and surgical lung biopsy or transbronchial lung cryobiopsy reveals a histopathological pattern of probable UIP. For idiopathic pulmonary fibrosis (continuation requests): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

For idiopathic pulmonary fibrosis: Prescribed by, or in consultation with, a pulmonologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For idiopathic pulmonary fibrosis (all requests): Will not be used in combination with other agents for the prescribed indication.

### PART B PREREQUISITE

Pending CMS Approval

### **EUCRISA\_NVT\_2026**

### **MEDICATION(S)**

**EUCRISA** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For atopic dermatitis: One of the following was ineffective or not tolerated: a) A topical corticosteroid or b) A topical calcineurin inhibitor

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

### **FANAPT NVT 2026**

### **MEDICATION(S)**

FANAPT, FANAPT TITRATION PACK A, FANAPT TITRATION PACK B, FANAPT TITRATION PACK C

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Two of the following were ineffective or not tolerated: a) aripiprazole, b) olanzapine, c) quetiapine, d) risperidone, e) ziprasidone, f) lurasidone, or g) asenapine.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

# FASENRA\_(UCARE)\_2026

### **MEDICATION(S)**

FASENRA, FASENRA PEN

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For asthma (initial requests): History within the last year of at least one asthma exacerbation requiring one of the following, despite regular use of inhaled corticosteroids plus an additional controller(s): a) treatment with systemic corticosteroids, b) emergency department visit or c) hospitalization. For eosinophilic granulomatosis with polyangiitis (EGPA)(initial requests): Trial of oral corticosteroid therapy was ineffective or not tolerated. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For asthma: Prescribed by, or in consultation with, an allergist, immunologist, or pulmonologist. For EGPA: Prescribed by, or in consultation with, a rheumatology specialist, allergist, pulmonologist, or immunologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For asthma (initial requests): Eosinophilic phenotype with baseline blood eosinophil concentration greater than or equal to 150 cells/microliter. For asthma (all requests): Will not be used in combination with another targeted immunomodulator product for the prescribed indication.

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

YES

Rending CMS Approval

**FINTEPLA** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Trial of at least one anti-epileptic medication was ineffective or not tolerated.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

### FIRMAGON\_NVT\_2026

### **MEDICATION(S)**

FIRMAGON, FIRMAGON (240 MG DOSE)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**FOTIVDA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**FRUZAQLA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

FYCOMPA 0.5 MG/ML SUSPENSION, PERAMPANEL

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For partial-onset seizures: Two of the following were ineffective or not tolerated: a) lamotrigine b) carbamazepine c) levetiracetam d) oxcarbazepine e) phenytoin f) topiramate or g) lacosamide. For primary generalized tonic-clonic seizures: Two of the following were ineffective or not tolerated: a) lamotrigine, b) levetiracetam, c) primidone or d) topiramate.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**GAVRETO** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**GILOTRIF** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# GLP-1\_AGONISTS\_(UCARE)\_2026

### **MEDICATION(S)**

MOUNJARO, OZEMPIC (0.25 OR 0.5 MG/DOSE) 2 MG/3ML SOLN PEN, OZEMPIC (1 MG/DOSE) 4 MG/3ML SOLN PEN, OZEMPIC (2 MG/DOSE), RYBELSUS, TRULICITY

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# CINIS REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

# **COVERAGE DURATION**

Approved for duration of 1 year

### OTHER CRITERIA

Continuation therapy (all FDA approved indications): Approve if the member has been using the requested medication within the past 180 days.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**GOMEKLI** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### **GROWTH HORMONES NVT 2026**

### MEDICATION(S)

OMNITROPE, SKYTROFA 11 MG CARTRIDGE, SKYTROFA 13.3 MG CARTRIDGE, SKYTROFA 3 MG CARTRIDGE, SKYTROFA 3.6 MG CARTRIDGE, SKYTROFA 4.3 MG CARTRIDGE, SKYTROFA 5.2 MG CARTRIDGE, SKYTROFA 6.3 MG CARTRIDGE, SKYTROFA 7.6 MG CARTRIDGE, SKYTROFA 9.1 MG CARTRIDGE

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Documentation is provided of failure to stimulate growth hormone secretion (peak growth hormone level of 10mcg/L or less) by one of the acceptable provocative tests.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, an endocrinologist

# **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

### HADLIMA\_(UCARE)\_2026

MEDICATION(S)

HADLIMA, HADLIMA PUSHTOUCH

PA INDICATION INDICATOR

N/A

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

### REQUIRED MEDICAL INFORMATION

For rheumatoid arthritis (initial requests): Trial of methotrexate at a dose of at least 20mg/week (or maximally tolerated dose) was ineffective or not tolerated. For polyarticular juvenile idiopathic arthritis (pJIA)(initial requests): Trial of methotrexate at a dose of at least 15 mg/week (or maximally tolerated dose) was ineffective or not tolerated. For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of at least 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For ankylosing spondylitis (AS)(all requests): Trial of other agents not required. For psoriatic arthritis (all requests): Trial of other agents not required. For ulcerative colitis (all requests): Trial of other agents not required. For hidradenitis suppurativa (initial requests): Member must have both of the following: a) At least 3 cysts and b) Trial of one oral antibiotic was ineffective or not tolerated. For uveitis (initial requests): Both of the following were ineffective or not tolerated: a) a corticosteroid and b) an immunosuppressant (methotrexate, mycophenolate mofetil, or cyclosporine). For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For rheumatoid arthritis, psoriatic arthritis, pJIA or ankylosing spondylitis: Prescribed by, or in consultation with, a rheumatology specialist. For plaque psoriasis or hidradenitis suppurativa: Prescribed by, or in consultation with, a dermatologist. For Crohn's disease and ulcerative colitis: Prescribed by, or in consultation with, a gastroenterologist. For uveitis: Prescribed by, or in consult

with, a rheumatology specialist or ophthalmologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

**OTHER CRITERIA** 

N/A

**PART B PREREQUISITE** 

N/A

Pending CMS Approval PREREQUISITE THERAPY REQUIRED

YES

### **HAE AGENTS NVT 2026**

# **MEDICATION(S)**

HAEGARDA, ICATIBANT ACETATE

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, an allergist or immunologist

### **COVERAGE DURATION**

Approved for duration of 1 years

### **OTHER CRITERIA**

For medications indicated for long-term prophylaxis (all requests): Will not be used in combination with another agent for long-term prophylaxis of hereditary angioedema attacks.

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

# HARVONI\_(UCARE)\_2026

### MEDICATION(S)

LEDIPASVIR-SOFOSBUVIR

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

All of the following: 1) Genotype is provided, 2) Current HCV-RNA titer is provided, 3) Member has one of the following: a) no cirrhosis, b) compensated cirrhosis, or c) decompensated cirrhosis, and 4) Member is intolerant to, or unable to use both of the following: a) Mavyret and b) Sofosbuvir-Velpatasvir.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a gastroenterologist, hepatologist, infectious disease or transplant specialist.

### COVERAGE DURATION

Coverage duration of 12 to 24 weeks. Applied consistent with current AASLD-IDSA guidance.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**HERNEXEOS** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**IBRANCE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**IBTROZI** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ICLUSIG** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**IDHIFA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

IMBRUVICA 140 MG CAP, IMBRUVICA 140 MG TAB, IMBRUVICA 280 MG TAB, IMBRUVICA 420 MG TAB, IMBRUVICA 70 MG CAP, IMBRUVICA 70 MG/ML SUSPENSION

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

CINIS REQUIRED MEDICAL INFORMATION

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for duration of 1 year

OTHER CRITERIA

N/A

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

**IMKELDI** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

REQUIRED MEDICAL INFORMATION

Member is unable to swallow solid dosage forms of imatinib.

AGE RESTRICTION

N/A

PRESCRIBER RESTRICT

//A

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**IMPAVIDO** 

PA INDICATION INDICATOR

N/A

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

A CINS APPROVAL **REQUIRED MEDICAL INFORMATION** 

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for 1 month.

