

VNS Health EasyCare Plus (HMO D-SNP) and

VNS Health Total (HMO D-SNP)

Prior Authorization Requirements

Effective: 09/01/2024

Updated: 08/26/24

ABATACEPT IV

Products Affected

• ORENCIA (WITH MALTOSE)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST.
Coverage Duration	RA, PJIA, PSA: INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS. AGVHD: 12 MONTHS.
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PJIA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ IR, RINVOQ. PSA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OTEZLA. RENEWAL: RA, PJIA, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION.

PA Criteria	Criteria Details
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ABATACEPT SQ

Products Affected

- ORENCIA
- ORENCIA CLICKJECT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PJIA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ IR, RINVOQ. PSA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OTEZLA. RENEWAL: RA, PJIA, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION.

PA Criteria	Criteria Details
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ABEMACICLIB

Products Affected

• VERZENIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ABIRATERONE

Products Affected

• abiraterone

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC HIGH-RISK CASTRATION-SENSITIVE PROSTATE CANCER (MCSPC), METASTATIC CASTRATION- RESISTANT PROSTATE CANCER (MCRPC): 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ABIRATERONE SUBMICRONIZED

Products Affected

• YONSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC CASTRATION-RESISTANT PROSTATE CANCER: 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ACALABRUTINIB

Products Affected

• CALQUENCE (ACALABRUTINIB MAL)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	TRIAL OF OR CONTRAINDICATION TO ONE OF THE PREFERRED AGENTS: BRUKINSA OR IMBRUVICA, WHERE INDICATIONS ALIGN.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ADAGRASIB

Products Affected

• KRAZATI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ADALIMUMAB

Products Affected

- HUMIRA PEN
- HUMIRA PEN CROHNS-UC-HS START
- HUMIRA PEN PSOR-UVEITS-ADOL HS
- HUMIRA SUBCUTANEOUS SYRINGE KIT 40 MG/0.8 ML
- HUMIRA(CF)

- HUMIRA(CF) PEDI CROHNS STARTER
- HUMIRA(CF) PEN
- HUMIRA(CF) PEN CROHNS-UC-HS
- HUMIRA(CF) PEN PEDIATRIC UC
- HUMIRA(CF) PEN PSOR-UV-ADOL HS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, FACE, OR GENITAL AREA.
Age Restrictions	
Prescriber Restrictions	INITIAL: RA, POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA), ANKYLOSING SPONDYLITIS (AS): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST. UVEITIS: PRESCRIBED BY OR IN CONSULTATION WITH AN OPTHALMOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: TRIAL OF OR CONTRAINDICATION TO 3 MONTHS OF TREATMENT WITH ONE DMARD (DISEASE- MODIFYING ANTIRHEUMATIC DRUG) - IF PATIENT TRIED METHOTREXATE, THEN TRIAL AT A DOSE GREATER THAN OR EQUAL TO 20 MG PER WEEK OR MAXIMALLY TOLERATED DOSE IS REQUIRED. PJIA, PSA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD. PSO: TRIAL OF OR CONTRAINDICATION TO ONE DMARD. PSO: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. CD, UC: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. UVEITIS: NO ISOLATED ANTERIOR UVEITIS. RENEWAL: RA, PJIA, PSA, AS, PSO, HIDRADENITIS SUPPURATIVA, UVEITIS: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AFATINIB

Products Affected

• GILOTRIF

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS.
Other Criteria	METASTATIC NSCLC WITH EGFR MUTATION: NOT ON CONCURRENT THERAPY WITH AN EGFR TYROSINE KINASE-INHIBITOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AGALSIDASE BETA

Products Affected

• FABRAZYME

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	FABRY DISEASE: INITIAL: 1) SYMPTOMATIC OR EVIDENCE OF INJURY FROM GL-3 TO THE KIDNEY, HEART, OR CENTRAL NERVOUS SYSTEM RECOGNIZED BY LABORATORY, HISTOLOGICAL, OR IMAGING FINDINGS, AND 2) ONE OF THE FOLLOWING: (A) FEMALES: GALACTOSIDASE ALPHA (GLA) GENE MUTATION VIA GENETIC TESTING, OR (B) MALES: ENZYME ASSAY INDICATING ALPHA GALACTOSIDASE A DEFICIENCY OR GLA GENE MUTATION VIA GENETIC TESTING.
Age Restrictions	
Prescriber Restrictions	FABRY DISEASE: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH NEPHROLOGIST, CARDIOLOGIST, OR SPECIALIST IN GENETICS OR INHERITED METABOLIC DISORDERS.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	FABRY DISEASE: INITIAL: NO CONCURRENT USE WITH ANOTHER FABRY DISEASE THERAPY. RENEWAL: 1) DEMONSTRATED IMPROVEMENT OR STABILIZATION, AND 2) NO CONCURRENT USE WITH ANOTHER FABRY DISEASE THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