**OTHER CRITERIA** 

N/A

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

**INCRELEX** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**INGREZZA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For tardive dyskinesia (initial requests): A) One of the following: i) Member has failed to respond to a change in current antidopaminergic therapy OR ii) Member is unable to switch current antidopaminergic therapy OR iii) Member has symptoms of tardive dyskinesia and is not using antidopaminergic therapy AND B) Member has a functional disability due to tardive dyskinesia. For chorea associated with Huntington's disease (initial requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist or psychiatrist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

pending and proval

**INLYTA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**INQOVI** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**INREBIC** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

ROPRIONAL MARIANTE MA Trial of Jakafi was ineffective or not tolerated.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**GEFITINIB** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ITOVEBI** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# ITRACONAZOLE\_(UCARE)\_2026

# **MEDICATION(S)**

ITRACONAZOLE 100 MG CAP

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for a duration of 6 months.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

GAMMAGARD, GAMMAGARD S/D LESS IGA, GAMUNEX-C, PRIVIGEN

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

Approval will be based off BvD coverage determination.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**IWILFIN** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

DEFERASIROX 180 MG TAB, DEFERASIROX 360 MG TAB, DEFERASIROX 90 MG TAB

SARRIONA

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by or in consultation with a hematologist

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**JAKAFI** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**JAYPIRCA** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# JOURNAVX\_(UCARE)\_2026

# **MEDICATION(S)**

**JOURNAVX** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

CMS REQUIRED MEDICAL INFORMATION

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for duration of 1 month

**OTHER CRITERIA** 

N/A

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

**KALYDECO** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**KERENDIA** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

KISQALI (200 MG DOSE), KISQALI (400 MG DOSE), KISQALI (600 MG DOSE), KISQALI FEMARA (200 MG DOSE), KISQALI FEMARA (400 MG DOSE), KISQALI FEMARA (600 MG DOSE)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

CINIS REQUIRED MEDICAL INFORMATION

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for duration of 1 year

OTHER CRITERIA

N/A

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

## KORLYM\_NVT\_2026

# **MEDICATION(S)**

MIFEPRISTONE 300 MG TAB

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**KOSELUGO** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**KRAZATI** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

# **KUVAN\_NVT\_2026**

### **MEDICATION(S)**

SAPROPTERIN DIHYDROCHLORIDE 100 MG PACKET, SAPROPTERIN DIHYDROCHLORIDE 500 MG PACKET

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For continuation requests: Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a medical geneticist or metabolic physician.

### **COVERAGE DURATION**

Initial approval of 3 months. Continuing therapy approved for 1 year.

## **OTHER CRITERIA**

N/A

# **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

**LAZCLUZE** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

LENVIMA (10 MG DAILY DOSE), LENVIMA (12 MG DAILY DOSE), LENVIMA (14 MG DAILY DOSE), LENVIMA (18 MG DAILY DOSE), LENVIMA (20 MG DAILY DOSE), LENVIMA (24 MG DAILY DOSE), LENVIMA (4 MG DAILY DOSE), LENVIMA (8 MG DAILY DOSE)

## **PA INDICATION INDICATOR**

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# CINIS ARRIVA REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

## **LETAIRIS NVT 2026**

# **MEDICATION(S)**

**AMBRISENTAN** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

Atheterization.

LER RESTRICTION

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

# LIDOCAINE\_PATCH\_(UCARE)\_2026

# **MEDICATION(S)**

LIDOCAINE 5% PATCH

### PA INDICATION INDICATOR

4 - All FDA-Approved Indications, Some Medically-Accepted Indications

### **OFF LABEL USES**

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERF

N/A

### PREREQUISITE THERAPY REQUIRED

## LIVTENCITY\_NVT\_2026

# **MEDICATION(S)**

LIVTENCITY

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Prescriber attests that the medication will not be used for CMV infection prophylaxis.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a hematologist, oncologist, transplant or infectious disease specialist.

# **COVERAGE DURATION**

Approved for 3 months.

### OTHER CRITERIA

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

# LONG\_ACTING\_OPIOIDS\_(UCARE)\_2026

### MEDICATION(S)

BUPRENORPHINE WEEKLY PATCH, FENTANYL 100 MCG/HR PATCH 72HR, FENTANYL 12 MCG/HR PATCH 72HR, FENTANYL 25 MCG/HR PATCH 72HR, FENTANYL 50 MCG/HR PATCH 72HR, FENTANYL 75 MCG/HR PATCH 72HR, METHADONE HCL 10 MG TAB, METHADONE HCL 10 MG/5ML SOLUTION, METHADONE HCL 5 MG TAB, METHADONE HCL 5 MG/5ML SOLUTION, MORPHINE SULFATE ER 100 MG TAB ER, MORPHINE SULFATE ER 15 MG TAB ER, MORPHINE SULFATE ER 200 MG TAB ER, MORPHINE SULFATE ER 30 MG TAB ER, MORPHINE SULFATE ER 60 MG TAB ER

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

Acute (i.e., non-chronic) pain

### REQUIRED MEDICAL INFORMATION

Pain type (chronic vs acute), prior pain medications/therapies tried, concurrent pain medications/therapies

### **AGE RESTRICTION**

N/A

# PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For pain severe enough to require daily, around-the-clock, long-term opioid treatment (initial and continuation): Approve if all of the following criteria are met: 1) member is not opioid naive, and 2) non-opioid therapies have been tried and are being used in conjunction with opioid therapy according to the prescribing physician, and 3) the prescribing physician has checked the patient's history of controlled

substance prescriptions using state prescription drug monitoring program (PDMP), and 4) the prescribing physician has discussed risks (e.g., addiction, overdose) and realistic benefits of opioid therapy with the patient, and 5) according to the prescriber physician there is a treatment plan (including goals for pain and function) in place and reassessments are scheduled at regular intervals. Patients with cancer, in hospice, sickle cell disease or who reside in a long term care facility are not required to meet above criteria.

### PART B PREREQUISITE

N/A

Pending CMS Approval PREREQUISITE THERAPY REQUIRED

YES

## LONSURF\_NVT\_2026

# **MEDICATION(S)**

**LONSURF** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

## LORBRENA NVT 2026

# **MEDICATION(S)**

**LORBRENA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

## **LUMAKRAS NVT 2026**

# **MEDICATION(S)**

**LUMAKRAS** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

LYNPARZA

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

LYTGOBI (12 MG DAILY DOSE), LYTGOBI (16 MG DAILY DOSE), LYTGOBI (20 MG DAILY DOSE)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERF

N/A

PREREQUISITE THERAPY REQUIRED

**MAVYRET** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

All of the following: 1) Current HCV-RNA titer is provided, 2) Member does not have decompensated cirrhosis, and 3) For prior treatment with a sofosbuvir-based regimen, all of the following: i) Member does not have genotype 3 and ii) No prior treatment with an NS3/4A protease inhibitor.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a gastroenterologist, hepatologist, infectious disease or transplant specialist.

# **COVERAGE DURATION**

Coverage duration of 8 to 16 weeks. Applied consistent with current AASLD-IDSA guidance.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

MEGESTROL ACETATE 40 MG/ML SUSPENSION, MEGESTROL ACETATE 400 MG/10ML SUSPENSION, MEGESTROL ACETATE 625 MG/5ML SUSPENSION, MEGESTROL ACETATE 800 MG/20ML SUSPENSION

## **PA INDICATION INDICATOR**

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# CINIS REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

MEGESTROL ACETATE 20 MG TAB, MEGESTROL ACETATE 40 MG TAB

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERF

N/A

PREREQUISITE THERAPY REQUIRED

**MEKINIST** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**MEKTOVI** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

## MIGRANAL NVT 2026

# **MEDICATION(S)**

DIHYDROERGOTAMINE MESYLATE 4 MG/ML SOLUTION

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For acute treatment of migraine: Trial of two different triptans was ineffective or not tolerated. Trial of triptans not required for patients with history of coronary artery disease, peripheral vascular disease, uncontrolled hypertension, or other vascular risk factors or disorders. Trial of second triptan not required for patients who did not tolerate initial triptan therapy.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

## **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

# MODAFINIL\_ARMODAFINIL\_(UCARE)\_2026

### MEDICATION(S)

ARMODAFINIL, MODAFINIL 100 MG TAB, MODAFINIL 200 MG TAB

### PA INDICATION INDICATOR

4 - All FDA-Approved Indications, Some Medically-Accepted Indications

### **OFF LABEL USES**

Excessive daytime sleepiness (EDS) associated with myotonic dystrophy - modafinil only. Adjunctive/augmentation for treatment of depression in adults - modafinil only. Fatigue due to multiple sclerosis - modafinil only. Idiopathic hypersomnia - modafinil only.

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

Fatigue due to multiple sclerosis and Idiopathic hypersomnia - 18 years of age and older. All others - 17 years of age and older.

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

Adjunctive/augmentation treatment for depression in adults: Approve modafinil if the member is concurrently receiving at least one other medication for the treatment of depression. Excessive daytime sleepiness associated with obstructive sleep apnea/hypopnea syndrome: Approve modafinil or armodafinil. Excessive daytime sleepiness associated with Narcolepsy: Approve if narcolepsy has been confirmed with polysomnography and a multiple sleep latency test (MSLT). Fatigue due to multiple sclerosis: Approve modafinil. Idiopathic hypersomnia: Approve modafinil. Excessive daytime sleepiness (EDS) associated with myotonic dystrophy: Approve modafinil.

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

N/A

Rending CMS Approval

**MODEYSO** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

AVONEX PEN, AVONEX PREFILLED, DIMETHYL FUMARATE 120 MG CAP DR, DIMETHYL FUMARATE 240 MG CAP DR, DIMETHYL FUMARATE STARTER PACK, FINGOLIMOD HCL, GLATIRAMER ACETATE, GLATOPA, KESIMPTA, PLEGRIDY, TERIFLUNOMIDE

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# CINIS REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

# **NEMLUVIO (UCARE) 2026**

## MEDICATION(S)

**NEMLUVIO** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For atopic dermatitis (initial requests): Two of the following were ineffective or not tolerated: a) A medium to very high potency topical steroid, b) A topical calcineurin inhibitor or c) An oral immunosuppressant. For prurigo nodularis (initial requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For atopic dermatitis or prurigo nodularis: Prescribed by, or in consultation with, an allergist, immunologist, or dermatologist.

### COVERAGE DURATION

Approved for duration of 1 year.

### OTHER CRITERIA

For atopic dermatitis (initial requests): Member has moderate to severe atopic dermatitis defined as: 1) One of the following: a) Body surface area involvement of 10 percent or more or b) Involvement of the face, head, neck, hands, feet, groin, or intertriginous areas and 2) At least two of the following: a) Intractable pruritus (itching), b) Cracking and oozing/bleeding of skin or c) Impaired activities of daily living. For prurigo nodularis (initial requests): Both of the following apply: a) Diagnosis has persisted for at least 6 weeks and b) Nodules present at baseline. For all indications (all requests): Will not be used in combination with another targeted immunomodulator product for the prescribed indication.

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

YES

Pending CMS Approval

**NERLYNX** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**SORAFENIB TOSYLATE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

# **NEXLETOL NVT 2026**

### **MEDICATION(S)**

NEXLETOL, NEXLIZET

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# DITHER CRITERIA A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

# **NEXVIAZYME UCARE 2026**

### **MEDICATION(S)**

**NEXVIAZYME** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

For Acid Alpha-Glucosidase Deficiency (Pompe Disease): 1 year of age or older

### PRESCRIBER RESTRICTION

For Acid Alpha-Glucosidase Deficiency (Pompe Disease): Prescribed by or in consultation with a geneticist, neurologist, a metabolic disorder sub-specialist, or a physician who specializes in the treatment of lysosomal storage disorders.

### **COVERAGE DURATION**

Approved for duration of 1 year.

# OTHER CRITERIA

For Acid Alpha-Glucosidase Deficiency (Pompe Disease): Approve if the member has late-onset acid alpha-glucosidase deficiency (late-onset Pompe disease) and the diagnosis is established by one of the following: a laboratory test demonstrating deficient acid alphaglucosidase activity in blood, fibroblasts, or muscle tissue or a molecular genetic test demonstrating acid alpha-glucosidase gene mutation.

### PART B PREREQUISITE

Pending CMS Approval

# NILUTAMIDE\_(UCARE)\_2026

# **MEDICATION(S)**

**NILUTAMIDE** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

For Prostate Cancer. Approve if nilutamide is used concurrently with a luteinizing hormone-releasing hormone (LHRH) agonist.

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

**NINLARO** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

DROXIDOPA

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

POSACONAZOLE 100 MG TAB DR

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# NUBEQA\_NVT\_2026

# **MEDICATION(S)**

**NUBEQA** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For metastatic hormone-sensitive prostate cancer: Trial of abiraterone was ineffective or not tolerated. For non-metastatic castration-resistant prostate cancer: Trial of other agents not required.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

# **COVERAGE DURATION**

Approved for duration of 1 year

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

# **NUEDEXTA\_NVT\_2026**

### **MEDICATION(S)**

**NUEDEXTA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For initial requests: A) Documentation is provided (in the form of chart notes or imaging) of structural neurological condition as the cause of pseudobulbar affect. For continuation requests, both of the following: A) Documentation is provided (in the form of chart notes or imaging) of structural neurological condition as the cause of pseudobulbar affect AND B) Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

# PRESCRIBER RESTRICTION

N/A

# **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**NUPLAZID** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# NURTEC\_(UCARE)\_2026

# **MEDICATION(S)**

NURTEC

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For initial requests for acute treatment of migraines: Member has tried at least two different triptan therapies (e.g., sumatriptan and rizatriptan) or has a contraindication to triptans according to the prescriber. For initial requests for the prevention of episodic migraines: Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

# **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

OCTREOTIDE ACETATE 100 MCG/ML SOLUTION, OCTREOTIDE ACETATE 1000 MCG/ML SOLUTION, OCTREOTIDE ACETATE 200 MCG/ML SOLUTION, OCTREOTIDE ACETATE 50 MCG/ML SOLUTION, OCTREOTIDE ACETATE 500 MCG/ML SOLUTION

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

CMS APPROVAL REQUIRED MEDICAL INFORMATION

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for duration of 1 year.

N/A

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

**ODOMZO** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**OFEV** 

PA INDICATION INDICATOR

N/A

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

### REQUIRED MEDICAL INFORMATION

For idiopathic pulmonary fibrosis (initial requests): Both of the following: 1) Diagnosis confirmed by one of the following: A) Surgical lung biopsy or transbronchial lung cryobiopsy revealing histopathological pattern of unspecified interstitial pneumonia (UIP), B) High-resolution computed tomography (HRCT) indicates definite UIP pattern C) Both of the following: HRCT indicates possible UIP pattern and surgical lung biopsy or transbronchial lung cryobiopsy reveals a histopathological pattern of probable UIP and 2) Trial of pirfenidone was ineffective or not tolerated. For systemic sclerosis-associated interstitial lung disease (ILD) (initial requests): All of the following: 1) Diagnosis confirmed with documentation provided of both of the following: A) HRCT scan and B) pulmonary function tests, 2) Trial of mycophenolate mofetil was ineffective or not tolerated and 3) Trial of Tyenne was ineffective or not tolerated. For chronic fibrosing ILDs with a progressive phenotype (initial requests): All of the following: 1) Disease is progressive, as defined by one of the following over the past 12 months, with no alternative explanation: A) Worsening respiratory symptoms, B) One of the following: i) Forced vital capacity (FVC) decline of 5% or more or ii) Absolute decline in, and diffusing capacity of, the lung for carbon monoxide (corrected for hemoglobin) of 10% predicted or greater or C) Radiological evidence of disease progression and 2) Progression occurred despite treatment with one of the following: i) azathioprine, ii) cyclosporine, iii) mycophenolate mofetil, iv) tacrolimus, v) oral corticosteroids equivalent to 20 mg or more per day of prednisone vi) cyclophosphamide, or vii) rituximab. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

For idiopathic pulmonary fibrosis and chronic fibrosing ILDs with a progressive phenotype: Prescribed by, or in consultation with, a pulmonologist. For systemic sclerosis-associated interstitial lung disease: Prescribed by, or in consultation with, a pulmonologist or rheumatologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

on with the property of the pr For idiopathic pulmonary fibrosis (all requests): Will not be used in combination with other agents for the prescribed indication.

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**OGSIVEO** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**OJEMDA** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

OJJAARA

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ONUREG** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**OPIPZA** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Both of the following: A) Member is unable to swallow aripiprazole tablet and B) Member is unable to use aripiprazole oral solution.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

# **COVERAGE DURATION**

Approved for duration of 1 year

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**OPSUMIT** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

Acheterization.

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERF

N/A

### PREREQUISITE THERAPY REQUIRED

**ORGOVYX** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ORKAMBI** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**ORSERDU** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# OTEZLA\_(UCARE)\_2026

# MEDICATION(S)

OTEZLA

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For oral ulcers associated with Behcet's disease (initial requests). Trial of topical triamcinolone 0.1% oral paste was ineffective or not tolerated. For psoriatic arthritis (all requests): Trial of other agents not required. For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For oral ulcers associated with Behcet's disease and psoriatic arthritis: Prescribed by, or in consultation with, a rheumatology specialist. For Plaque Psoriasis: Prescribed by, or in consultation with, a dermatologist (dermatologist not required for mild plaque psoriasis).

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For oral ulcers associated with Behcet's disease (initial requests): Diagnosis confirmed by the presence of oral ulcers and at least two of the following: recurrent genital ulceration, eye lesions, skin lesions, positive pathergy test. For psoriatic arthritis and plaque psoriasis (all requests): Will not be used in combination with biologic therapy for the prescribed indication.