ALECTINIB

Products Affected

• ALECENSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ALPELISIB-PIQRAY

Products Affected

 PIQRAY ORAL TABLET 200 MG/DAY (200 MG X 1), 250 MG/DAY (200 MG X1-50 MG X1), 300 MG/DAY (150 MG X 2)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AMBRISENTAN

Products Affected

• ambrisentan

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	PAH: INITIAL: DOES NOT HAVE IDIOPATHIC PULMONARY FIBROSIS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AMIVANTAMAB-VMJW

Products Affected

• RYBREVANT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ANAKINRA

Products Affected

• KINERET

PA Criteria	Criteria Details
Exclusion Criteria	CORONAVIRUS DISEASE 2019 (COVID-19) IN HOSPITALIZED ADULTS.
Criteria	HOSPITALIZED ADULTS.
Required Medical Information	
Age Restrictions	
Prescriber	INITIAL: RHEUMATOID ARTHRITIS (RA): PRESCRIBED BY
Restrictions	OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage	RA: INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS. ALL
Duration	OTHERS: 12 MONTHS.
Other Criteria	RA: INITIAL: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, RINVOQ, ENBREL, XELJANZ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

APALUTAMIDE

Products Affected

• ERLEADA ORAL TABLET 240 MG, 60 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: NON-METASTATIC CASTRATION-RESISTANT PROSTATE CANCER (NMCRPC): HIGH RISK PROSTATE CANCER (I.E., RAPIDLY INCREASING PROSTATE SPECIFIC ANTIGEN [PSA] LEVELS). NMCRPC, METASTATIC CASTRATION-SENSITIVE PROSTATE CANCER (MCSPC): 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG. RENEWAL: NMCRPC, MCSPC: 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GNRH ANALOG.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

APOMORPHINE

Products Affected

• apomorphine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	PARKINSONS DISEASE (PD): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	PD: INITIAL: PHYSICIAN HAS OPTIMIZED DRUG THERAPY FOR PD. RENEWAL: IMPROVEMENT WITH MOTOR FLUCTUATIONS DURING OFF EPISODES WHILE ON THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

APOMORPHINE - SL

Products Affected

 KYNMOBI SUBLINGUAL FILM 10 MG, 10-15-20-25-30 MG, 15 MG, 20 MG, 25 MG, 30 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	PARKINSONS DISEASE (PD): INITIAL: 18 YEARS OF AGE OR OLDER.
Prescriber Restrictions	PD: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	PD: INITIAL: PHYSICIAN HAS OPTIMIZED DRUG THERAPY FOR PARKINSONS DISEASE. RENEWAL: IMPROVEMENT WITH MOTOR FLUCTUATIONS DURING OFF EPISODES WITH THE USE OF THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

APREMILAST

Products Affected

• OTEZLA ORAL TABLET 30 MG

(4)-30 MG(19)