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

YES

Pending CMS Approval

**PANRETIN** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

ACETYLCYSTEINE 10 % SOLUTION, ACETYLCYSTEINE 20 % SOLUTION, ACYCLOVIR SODIUM, ALBUTEROL SULFATE (2.5 MG/3ML) 0.083% NEBU SOLN, ALBUTEROL SULFATE (5 MG/ML) 0.5% NEBU SOLN, ALBUTEROL SULFATE 0.63 MG/3ML NEBU SOLN, ALBUTEROL SULFATE 1.25 MG/3ML NEBU SOLN, ALBUTEROL SULFATE 2.5 MG/0.5ML NEBU SOLN, AMPHOTERICIN B 50 MG RECON SOLN, AMPHOTERICIN B LIPOSOME, APREPITANT, ARFORMOTEROL TARTRATE, AZATHIOPRINE 50 MG TAB, BUDESONIDE 0.25 MG/2ML SUSPENSION, BUDESONIDE 0.5 MG/2ML SUSPENSION, BUDESONIDE 1 MG/2ML SUSPENSION, CROMOLYN SODIUM 20 MG/2ML NEBU SOLN, CYCLOPHOSPHAMIDE 25 MG TAB, CYCLOPHOSPHAMIDE 50 MG TAB, CYCLOPHOSPHAMIDE 25 MG CAP, CYCLOPHOSPHAMIDE 50 MG CAP, CYCLOSPORINE 100 MG CAP, CYCLOSPORINE 25 MG CAP, CYCLOSPORINE MODIFIED, DIPHTHERIA-TETANUS TOXOIDS DT, ENGERIX-B, ENVARSUS XR, EVEROLIMUS 0.25 MG TAB, EVEROLIMUS 0.5 MG TAB, EVEROLIMUS 0.75 MG TAB, EVEROLIMUS 1 MG TAB, FORMOTEROL FUMARATE 20 MCG/2ML NEBU SOLN, GRANISETRON HCL 1 MG TAB, HEPLISAV-B, HUMULIN R U-500 (CONCENTRATED), IMOVAX RABIES, INSULIN ASPART, IPRATROPIUM BROMIDE 0.02 % SOLUTION, IPRATROPIUM-ALBUTEROL, JYNNEOS, LEVALBUTEROL HCL 0.31 MG/3ML NEBU SOLN, LEVALBUTEROL HCL 0.63 MG/3ML NEBU SOLN, LEVALBUTEROL HCL 1.25 MG/0.5ML NEBU SOLN, LEVALBUTEROL HCL 1.25 MG/3ML NEBU SOLN, METHYLPREDNISOLONE 16 MG TAB, METHYLPREDNISOLONE 32 MG TAB, METHYLPREDNISOLONE 4 MG TAB, METHYLPREDNISOLONE 8 MG TAB, MYCOPHENOLATE MOFETIL 200 MG/ML RECON SUSP, MYCOPHENOLATE MOFETIL 250 MG CAP, MYCOPHENOLATE MOFETIL 500 MG TAB, MYCOPHENOLATE SODIUM, MYCOPHENOLIC ACID, NOVOLOG, NOVOLOG RELION, ONDANSETRON 4 MG TAB DISP, ONDANSETRON 8 MG TAB DISP, ONDANSETRON HCL 4 MG TAB, ONDANSETRON HCL 4 MG/5ML SOLUTION, ONDANSETRON HCL 8 MG TAB, PENTAMIDINE ISETHIONATE FOR NEBULIZATION SOLUTION, PLENAMINE, PREDNISOLONE 15 MG/5ML SOLUTION, PREDNISOLONE SODIUM PHOSPHATE 15 MG/5ML SOLUTION, PREDNISOLONE SODIUM PHOSPHATE 25 MG/5ML SOLUTION, PREDNISOLONE SODIUM PHOSPHATE 6.7 (5 BASE) MG/5ML SOLUTION, PREDNISONE 1 MG TAB, PREDNISONE 10 MG TAB, PREDNISONE 2.5 MG TAB, PREDNISONE 20 MG TAB, PREDNISONE 5 MG TAB, PREDNISONE 5 MG/5ML SOLUTION, PREDNISONE 50 MG TAB, PREDNISONE INTENSOL, PREHEVBRIO, PROGRAF 0.2 MG PACKET, PROGRAF 1 MG PACKET, PULMOZYME, RABAVERT, RECOMBIVAX HB, SIROLIMUS 0.5 MG TAB, SIROLIMUS 1 MG TAB, SIROLIMUS 1 MG/ML SOLUTION, SIROLIMUS 2 MG TAB, TACROLIMUS 0.5 MG CAP, TACROLIMUS 1 MG CAP, TACROLIMUS 5 MG CAP, TDVAX, TENIVAC, TETANUS-DIPHTHERIA TOXOIDS TD

### **DETAILS**

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

Pending CMS Approval

# PEGASYS\_(UCARE)\_2026

# **MEDICATION(S)**

**PEGASYS** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# DITHER CRITERIA A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**PEMAZYRE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# PENICILLAMINE\_(UCARE)\_2026

# **MEDICATION(S)**

PENICILLAMINE 250 MG TAB

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Diagnosis

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

For Wilson's Disease: Prescribed by or in consultation with a gastroenterologist, hepatologist or liver transplant physician

# **COVERAGE DURATION**

Approved for duration of 1 year

# **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

# PHENYLBUTYRATE\_(UCARE)\_2026

# **MEDICATION(S)**

SODIUM PHENYLBUTYRATE 500 MG TAB

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

Concomitant use of other phenylbutyrate products (e.g., Ravicti, Buphenyl, Pheburane, Olpruva)

### REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by or in consultation with a metabolic disease specialist (or specialist who focuses in the treatment of metabolic diseases)

# **COVERAGE DURATION**

Approved for duration of 1 year

### OTHER CRITERIA

For urea cycle disorders: Approve if genetic testing confirmed a mutation resulting in a urea cycle disorder or if the member has hyperammonemia.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

PIQRAY (200 MG DAILY DOSE), PIQRAY (250 MG DAILY DOSE), PIQRAY (300 MG DAILY DOSE)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

# COVERAGE DURATION Approved for duration of 1 year OTHER CRITERIA /A RT B PRERF

N/A

### PREREQUISITE THERAPY REQUIRED

**POMALYST** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

PREVYMIS 120 MG PACKET, PREVYMIS 240 MG TAB, PREVYMIS 480 MG TAB

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

Member will/has initiated Prevymis within 30 days after an allogeneic hematopoietic stem cell transplant or 7 days after kidney transplant.

## **AGE RESTRICTION**

N/A

## PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a hematologist, oncologist, transplant or infectious disease specialist.

## **COVERAGE DURATION**

Approved for 8 months for hematopoietic stem cell transplant or 8 months for kidney transplant.

## OTHER CRITERIA

N/A

## PART B PREREQUISITE

N/A

## PREREQUISITE THERAPY REQUIRED

# PROMACTA\_NVT\_2026

# **MEDICATION(S)**

**ELTROMBOPAG OLAMINE** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

## PREREQUISITE THERAPY REQUIRED

QINLOCK

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

QUININE SULFATE 324 MG CAP

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

ARPROVAL SIRPO REQUIRED MEDICAL INFORMATION

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for 1 month.

**OTHER CRITERIA** 

N/A

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

# RADICAVA\_NVT\_2026

# **MEDICATION(S)**

RADICAVA ORS, RADICAVA ORS STARTER KIT

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

For initial requests: Member has a score of two or greater for each individual item on the Amyotrophic Lateral Sclerosis Functional Rating Scale-Revised (ALSFRS-R). For continuation requests: Member has benefited with use of this medication.

## **AGE RESTRICTION**

N/A

## PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

# **COVERAGE DURATION**

Approved for duration of 1 year.

## OTHER CRITERIA

N/A

## PART B PREREQUISITE

N/A

## PREREQUISITE THERAPY REQUIRED

**RALDESY** 

## PA INDICATION INDICATOR

N/A

## **OFF LABEL USES**

REQUIRED MEDICAL INFORMATION

Member is unable to swallow solid dosage forms of trazodone.

AGE RESTRICTION

V/A

PRESCRIBEP

N/A

## **COVERAGE DURATION**

Approved for duration of 1 year.

## **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

## PREREQUISITE THERAPY REQUIRED

YES

# RETACRIT\_NVT\_2026

# **MEDICATION(S)**

**RETACRIT** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# DITHER CRITERIA A RT B PRERE

N/A

## PREREQUISITE THERAPY REQUIRED

RETEVMO 120 MG TAB, RETEVMO 160 MG TAB, RETEVMO 40 MG TAB, RETEVMO 80 MG TAB

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

# APPROVAL. REQUIRED MEDICAL INFORMATION

N/A

## **AGE RESTRICTION**

N/A

## PRESCRIBER RESTRICTION

N/A

## **COVERAGE DURATION**

Approved for duration of 1 year.

## **OTHER CRITERIA**

N/A

## PART B PREREQUISITE

N/A

## PREREQUISITE THERAPY REQUIRED

# **REVATIO\_NVT\_2026**

# **MEDICATION(S)**

SILDENAFIL CITRATE 20 MG TAB

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## **OFF LABEL USES**

JER RESTRICTION

COVERAGE DURATION
Approved for duration of 1 years

OTHER CRITERIA
/A

RT B PRERF

N/A

## PREREQUISITE THERAPY REQUIRED

## **REVCOVI NVT 2026**

## **MEDICATION(S)**

**REVCOVI** 

## PA INDICATION INDICATOR

N/A

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

For adenosine deaminase severe combined immune deficiency (ADA-SCID)(initial requests): Diagnosis confirmed by one of the following: a) Absent or very low (less than 1 percent of normal) ADA activity in red blood cells, b) Increased levels of deoxyadenosine triphosphate in erythrocyte lysates compared to laboratory standards, c) Significantly decreased concentration of adenosine triphosphate in red blood cells, d) Absent or very low (less than 5 percent of normal) levels of sadenosylhomocysteine hydrolase in red blood cells, e) Elevated levels of 2-deoxyadensoine in plasma, urine, or dried blood spots, or f) Presence of biallelic pathogenic mutations in the ADA gene. For ADA-SCID (continuation requests): Member has benefited with use of this medication.

## AGE RESTRICTION

Elevated levels of 2-deoxyadensoine in plasma, urine, or dried blood spots, or f) Presence of biallelic pathogenic mutations in the ADA gene. For ADA-SCID (continuation requests): Member has benefited with use of this medication.

# PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, an immunologist or provider who specializes ADA-SCID.

## COVERAGE DURATION

Approved for duration of 1 year.

## **OTHER CRITERIA**

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

N/A

Pending CMS Approval

**LENALIDOMIDE** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# **REVUFORJ\_NVT\_2026**

# **MEDICATION(S)**

**REVUFORJ** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# REZDIFFRA\_NVT\_2026

## **MEDICATION(S)**

**REZDIFFRA** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

For noncirrhotic nonalcoholic steatohepatitis (initial requests): 1) Stage F2 or F3 fibrosis confirmed by one of the following: a) Liver biopsy or b) Both of the following: i) Fibrosis-4 score greater than or equal to 1.3 and ii) One of the following: Vibration-controlled transient elastography greater than or equal to 8 kPa, magnetic resonance elastography greater than or equal to 3.63 kPa, or enhanced liver fibrosis test greater than or equal to 7.7 and 2) Attestation that the medication will be used in conjunction with diet and exercise. For noncirrhotic nonalcoholic steatohepatitis (continuation requests): Member has benefited with use of this medication.

## **AGE RESTRICTION**

N/A

# PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a hepatologist or gastroenterologist.

## **COVERAGE DURATION**

Approved for duration of 1 year.

## OTHER CRITERIA

N/A

## PART B PREREQUISITE

Pending CMS Approval

**REZLIDHIA** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# REZUROCK\_NVT\_2026

# **MEDICATION(S)**

REZUROCK

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# RINVOQ\_(UCARE)\_2026

## MEDICATION(S)

RINVOQ, RINVOQ LQ

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

For rheumatoid arthritis (initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For psoriatic arthritis (initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For atopic dermatitis (initial requests): Two of the following were ineffective or not tolerated: a) A medium to very high potency topical steroid, b) A topical calcineurin inhibitor or c) An oral immunosuppressant. For ulcerative colitis (initial requests): Trial of a TNF antagonist was ineffective or not tolerated. For ankylosing spondylitis (initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For non-radiographic axial spondyloarthritis (initial requests): Trial of two non-steroidal anti-inflammatory drugs (NSAIDs) was ineffective or not tolerated. For Crohn's disease (initial requests): Trial of Hadlima or Simlandi was ineffective or not tolerated. For polyarticular juvenile idiopathic arthritis (pJIA)(initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For giant cell arteritis (initial requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

## AGE RESTRICTION

N/A

## PRESCRIBER RESTRICTION

For rheumatoid arthritis, psoriatic arthritis, ankylosing spondylitis, pJIA, non-radiographic axial spondyloarthritis, or giant cell arteritis: Prescribed by, or in consultation with, a rheumatology specialist. For atopic dermatitis: Prescribed by, or in consultation with, an allergist, immunologist, or dermatologist. For Crohn's disease or ulcerative colitis: Prescribed by, or in consultation with a gastroenterologist.

## **COVERAGE DURATION**

Approved for duration of 1 year.

## **OTHER CRITERIA**

For atopic dermatitis (initial requests): Member has moderate to severe atopic dermatitis defined as: 1) One of the following: a) Body surface area involvement of 10 percent or more or b) Involvement of the face, head, neck, hands, feet, groin, or intertriginous areas and 2) At least two of the following: a) Intractable pruritus (itching), b) Cracking and oozing/bleeding of skin or c) Impaired activities of daily Pendino CMS Approval living. For atopic dermatitis (all requests): Will not be used in combination with another targeted immunomodulator product for the prescribed indication.

## PART B PREREQUISITE

N/A

## PREREQUISITE THERAPY REQUIRED

**ROMVIMZA** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# **ROZLYTREK NVT 2026**

# **MEDICATION(S)**

**ROZLYTREK** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

## PREREQUISITE THERAPY REQUIRED

# **RUBRACA NVT 2026**

# **MEDICATION(S)**

RUBRACA

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**RYDAPT** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# DITHER CRITERIA A RT B PRERE

N/A

## PREREQUISITE THERAPY REQUIRED

VIGABATRIN, VIGPODER

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

N/A

## **AGE RESTRICTION**

N/A

## PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

## **COVERAGE DURATION**

Approved for duration of 1 year.

## **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

## PREREQUISITE THERAPY REQUIRED

## **SAXENDA**

# MEDICATION(S)

LIRAGLUTIDE -WEIGHT MANAGEMENT, SAXENDA

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

## OFF LABEL USES

N/A

## **EXCLUSION CRITERIA**

Concurrent use of other weight loss medications (other than phentermine) and glucagon-like peptide-1 (GLP-1) medications.

## REQUIRED MEDICAL INFORMATION

N/A

## **AGE RESTRICTION**

N/A

## PRESCRIBER RESTRICTION

N/A

## **COVERAGE DURATION**

Initial: Approved for 6 months. Continuation: Approved for 1 year.

## OTHER CRITERIA

For weight loss (initial requests): 1) The member has a baseline body mass index (BMI) of 30 kg/m2 or greater or has a BMI of 27 kg/m2 or greater with at least one weight-related comorbidity AND 2) documentation of the following has been provided: a) baseline BMI and weight, and b) evidence the member is utilizing a comprehensive weight management plan that includes a reduced calorie diet or care of a registered dietitian, AND 3) evidence the member is utilizing a comprehensive weight management plan that includes initiation of or ongoing regimen of increased physical activity, unless medically contraindicated, AND 4) the member has been utilizing a comprehensive weight management plan for at least 6 months or greater. For weight loss (continuation) - 1) Documentation of the following has been provided: a) the member's baseline and current weight, and b) evidence the member will continue to utilize a comprehensive weight management plan including reduced calorie

diet or care of registered dietitian and increased physical activity, AND 2) (for adults) the member has maintained at least a 5% reduction in body weight from baseline OR (for pediatrics) the member has maintained at least five percent (5%) reduction in body mass index (BMI) from baseline.

## **PART B PREREQUISITE**

N/A

## PREREQUISITE THERAPY REQUIRED



**SCEMBLIX** 

## PA INDICATION INDICATOR

N/A

## **OFF LABEL USES**

REQUIRED MEDICAL INFORMATION
For T315I mutation: failure of or intolerance to Iclusig required.

AGE RESTRICTION

V/A

PRESCRIBEP PF

N/A

## **COVERAGE DURATION**

Approved for duration of 1 year.

## **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

## PREREQUISITE THERAPY REQUIRED

YES

**SECUADO** 

## PA INDICATION INDICATOR

N/A

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

Two of the following were ineffective or not tolerated: a) aripiprazole, b) olanzapine, c) quetiapine, d) risperidone, e) ziprasidone, f) lurasidone, or g) oral asenapine.

## **AGE RESTRICTION**

N/A

## PRESCRIBER RESTRICTION

N/A

# **COVERAGE DURATION**

Approved for duration of 1 year

## **OTHER CRITERIA**

N/A

## **PART B PREREQUISITE**

N/A

## PREREQUISITE THERAPY REQUIRED

YES

# SENSIPAR (UCARE) 2026

## **MEDICATION(S)**

CINACALCET HCL

## PA INDICATION INDICATOR

4 - All FDA-Approved Indications, Some Medically-Accepted Indications

## OFF LABEL USES

Hyperparathyroidism in post-renal transplant patients

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

N/A

## AGE RESTRICTION

N/A

## PRESCRIBER RESTRICTION

Hypercalcemia due to parathyroid carcinoma: Prescribed by, or in consultation with, an oncologist or endocrinologist. Hypercalcemia with primary hyperparathyroidism: Prescribed by, or in consultation with, a nephrologist or endocrinologist. Hyperparathyroidism in post-renal transplant: Prescribed by, or in consultation with, a transplant physician, nephrologist or endocrinologist.