 OTEZLA STARTER ORAL TABLETS, DOSE PACK 10 MG (4)-20 MG (4)-30 MG (47), 10 MG (4)-20 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: MILD PLAQUE PSORIASIS (PSO): ONE OF THE FOLLOWING: 1) PSORIASIS COVERING 2 PERCENT OF BODY SURFACE AREA (BSA), 2) STATIC PHYSICIAN GLOBAL ASSESSMENT (SPGA) SCORE OF 2, OR 3) PSORIASIS AREA AND SEVERITY INDEX (PASI) SCORE OF 2 TO 9. MODERATE TO SEVERE PSO: 1) PSORIASIS COVERING 3 PERCENT OR MORE OF BSA, OR 2) PSORIATIC LESIONS AFFECTING THE HANDS, FEET, FACE, OR GENITAL AREA.
Age Restrictions	
Prescriber Restrictions	INITIAL: PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. BEHCETS DISEASE: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
PA Criteria Other Criteria	Criteria Details INITIAL: PSA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OR 2) TRIAL OF OR CONTRAINDICATION TO ONE DMARD. MILD PSO: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL SYSTEMIC THERAPY (E.G., METHOTREXATE, ACITRETIN, CYCLOSPORINE) OR ONE CONVENTIONAL TOPICAL THERAPY (E.G., PUVA, UVB, TOPICAL CORTICOSTEROIDS). MODERATE TO SEVERE PSO: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: COSENTYX, ENBREL, HUMIRA, STELARA, SKYRIZI, TREMFYA, OR 2) TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. BEHCETS DISEASE: 1) HAS ORAL ULCERS OR A HISTORY OF RECURRENT ORAL ULCERS BASED ON
	CLINICAL SYMPTOMS, AND 2) TRIAL OF OR CONTRAINDICATION TO ONE OR MORE CONSERVATIVE TREATMENTS (E.G., COLCHICINE, TOPICAL CORTICOSTEROID, ORAL CORTICOSTEROID). RENEWAL: PSA, PSO, BEHCETS DISEASE: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ASCIMINIB

Products Affected

• SCEMBLIX ORAL TABLET 100 MG, 20 MG, 40 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PHILADELPHIA CHROMOSOME-POSITIVE CHRONIC MYELOID LEUKEMIA (Ph+ CML): MUTATIONAL ANALYSIS PRIOR TO INITIATION AND SCEMBLIX IS APPROPRIATE PER NCCN GUIDELINE TABLE FOR TREATMENT RECOMMENDATIONS BASED ON BCR-ABL1 MUTATION PROFILE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ASFOTASE ALFA

Products Affected

• STRENSIQ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	HYPOPHOSPHATASIA (HPP): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST, GENETICIST, OR METABOLIC SPECIALIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: PERINATAL/INFANTILE-ONSET HPP: 1) 6
	MONTHS OF AGE OR YOUNGER AT ONSET OF HPP, AND 2)
	POSITIVE FOR A TISSUE NON-SPECIFIC ALKALINE
	PHOSPHATASE (TNSALP) (ALPL) GENE MUTATION AS
	CONFIRMED BY GENETIC TESTING OR TWO OF THE
	FOLLOWING: (A) SERUM ALKALINE PHOSPHATASE (ALP)
	LEVEL BELOW THAT OF NORMAL RANGE FOR PATIENT
	AGE, (B) ELEVATED SERUM PYRIDOXAL-5'-PHOSPHATE
	(PLP) LEVELS AND NO VITAMIN B6 SUPPLEMENTATION IN
	THE PREVIOUS WEEK, (C) URINE
	PHOSPHOETHANOLAMINE (PEA) LEVEL ABOVE THAT OF
	NORMAL RANGE FOR PATIENT AGE, (D) RADIOGRAPHIC
	EVIDENCE OF HPP, (E) AT LEAST TWO OF THE
	FOLLOWING: (I) RACHITIC CHEST DEFORMITY, (II)
	CRANIOSYNOSTOSIS, (III) DELAY IN SKELETAL GROWTH
	RESULTING IN DELAY OF MOTOR DEVELOPMENT, (IV)
	HISTORY OF VITAMIN B6 DEPENDENT SEIZURES, (V)
	NEPHROCALCINOSIS OR HISTORY OF ELEVATED SERUM
	CALCIUM, (VI) HISTORY OR PRESENCE OF NON-
	TRAUMATIC POSTNATAL FRACTURE AND DELAYED
	FRACTURE HEALING. JUVENILE-ONSET HPP: 1) 18 YEARS
	OF AGE OR YOUNGER AT ONSET OF HPP, AND 2) POSITIVE
	FOR A TNSALP ALPL GENE MUTATION AS CONFIRMED
	BY GENETIC TESTING OR TWO OF THE FOLLOWING: (A)
	SERUM ALP LEVEL BELOW THAT OF NORMAL RANGE
	FOR PATIENT AGE, (B) ELEVATED SERUM PLP LEVELS
	AND NO VITAMIN B6 SUPPLEMENTATION IN THE
	PREVIOUS WEEK, (C) URINE PEA LEVEL ABOVE THAT OF
	NORMAL RANGE FOR PATIENT AGE, (D) RADIOGRAPHIC
	EVIDENCE OF HPP, (E) AT LEAST TWO OF THE
	FOLLOWING: (I) RACHITIC DEFORMITIES, (II)
	PREMATURE LOSS OF PRIMARY TEETH PRIOR TO 5
	YEARS OF AGE, (III) DELAY IN SKELETAL GROWTH
	RESULTING IN DELAY OF MOTOR DEVELOPMENT, (IV)

PA Criteria	Criteria Details
	HISTORY OR PRESENCE OF NON-TRAUMATIC FRACTURES OR DELAYED FRACTURE HEALING. ALL INDICATIONS: 1) NOT CURRENTLY RECEIVING TREATMENT WITH A BISPHOSPHONATE, 2) CALCIUM OR PHOSPHATE LEVELS ARE NOT BELOW THE NORMAL RANGE, 3) NOT HAVE A TREATABLE FORM OF RICKETS. RENEWAL: ALL INDICATIONS: 1) IMPROVEMENT IN THE SKELETAL CHARACTERISTICS OF HPP, AND 2) NOT CURRENTLY RECEIVING TREATMENT WITH A BISPHOSPHONATE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ATEZOLIZUMAB

Products Affected

• TECENTRIQ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ATOGEPANT

Products Affected

• QULIPTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	MIGRAINE PREVENTION: INITIAL: 1) TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING PREVENTIVE MIGRAINE TREATMENTS: DIVALPROEX SODIUM, TOPIRAMATE, PROPRANOLOL, TIMOLOL, AND 2) NO CONCURRENT USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION. RENEWAL: 1) REDUCTION IN MIGRAINE OR HEADACHE FREQUENCY, MIGRAINE SEVERITY, OR MIGRAINE DURATION WITH THERAPY, AND 2) NO CONCURRENT USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AVAPRITINIB

Products Affected

• AYVAKIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AVATROMBOPAG

Products Affected

- DOPTELET (10 TAB PACK)
- DOPTELET (15 TAB PACK)
- DOPTELET (30 TAB PACK)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: CHRONIC LIVER DISEASE (CLD): PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST, GASTROENTEROLOGIST, HEPATOLOGIST, IMMUNOLOGIST, ENDOCRINOLOGIST, OR A SURGEON. CHRONIC IMMUNE THROMBOCYTOPENIA (ITP): PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR IMMUNOLOGIST.
Coverage Duration	CLD: 1 MONTH. CHRONIC ITP: INITIAL: 2 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: CLD: 1) PLANNED PROCEDURE 10 TO 13 DAYS AFTER INITIATION OF DOPTELET, AND 2) NOT RECEIVING OTHER THROMBOPOIETIN RECEPTOR AGONISTS (E.G., ROMIPLOSTIM, ELTROMBOPAG, ETC.). CHRONIC ITP: TRIAL OF OR CONTRAINDICATION TO CORTICOSTEROIDS OR IMMUNOGLOBULINS OR INSUFFICIENT RESPONSE TO SPLENECTOMY. RENEWAL: CHRONIC ITP: PATIENT HAD A CLINICAL RESPONSE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AXITINIB

Products Affected

• INLYTA ORAL TABLET 1 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AZACITIDINE

Products Affected

• ONUREG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

AZTREONAM INHALED

Products Affected

• CAYSTON

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	7 YEARS OF AGE OR OLDER
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BARICITINIB

Products Affected

• OLUMIANT

PA Criteria	Criteria Details
Exclusion	CORONAVIRUS DISEASE 2019 (COVID-19) IN
Criteria	HOSPITALIZED ADULTS.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. SEVERE ALOPECIA AREATA: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST.
Coverage Duration	RA, SEVERE ALOPECIA AREATA: INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: RA: TRIAL OF OR CONTRAINDICATION TO ANY TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ. SEVERE ALOPECIA AREATA: 1) AT LEAST 50 PERCENT SCALP HAIR LOSS AS MEASURED BY THE SEVERITY OF ALOPECIA TOOL (SALT) FOR MORE THAN 6 MONTHS, AND 2) NO CONCURRENT USE WITH OTHER SYSTEMIC BIOLOGICS FOR SEVERE ALOPECIA AREATA OR OTHER JAK INHIBITORS FOR ANY INDICATION. RENEWAL: RA: CONTINUES TO BENEFIT FROM THE MEDICATION. SEVERE ALOPECIA AREATA: 1) CONTINUES TO BENEFIT FROM THE MEDICATION, AND 2) NO CONCURRENT USE WITH OTHER SYSTEMIC BIOLOGICS FOR SEVERE ALOPECIA AREATA OR OTHER JAK INHIBITORS FOR ANY INDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