## **COVERAGE DURATION**

Approved for duration of 1 year.

## OTHER CRITERIA

For hypercalcemia due to parathyroid carcinoma: Approve. For hypercalcemia in patients with primary hyperparathyroidism: Approve if the member has failed or is unable to undergo a parathyroidectomy due to a contraindication, as determined by the prescriber. For hyperparathyroidism in post-renal transplant patients: Approve if the baseline (prior to starting cinacalcet therapy) calcium and intact parathyroid hormone (iPTH) levels are above the normal range, as defined by the laboratory reference values. For secondary hyperparathyroidism in patients with chronic kidney disease on dialysis: Deny under Medicare Part D (claim should be submitted under the end stage renal disease (ESRD) bundle payment benefit).

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

N/A

Pendino CMS Approval

**SIGNIFOR** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

## PREREQUISITE THERAPY REQUIRED

# SIMLANDI (UCARE) 2026

## MEDICATION(S)

SIMLANDI (1 PEN), SIMLANDI (1 SYRINGE), SIMLANDI (2 PEN), SIMLANDI (2 SYRINGE)

## PA INDICATION INDICATOR

N/A

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

For rheumatoid arthritis (initial requests): Trial of methotrexate at a dose of at least 20mg/week (or maximally tolerated dose) was ineffective or not tolerated. For polyarticular juvenile idiopathic arthritis (pJIA)(initial requests): Trial of methotrexate at a dose of at least 15 mg/week (or maximally tolerated dose) was ineffective or not tolerated. For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of at least 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For ankylosing spondylitis (AS)(all requests): Trial of other agents not required. For psoriatic arthritis (all requests): Trial of other agents not required. For bidradenitis not required. For Crohn's disease (all requests): Trial of other agents not required. For hidradenitis suppurativa (initial requests): Member must have both of the following: a) At least 3 cysts and b) Trial of one oral antibiotic was ineffective or not tolerated. For uveitis (initial requests): Both of the following were ineffective or not tolerated: a) a corticosteroid and b) an immunosuppressant (methotrexate, mycophenolate mofetil, or cyclosporine). For continuation requests (all diagnoses): Member has benefited with use of this medication.

## AGE RESTRICTION

N/A

## PRESCRIBER RESTRICTION

For rheumatoid arthritis, psoriatic arthritis, pJIA or ankylosing spondylitis: Prescribed by, or in consultation with, a rheumatology specialist. For plaque psoriasis or hidradenitis suppurativa: Prescribed by, or in consultation with, a dermatologist. For Crohn's disease and ulcerative colitis: Prescribed by, or in consultation with, a gastroenterologist. For uveitis: Prescribed by, or in consult

with, a rheumatology specialist or ophthalmologist.

## **COVERAGE DURATION**

Approved for duration of 1 year.

**OTHER CRITERIA** 

N/A

**PART B PREREQUISITE** 

N/A

Pending CMS Approval PREREQUISITE THERAPY REQUIRED

YES

# SKYRIZI (UCARE) 2026

## MEDICATION(S)

SKYRIZI 150 MG/ML SOLN PRSYR, SKYRIZI 180 MG/1.2ML SOLN CART, SKYRIZI 360 MG/2.4ML SOLN CART, SKYRIZI PEN

## PA INDICATION INDICATOR

N/A

## **OFF LABEL USES**

N/A

## **EXCLUSION CRITERIA**

N/A

## REQUIRED MEDICAL INFORMATION

For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of at least 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For psoriatic arthritis (all requests): Trial of other agents not required. For Crohn's disease (all requests): Trial of other agents not required. For ulcerative colitis (all requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

## **AGE RESTRICTION**

N/A

# PRESCRIBER RESTRICTION

For psoriatic arthritis: Prescribed by, or in consultation with, a rheumatology specialist. For plaque psoriasis: Prescribed by, or in consultation with, a dermatologist. For Crohn's disease and ulcerative colitis: Prescribed by, or in consultation with, a gastroenterologist.

## **COVERAGE DURATION**

Approved for duration of 1 year.

## OTHER CRITERIA

N/A

## PART B PREREQUISITE

## PREREQUISITE THERAPY REQUIRED

YES

pending chis Approval

# **SOLARAZE NVT 2026**

# **MEDICATION(S)**

**DICLOFENAC SODIUM 3 % GEL** 

## PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**SOMAVERT** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**DASATINIB** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# STELARA\_(UCARE)\_2026

### MEDICATION(S)

STELARA 45 MG/0.5ML SOLN PRSYR, STELARA 45 MG/0.5ML SOLUTION, STELARA 90 MG/ML SOLN PRSYR, USTEKINUMAB 45 MG/0.5ML SOLN PRSYR, USTEKINUMAB 45 MG/0.5ML SOLUTION, USTEKINUMAB 90 MG/ML SOLN PRSYR

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# REQUIRED MEDICAL INFORMATION

For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of at least 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For psoriatic arthritis (all requests): Trial of other agents not required. For ulcerative colitis (all requests): Trial of other agents not required. For Crohn's disease (all requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

# PRESCRIBER BESTRICTION

For psoriatic arthritis: Prescribed by, or in consultation with, a rheumatology specialist. For Crohn's disease and ulcerative colitis: Prescribed by, or in consultation with, a gastroenterologist. For plaque psoriasis: Prescribed by, or in consultation with, a dermatologist.

### COVERAGE DURATION

Approved for duration of 1 year.

### **OTHER CRITERIA**

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

YES

Pending CMS Approval

# STEQEYMA\_(UCARE)\_2026

### **MEDICATION(S)**

STEQEYMA 45 MG/0.5ML SOLN PRSYR, STEQEYMA 90 MG/ML SOLN PRSYR

### PA INDICATION INDICATOR

N/A

### OFF LABEL USES

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For plaque psoriasis (initial requests): One of the following was ineffective or not tolerated: a) methotrexate at a dose of at least 10mg/week (or maximally tolerated dose) or b) acitretin (trial of other agents not required for patients under 18 years of age). For psoriatic arthritis (all requests): Trial of other agents not required. For ulcerative colitis (all requests): Trial of other agents not required. For Crohn's disease (all requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

For psoriatic arthritis: Prescribed by, or in consultation with, a rheumatology specialist. For Crohn's disease and ulcerative colitis: Prescribed by, or in consultation with, a gastroenterologist. For plaque psoriasis: Prescribed by, or in consultation with, a dermatologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

Pending CMS Approval

**STIVARGA** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

SUNOSI

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

One of the following was ineffective or not tolerated: a) modafinit or b) armodafinil.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

A nocturnal polysomnogram was used to confirm diagnosis.

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

SUNITINIB MALATE

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

# SYPRINE\_NVT\_2026

# **MEDICATION(S)**

TRIENTINE HCL 250 MG CAP

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

### **TABRECTA NVT 2026**

# **MEDICATION(S)**

**TABRECTA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**TAFINLAR** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**TAGRISSO** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**TALZENNA** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**ERLOTINIB HCL** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# **TARGRETIN\_NVT\_2026**

### **MEDICATION(S)**

**BEXAROTENE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

NILOTINIB HCL

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# TAZORAC\_NVT\_2026

# **MEDICATION(S)**

**TAZAROTENE 0.1 % CREAM** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

# PREREQUISITE THERAPY REQUIRED

**TAZVERIK** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**TEPMETKO** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# TERIPARATIDE\_(UCARE)\_2026

### **MEDICATION(S)**

**TERIPARATIDE** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

Concomitant use with other medications for osteoporosis

### REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

High risk for fracture: 2 years. Not high risk for fracture: Max of 2 years therapy per lifetime.

### OTHER CRITERIA

For postmenopausal osteoporosis: Approve if the member has tried one oral bisphosphonate (e.g., alendronate and ibandronate) or the member cannot take an oral bisphosphonate due to the inability to swallow or difficulty swallowing or the member cannot remain in an upright position following oral bisphosphonate administration or the member has a pre-existing gastrointestinal medical condition (e.g., esophageal lesions, esophageal ulcers, or abnormalities of the esophagus that delay esophageal emptying [stricture, achalasia]), or the member has tried an IV bisphosphonate (e.g., ibandronate or zoledronic acid), or the member has severe renal impairment (creatinine clearance less than 35 mL/min), has chronic kidney disease or the member has had an osteoporotic fracture or fragility fracture. For increasing bone mass in men (a man is defined as an individual with the biological traits of a man, regardless of the individual's gender identity or gender expression) with primary or hypogonadal osteoporosis or for the treatment of glucocorticoid induced osteoporosis: Approve if the member has

tried one oral bisphosphonate (e.g., alendronate and ibandronate) or the member cannot take an oral bisphosphonate due to the inability to swallow or difficulty swallowing or the member cannot remain in an upright position following oral bisphosphonate administration or the member has a pre-existing gastrointestinal medical condition (e.g., esophageal lesions, esophageal ulcers, or abnormalities of the esophagus that delay esophageal emptying [stricture, achalasia]), or the member has tried zoledronic acid (Reclast), or the member has severe renal impairment (CrCL less than 35 mL/min), has chronic kidney disease or the member has had an osteoporotic fracture or fragility fracture. Patients who have Pending CMS Approval already taken teriparatide (Forteo) for 2 years: Approve if the member is at high risk for fracture.

### PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

YES

# **TESTOSTERONE\_NVT\_2026**

### MEDICATION(S)

TESTOSTERONE 1.62 % GEL, TESTOSTERONE 12.5 MG/ACT (1%) GEL, TESTOSTERONE 20.25 MG/ACT (1.62%) GEL, TESTOSTERONE 25 MG/2.5GM (1%) GEL, TESTOSTERONE 30 MG/ACT SOLUTION, TESTOSTERONE 50 MG/5GM (1%) GEL, TESTOSTERONE CYPIONATE 100 MG/ML SOLUTION, TESTOSTERONE CYPIONATE 200 MG/ML SOLUTION, TESTOSTERONE ENANTHATE 200 MG/ML SOLUTION

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For initial requests (all diagnoses): A) Attestation that hypogonadism is not age-related and B) Documentation is provided of two morning fasting testosterone levels (from two separate days) that fall below the normal range for a healthy adult male. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

# PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE



**TIBSOVO** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

TOBRAMYCIN 300 MG/5ML NEBU SOLN

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

Approval will be based off BvD coverage determination.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

# **TOLVAPTAN\_NVT\_2026**

# **MEDICATION(S)**

**TOLVAPTAN** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Member has an eGFR of 25 ml/min/1.73m2 or greater (does not apply to generic Samsca equivalent).

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a nephrologist (does not apply to generic Samsca equivalent).

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

# TRACLEER\_NVT\_2026

# **MEDICATION(S)**

BOSENTAN 125 MG TAB, BOSENTAN 62.5 MG TAB

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

Acrestriction.

COVERAGE DURATION
Approved for duration of 1 years

OTHER CRITERIA
/A

RT B PRERF

N/A

### PREREQUISITE THERAPY REQUIRED

**TRIKAFTA** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**TRUQAP** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**TUKYSA** 

# PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

# TURALIO\_NVT\_2026

# **MEDICATION(S)**

**TURALIO 125 MG CAP** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

# **TYENNE (UCARE) 2026**

### **MEDICATION(S)**

TYENNE 162 MG/0.9ML SOLN A-INJ, TYENNE 162 MG/0.9ML SOLN PRSYR

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

Systemic sclerosis-associated interstitial lung disease

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For rheumatoid arthritis (initial requests): Two of the following were ineffective or not tolerated: a) Enbrel, b) Hadlima or Simlandi, c) Rinvoq or d) Xeljanz. For polyarticular juvenile idiopathic arthritis (pJIA)(initial requests): Two of the following were ineffective or not tolerated: a) Hadlima or Simlandi, b) Enbrel, c) Xeljanz, or d) Rinvoq. For giant cell arteritis (all requests): Trial of Rinvoq was ineffective or not tolerated. For systemic sclerosis-associated interstitial lung disease (initial requests): Both of the following: a) Diagnosis is confirmed with documentation provided of both of the following: i) HRCT scan and ii) pulmonary function tests and b) Trial of mycophenolate was ineffective or not tolerated. For systemic juvenile idiopathic arthritis (all requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

For rheumatoid arthritis, pJIA, systemic juvenile idiopathic arthritis, or giant cell arteritis: Prescribed by, or in consultation with, a rheumatology specialist. For systemic sclerosis-associated interstitial lung disease: Prescribed by, or in consultation with, a pulmonologist or rheumatologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

YES

Pending CMS Approval

LAPATINIB DITOSYLATE

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

# TYRVAYA\_(UCARE)\_2026

# **MEDICATION(S)**

**TYRVAYA** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For Dry Eye Disease: Trial of cyclosporine 0.05% eye emulsion was ineffective or not tolerated

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**BUDESONIDE ER 9 MG** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Trial of mesalamine was ineffective or not tolerated.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**VALCHLOR** 

### **PA INDICATION INDICATOR**

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**VANFLYTA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### **VENCLEXTA NVT 2026**

# **MEDICATION(S)**

VENCLEXTA, VENCLEXTA STARTING PACK

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**VERZENIO** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**VIGAFYDE** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Both of the following: a) Member is unable to swallow vigabatrin tablet and b) Member is unable to use vigabatrin powder for oral solution.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year

### **OTHER CRITERIA**

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**VITRAKVI** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**VIZIMPRO** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**VONJO** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### **REQUIRED MEDICAL INFORMATION**

Two of the following were ineffective or not tolerated: a) Jakafi, b) Inrebic, or c) Ojjaara.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

# VOQUEZNA\_(UCARE)\_2026

# **MEDICATION(S)**

**VOQUEZNA** 

### PA INDICATION INDICATOR

N/A

**OFF LABEL USES** 

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Trial of at least one generic proton pump inhibitor (PPI) medication was ineffective or not tolerated.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

**VORANIGO** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

VORICONAZOLE 200 MG TAB, VORICONAZOLE 50 MG TAB, VORICONAZOLE 200 MG RECON SOLN, VORICONAZOLE 40 MG/ML RECON SUSP

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

TION CANS ARRIVA REQUIRED MEDICAL INFORMATION

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for 6 months.

N/A

**PART B PREREQUISITE** 

N/A

PREREQUISITE THERAPY REQUIRED

**VOSEVI** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

All of the following: 1) Current HCV-RNA titer is provided, 2) Member does not have decompensated cirrhosis, and 3) Previous hepatitis C treatments are provided.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a gastroenterologist, hepatologist, infectious disease or transplant specialist.

### **COVERAGE DURATION**

Coverage duration of 12 weeks.

# OTHER CRITERIA

Treatment regimen will be approved based on previous treatment experience as defined by current AASLD guidelines.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

PAZOPANIB HCL

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**VOWST** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# " CINS ARPRIONAL REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for 1 month.

### **OTHER CRITERIA**

For all requests: Will not be used in combination with fecal microbiota, live for rectal use (Rebyota) or bezlotoxumab (Zinplava).

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

# WEGOVY\_MEDD\_(UCARE)\_2026

### **MEDICATION(S)**

**WEGOVY** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

Weight loss

# "Wo Wookong" REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

1 year

### **OTHER CRITERIA**

Reduction of major adverse cardiovascular events - Approve if the patient has established cardiovascular disease defined as a prior myocardial infarction, prior stroke, or peripheral arterial disease AND patient has a BMI of 27 or greater AND patient does not have type 1 or type 2 diabetes.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

# **WEIGHT LOSS DRUGS UCARE 2026**

### MEDICATION(S)

SAXENDA, WEGOVY

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

Concurrent use of other weight loss medications (other than phentermine) and glucagon-like peptide-1 (GLP-1) medications.

### REQUIRED MEDICAL INFORMATION

N/A

### AGE RESTRICTION

N/A

### PRESCRIBER RESTRICTION

N/A

### COVERAGE DURATION

Initial: Approved for 6 months. Continuation: Approved for 1 year.

### OTHER CRITERIA

For weight loss (initial requests): 1) The member has a baseline body mass index (BMI) of 30 kg/m2 or greater or has a BMI of 27 kg/m2 or greater with at least one weight-related comorbidity AND 2) documentation of the following has been provided: a) baseline BMI and weight, and b) evidence the member is utilizing a comprehensive weight management plan that includes a reduced calorie diet or care of a registered dietitian, AND 3) evidence the member is utilizing a comprehensive weight management plan that includes initiation of or ongoing regimen of increased physical activity, unless medically contraindicated, AND 4) the member has been utilizing a comprehensive weight management plan for at least 6 months or greater. For weight loss (continuation) - 1) Documentation of the following has been provided: a) the member's baseline and current weight, and b) evidence the member will continue to utilize a comprehensive weight management plan including reduced calorie diet or care of registered dietitian and increased physical activity, AND 2)

(for adults) the member has maintained at least a 5% reduction in body weight from baseline OR (for pediatrics) the member has maintained at least five percent (5%) reduction in body mass index (BMI) from baseline.

# **PART B PREREQUISITE**

**WELIREG** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**WINREVAIR** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

atheterization.