BECAPLERMIN

Products Affected

• REGRANEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	DIABETIC NEUROPATHIC ULCERS: PRESCRIBED BY OR IN CONSULTATION WITH A VASCULAR SURGEON, PODIATRIST, ENDOCRINOLOGIST, PHYSICIAN PRACTICING IN A SPECIALTY WOUND CLINIC OR INFECTIOUS DISEASE SPECIALIST.
Coverage Duration	3 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BEDAQUILINE

Products Affected

• SIRTURO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	24 WEEKS
Other Criteria	PULMONARY MULTI-DRUG RESISTANT TUBERCULOSIS (MDR-TB): SIRTURO USED IN COMBINATION WITH AT LEAST 3 OTHER ANTIBIOTICS FOR THE TREATMENT OF PULMONARY MDR-TB.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BELIMUMAB

- BENLYSTA INTRAVENOUS
- BENLYSTA SUBCUTANEOUS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: SYSTEMIC LUPUS ERYTHEMATOSUS (SLE): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. LUPUS NEPHRITIS (LN): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR NEPHROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	INITIAL: SLE: CURRENTLY TAKING CORTICOSTEROIDS, ANTIMALARIALS, NSAIDS, OR IMMUNOSUPPRESSIVE AGENTS. RENEWAL: SLE: PATIENT HAD CLINICAL IMPROVEMENT. LN: IMPROVEMENT IN RENAL RESPONSE FROM BASELINE LABORATORY VALUES (I.E., EGFR OR PROTEINURIA) AND/OR CLINICAL PARAMETERS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BELUMOSUDIL

Products Affected

• REZUROCK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BELZUTIFAN

Products Affected

• WELIREG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BENDAMUSTINE

- bendamustine intravenous recon soln
- BENDAMUSTINE INTRAVENOUS
- SOLUTION
- BENDEKA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BENRALIZUMAB

- FASENRA
- FASENRA PEN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	ASTHMA: INITIAL: BLOOD EOSINOPHIL LEVEL GREATER THAN OR EQUAL TO 150 CELLS/MCL WITHIN THE PAST 12 MONTHS.
Age Restrictions	
Prescriber Restrictions	ASTHMA: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN ALLERGY OR PULMONARY MEDICINE.
Coverage Duration	INITIAL: 4 MONTHS, RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	ASTHMA: INITIAL: 1) CONCURRENT THERAPY WITH A
	MEDIUM, HIGH-DOSE, OR MAXIMALLY TOLERATED
	DOSE OF AN INHALED CORTICOSTEROID (ICS) AND ONE
	OTHER MAINTENANCE MEDICATION, AND 2) ONE
	ASTHMA EXACERBATION REQUIRING SYSTEMIC
	CORTICOSTEROID BURST LASTING 3 OR MORE DAYS
	WITHIN THE PAST 12 MONTHS, OR AT LEAST ONE
	SERIOUS EXACERBATION REQUIRING HOSPITALIZATION
	OR ER VISIT WITHIN THE PAST 12 MONTHS, OR POOR
	SYMPTOM CONTROL DESPITE CURRENT THERAPY AS
	EVIDENCED BY AT LEAST THREE OF THE FOLLOWING
	WITHIN THE PAST 4 WEEKS: DAYTIME ASTHMA
	SYMPTOMS MORE THAN TWICE/WEEK, ANY NIGHT
	WAKING DUE TO ASTHMA, SABA RELIEVER FOR
	SYMPTOMS MORE THAN TWICE/WEEK, ANY ACTIVITY
	LIMITATION DUE TO ASTHMA, AND 3) NO CONCURRENT
	USE WITH XOLAIR, DUPIXENT OR OTHER ANTI-IL5
	BIOLOGICS WHEN USED FOR ASTHMA. RENEWAL: 1) NO
	CONCURRENT USE WITH XOLAIR, DUPIXENT OR OTHER
	ANTI-IL5 BIOLOGICS WHEN USED FOR ASTHMA, 2)
	CONTINUED USE OF ICS AND ONE OTHER MAINTENANCE
	MEDICATION, AND 3) CLINICAL RESPONSE AS
	EVIDENCED BY: (A) REDUCTION IN ASTHMA
	EXACERBATIONS FROM BASELINE, (B) DECREASED
	UTILIZATION OF RESCUE MEDICATIONS, (C) INCREASE
	IN PERCENT PREDICTED FEV1 FROM PRETREATMENT
	BASELINE, OR (D) REDUCTION IN SEVERITY OR
	FREQUENCY OF ASTHMA-RELATED SYMPTOMS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B	No
Prerequisite	

BETAINE

Products Affected

• betaine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BEVACIZUMAB-ADCD

Products Affected

• VEGZELMA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BEVACIZUMAB-AWWB

Products Affected

• MVASI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BEVACIZUMAB-BVZR

Products Affected

• ZIRABEV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BEXAROTENE

Products Affected

• bexarotene

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BINIMETINIB

Products Affected

• MEKTOVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BORTEZOMIB

- bortezomib injection VELCADE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BOSENTAN

- bosentan
- TRACLEER ORAL TABLET FOR SUSPENSION

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	PAH: INITIAL: 1) DOES NOT HAVE ELEVATED LIVER ENZYMES (ALT, AST) MORE THAN 3 TIMES UPPER LIMIT OF NORMAL (ULN) OR INCREASE IN BILIRUBIN BY 2 OR MORE TIMES ULN, AND 2) NOT CONCURRENTLY TAKING CYCLOSPORINE A OR GLYBURIDE. RENEWAL: NOT CONCURRENTLY TAKING CYCLOSPORINE A OR GLYBURIDE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BOSUTINIB

- BOSULIF ORAL CAPSULE 100 MG, 50 MG
- BOSULIF ORAL TABLET 100 MG, 400 MG, 500 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PREVIOUSLY TREATED (Ph+ CML): MUTATIONAL ANALYSIS PRIOR TO INITIATION AND BOSULIF IS APPROPRIATE PER NCCN GUIDELINE TABLE FOR TREATMENT RECOMMENDATIONS BASED ON BCR-ABL1 MUTATION PROFILE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

BRIGATINIB

- ALUNBRIG ORAL TABLET 180 MG, 30 MG, 90 MG
- ALUNBRIG ORAL TABLETS, DOSE PACK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

C1 ESTERASE INHIBITOR-CINRYZE

Products Affected

• CINRYZE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HEREDITARY ANGIOEDEMA (HAE): INITIAL: DIAGNOSIS CONFIRMED BY COMPLEMENT TESTING.
Age Restrictions	
Prescriber Restrictions	HAE: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST, IMMUNOLOGIST, OR ALLERGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	HAE: INITIAL: NOT ON CONCURRENT TREATMENT WITH ALTERNATIVE PROPHYLACTIC AGENT FOR HAE ATTACKS. RENEWAL: 1) IMPROVEMENT IN HAE ATTACKS (I.E., REDUCTIONS IN ATTACK FREQUENCY OR ATTACK SEVERITY), AND 2) NOT ON CONCURRENT TREATMENT WITH ALTERNATIVE PROPHYLACTIC AGENT FOR HAE ATTACKS. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

C1 ESTERASE INHIBITOR-HAEGARDA

Products Affected

• HAEGARDA SUBCUTANEOUS RECON SOLN 2,000 UNIT, 3,000 UNIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HEREDITARY ANGIOEDEMA (HAE): INITIAL: DIAGNOSIS CONFIRMED BY COMPLEMENT TESTING.
Age Restrictions	
Prescriber Restrictions	HAE: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST, IMMUNOLOGIST, OR ALLERGIST.
Coverage Duration	INITIAL AND RENEWAL: 12 MONTHS.
Other Criteria	HAE: INITIAL: NOT ON CONCURRENT TREATMENT WITH ALTERNATIVE PROPHYLACTIC AGENT FOR HAE ATTACKS. RENEWAL: 1) IMPROVEMENT COMPARED TO BASELINE IN HAE ATTACKS (I.E., REDUCTIONS IN ATTACK FREQUENCY OR ATTACK SEVERITY), AND 2) NOT ON CONCURRENT TREATMENT WITH ALTERNATIVE PROPHYLACTIC AGENT FOR HAE ATTACKS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CABOZANTINIB CAPSULE