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERF

N/A

### PREREQUISITE THERAPY REQUIRED

**WYOST** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

DITHER CRITERIA

A

RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

XALKORI

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

### XCOPRI\_NVT\_2026

# **MEDICATION(S)**

XCOPRI, XCOPRI (250 MG DAILY DOSE), XCOPRI (350 MG DAILY DOSE)

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Two of the following were ineffective or not tolerated: a) lamotrigine b) carbamazepine c) levetiracetam d) oxcarbazepine e) phenytoin f) topiramate or g) lacosamide.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

### **COVERAGE DURATION**

Approved for duration of 1 year

# OTHER CRITERIA

N/A

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

YES

# XDEMVY\_(UCARE)\_2026

# **MEDICATION(S)**

**XDEMVY** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

For demodex blepharitis: Prescribed by or in consultation with an optometrist or ophthalmologist.

### **COVERAGE DURATION**

Approved for duration of 6 weeks.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

# XELJANZ\_(UCARE)\_2026

### **MEDICATION(S)**

XELJANZ, XELJANZ XR

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For rheumatoid arthritis (initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For polyarticular juvenile idiopathic arthritis (pJIA)(initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For psoriatic arthritis (initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For ankylosing spondylitis (initial requests): One of the following was ineffective or not tolerated: a) Hadlima or Simlandi or b) Enbrel. For ulcerative colitis (initial requests): Trial of a TNF antagonist was ineffective or not tolerated. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

# PRESCRIBER RESTRICTION

For rheumatoid arthritis, pJIA, ankylosing spondylitis, or psoriatic arthritis: Prescribed by, or in consultation with a rheumatology specialist. For ulcerative colitis: Prescribed by, or in consultation with a gastroenterologist.

### COVERAGE DURATION

Approved for duration of 1 year.

### **OTHER CRITERIA**

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

N/A

Pendino CMS Approval

**XERMELO** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

XIFAXAN 550 MG TAB

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

For diagnosis of IBS-D, approval will increase quantity limit to 42 tablets over 14 days, maximum of three fills per 1 year.

### **PART B PREREQUISITE**

N/A

### PREREQUISITE THERAPY REQUIRED

# XOLAIR\_(UCARE)\_2026

MEDICATION(S)

**XOLAIR** 

PA INDICATION INDICATOR

N/A

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

### REQUIRED MEDICAL INFORMATION

For asthma (initial requests): History within the last year of at least one asthma exacerbation requiring one of the following, despite regular use of inhaled corticosteroids plus an additional controller(s): a) Treatment with systemic corticosteroids, b) Emergency department visit or c) Hospitalization. For chronic spontaneous urticaria (initial requests): Both of the following: 1) One of the following: a) Patient remains symptomatic despite H1 antihistamine treatment or b) Has intolerance or contraindication to H1 antihistamine treatment and 2) Trial of Dupixent was ineffective or not tolerated. For nasal polyps (initial requests): Dupixent was ineffective or not tolerated. For IgE-mediated food allergy (initial requests): Trial of other agents not required. For continuation requests (all diagnoses): Member has benefited with use of this medication.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

For asthma: Prescribed by, or in consultation with an allergist, pulmonologist, or immunologist. For chronic spontaneous urticaria: Prescribed by, or in consultation with an allergist, dermatologist, or immunologist. For nasal polyps: Prescribed by, or in consultation with, an allergist, immunologist, or otolaryngologist. For IgE-mediated food allergy: Prescribed by, or in consultation with an allergist or immunologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

For asthma (initial requests): Documentation of diagnosis via skin test or RAST for specific allergy sensitivity. For nasal polyps (initial requests): All of the following: a) Diagnosis of chronic rhinosinusitis with nasal polyposis, lasting at least 12 weeks, b) Bilateral nasal polyposis confirmed with sinus CT scan, and c) Moderate to severe symptoms of nasal congestion, blockage, or obstruction (such as loss of smell, rhinorrhea, or facial pain). For IgE-mediated food allergy (initial requests): Both of the following: a) Diagnosis supported by one of the following: i) Positive skin prick test or ii) Positive serum IgE test and b) Diagnosis confirmed by one of the following: i) Positive oral food challenge or ii) History of anaphylaxis to the suspected food allergen. For asthma (all requests): Will not be used in combination with another targeted immunomodulator product for the prescribed indication. For IgEmediated food allergy (all requests): Will not be used in combination with peanut allergen powder (Palforzia).

### PART B PREREQUISITE

N/A

Pendino Chis PREREQUISITE THERAPY REQUIRED

YES

### XOSPATA\_NVT\_2026

# **MEDICATION(S)**

**XOSPATA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

XPOVIO (100 MG ONCE WEEKLY) 50 MG TAB THPK, XPOVIO (40 MG ONCE WEEKLY) 10 MG TAB THPK, XPOVIO (40 MG ONCE WEEKLY) 40 MG TAB THPK, XPOVIO (40 MG TWICE WEEKLY) 40 MG TAB THPK, XPOVIO (60 MG ONCE WEEKLY) 60 MG TAB THPK, XPOVIO (60 MG TWICE WEEKLY), XPOVIO (80 MG ONCE WEEKLY) 40 MG TAB THPK, XPOVIO (80 MG TWICE WEEKLY)

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

# CINIS REQUIRED MEDICAL INFORMATION

N/A

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**XTANDI** 

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

Homologous recombination repair gene-mutated metastatic castration-resistant prostate cancer in combination with Talzenna.

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For metastatic castration-resistant prostate cancer and metastatic castration-sensitive prostate cancer: Trial of abiraterone was ineffective or not tolerated. For nonmetastatic castration-resistant prostate cancer: Both of the following were ineffective or not tolerated: a) Nubeqa and b) Erleada. For non metastatic castration sensitive prostate cancer with biochemical recurrence at high risk for metastasis: Trial of other agents not required. For homologous recombination repair gene-mutated metastatic castration-resistant prostate cancer in combination with Talzenna: Trial of other agents not required.

### AGE RESTRICTION

N/A

# PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

### OTHER CRITERIA

N/A

### PART B PREREQUISITE

Pending CMS Approval

### XYREM\_NVT\_2026

### **MEDICATION(S)**

SODIUM OXYBATE

### PA INDICATION INDICATOR

N/A

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

For excessive daytime sleepiness with narcolepsy in adults. Both of the following were ineffective or not tolerated: a) Sunosi and b) either modafinil or armodafinil. Trial of other agents not required for patients aged 7 to 17 years. For cataplexy with narcolepsy: Trial of other agents not required.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

N/A

### **COVERAGE DURATION**

Approved for duration of 1 year.

# OTHER CRITERIA

For excessive daytime sleepiness with narcolepsy: A nocturnal polysomnogram was used to confirm diagnosis. For cataplexy with narcolepsy: One of the following was used to confirm diagnosis: a) nocturnal polysomnogram or b) low cerebrospinal fluid orexin-A concentration.

### PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

YES

ZEJULA 100 MG TAB, ZEJULA 200 MG TAB, ZEJULA 300 MG TAB

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

# COVERAGE DURATION Approved for duration of 1 year. OTHER CRITERIA /A RT B PRERE

N/A

### PREREQUISITE THERAPY REQUIRED

**ZELBORAF** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

ZOLINZA

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

# COVERAGE DURATION Approved for duration of 1 year OTHER CRITERIA /A RT B PREREF

N/A

### PREREQUISITE THERAPY REQUIRED

**ZTALMY** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

### **OFF LABEL USES**

N/A

### **EXCLUSION CRITERIA**

N/A

### REQUIRED MEDICAL INFORMATION

Documentation is provided of a CDKL5 gene mutation.

### **AGE RESTRICTION**

N/A

### PRESCRIBER RESTRICTION

Prescribed by, or in consultation with, a neurologist.

### **COVERAGE DURATION**

Approved for duration of 1 year.

### **OTHER CRITERIA**

N/A

# PART B PREREQUISITE

N/A

### PREREQUISITE THERAPY REQUIRED

**ZURZUVAE** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

**OFF LABEL USES** 

N/A

**EXCLUSION CRITERIA** 

N/A

ino chis REQUIRED MEDICAL INFORMATION

N/A

**AGE RESTRICTION** 

N/A

PRESCRIBER RESTRICTION

N/A

**COVERAGE DURATION** 

Approved for 1 month.

**OTHER CRITERIA** 

N/A

PART B PREREQUISITE

N/A

PREREQUISITE THERAPY REQUIRED

**ZYDELIG** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year.

OTHER CRITERIA
/A
RT B PRERE

N/A

PREREQUISITE THERAPY REQUIRED

**ZYKADIA** 

### PA INDICATION INDICATOR

1 - All FDA-Approved Indications

COVERAGE DURATION
Approved for duration of 1 year

OTHER CRITERIA
/A

RT B PREREF

N/A

PREREQUISITE THERAPY REQUIRED