Products Affected

 COMETRIQ ORAL CAPSULE 100 MG/DAY(80 MG X1-20 MG X1), 140 MG/DAY(80 MG X1-20 MG X3), 60 MG/DAY (20 MG X 3/DAY)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CABOZANTINIB TABLET

Products Affected

• CABOMETYX ORAL TABLET 20 MG, 40 MG, 60 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CANAKINUMAB

Products Affected

• ILARIS (PF)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: CRYOPYRIN-ASSOCIATED PERIODIC SYNDROMES (CAPS): 1) ONE OF THE FOLLOWING: (A) GENETIC TEST FOR GAIN-OF-FUNCTION MUTATIONS IN THE NLRP3 GENE, OR (B) HAS INFLAMMATORY MARKERS (I.E., ELEVATED CRP, ESR, SERUM AMYLOID A PROTEIN (SAA) OR S100 PROTEINS), AND 2) TWO OF THE FOLLOWING: URTICARIAL-LIKE RASH (NEUTROPHILIC DERMATITIS), COLD-TRIGGERED EPISODES, SENSORINEURAL HEARING LOSS, MUSCULOSKELETAL SYMPTOMS, CHRONIC ASEPTIC MENINGITIS, SKELETAL ABNORMALITIES.
Age Restrictions	
Prescriber Restrictions	INITIAL: SYSTEMIC JUVENILE IDIOPATHIC ARTHRITIS (SJIA), ADULT-ONSET STILLS DISEASE (AOSD): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST, DERMATOLOGIST, OR IMMUNOLOGIST. GOUT: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. RENEWAL: GOUT: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage Duration	INITIAL: AOSD/SJIA: 6 MO, CAPS: LIFETIME, ALL OTHER DIAGNOSES: 12 MO. RENEWAL: AOSD/SJIA/GOUT: 12 MO

PA Criteria	Criteria Details
Other Criteria	INITIAL: CAPS: NO CONCURRENT USE WITH OTHER IL-1 INHIBITORS. AOSD, SJIA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE- MODIFYING ANTIRHEUMATIC DRUG). GOUT: NO CONCURRENT USE WITH OTHER IL-1 INHIBITORS. RENEWAL: AOSD, SJIA: CONTINUES TO BENEFIT FROM THE MEDICATION. GOUT: 1) NO CONCURRENT USE WITH OTHER IL-1 INHIBITORS, AND 2) IMPROVEMENT IN GOUT FLARES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CANNABIDIOL

Products Affected

• EPIDIOLEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	DRAVET SYNDROME (DS), LENNOX-GASTAUT SYNDROME (LGS), TUBEROUS SCLEROSIS COMPLEX (TSC): PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	INITIAL: 12 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: LENNOX-GASTAUT SYNDROME (LGS): TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING ANTIEPILEPTIC MEDICATIONS: RUFINAMIDE, FELBAMATE, CLOBAZAM, TOPIRAMATE, LAMOTRIGINE, CLONAZEPAM. RENEWAL: DS, LGS, TSC: CONFIRMATION OF DIAGNOSIS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CAPIVASERTIB

Products Affected

• TRUQAP

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CAPLACIZUMAB YHDP

Products Affected

• CABLIVI INJECTION KIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	ACQUIRED THROMBOTIC THROMBOCYTOPENIA PURPURA (ATTP): PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST
Coverage Duration	12 MONTHS
Other Criteria	ATTP: CABLIVI WAS PREVIOUSLY INITIATED AS PART OF THE FDA APPROVED TREATMENT REGIMEN IN COMBINATION WITH PLASMA EXCHANGE AND IMMUNOSUPPRESSIVE THERAPY WITHIN AN INPATIENT SETTING. THE PATIENT HAS NOT EXPERIENCED MORE THAN TWO RECURRENCES OF ATTP WHILE ON CABLIVI THERAPY (I.E., NEW DROP IN PLATELET COUNT REQUIRING REPEAT PLASMA EXCHANGE DURING 30 DAYS POST-PLASMA EXCHANGE THERAPY [PEX] AND UP TO 28 DAYS OF EXTENDED THERAPY).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CAPMATINIB

Products Affected

• TABRECTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CARGLUMIC ACID

Products Affected

• carglumic acid

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: ACUTE OR CHRONIC HYPERAMMONEMIA (HA) DUE TO N ACETYLGLUTAMATE SYNTHASE (NAGS) DEFICIENCY: NAGS GENE MUTATION IS CONFIRMED BY BIOCHEMICAL OR GENETIC TESTING. ACUTE HA DUE TO PROPIONIC ACIDEMIA (PA): 1) CONFIRMED BY ELEVATED METHYLCITRIC ACID AND NORMAL METHYLMALONIC ACID, OR 2) GENETIC TESTING CONFIRMS MUTATION IN THE PCCA OR PCCB GENE. ACUTE HA DUE TO METHYLMALONIC ACIDEMIA (MMA): 1) CONFIRMED BY ELEVATED METHYLMALONIC ACID, METHYLCITRIC ACID, OR 2) GENETIC TESTING CONFIRMS MUTATION IN THE METHYLMALONIC ACID, METHYLCITRIC ACID, OR 2) GENETIC TESTING CONFIRMS MUTATION IN THE MMUT, MMA, MMAB OR MMADHC GENES.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	ACUTE HA DUE TO NAGS/PA/MMA: 7 DAYS. CHRONIC HA DUE TO NAGS: INITIAL: 6 MOS, RENEWAL: 12 MOS.
Other Criteria	RENEWAL: CHRONIC HA DUE TO NAGS: PATIENT HAS SHOWN CLINICAL IMPROVEMENT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CERITINIB

Products Affected

• ZYKADIA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CETUXIMAB

Products Affected

• ERBITUX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CLADRIBINE

- MAVENCLAD (10 TABLET PACK)
- MAVENCLAD (4 TABLET PACK)
- MAVENCLAD (5 TABLET PACK)
- MAVENCLAD (6 TABLET PACK)
- MAVENCLAD (7 TABLET PACK)
- MAVENCLAD (8 TABLET PACK)
- MAVENCLAD (9 TABLET PACK)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL: 48 WEEKS.
Other Criteria	RELAPSING FORM OF MULTIPLE SCLEROSIS (MS): INITIAL: HAS NOT RECEIVED A TOTAL OF TWO YEARS OF MAVENCLAD TREATMENT (I.E., TWO YEARLY TREATMENT COURSES OF TWO CYCLES IN EACH). RENEWAL: 1) HAS DEMONSTRATED CLINICAL BENEFIT COMPARED TO PRE-TREATMENT BASELINE, 2) DOES NOT HAVE LYMPHOPENIA, AND 3) HAS NOT RECEIVED A TOTAL OF TWO YEARS OF MAVENCLAD TREATMENT (I.E., TWO YEARLY TREATMENT COURSES OF TWO CYCLES IN EACH).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CLOBAZAM-SYMPAZAN

Products Affected

• SYMPAZAN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	LENNOX-GASTAUT SYNDROME (LGS): THERAPY IS PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	LGS: 1) UNABLE TO TAKE TABLETS OR SUSPENSIONS, AND 2) TRIAL OF OR CONTRAINDICATION TO A FORMULARY VERSION OF CLOBAZAM.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

COBIMETINIB

Products Affected

• COTELLIC

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CORTICOTROPIN

Products Affected

• ACTHAR

• CORTROPHIN GEL

 ACTHAR SELFJECT SUBCUTANEOUS PEN INJECTOR 40 UNIT/0.5 ML, 80 UNIT/ML

PA Criteria	Criteria Details
Exclusion Criteria	INITIAL: NOT APPROVED FOR DIAGNOSTIC PURPOSES.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: ALL FDA APPROVED INDICATIONS EXCEPT INFANTILE SPASMS AND MULTIPLE SCLEROSIS (MS): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST, DERMATOLOGIST, ALLERGIST/IMMUNOLOGIST, OPHTHALMOLOGIST, PULMONOLOGIST OR NEPHROLOGIST.
Coverage Duration	INFANTILE SPASMS AND MS: 28 DAYS. OTHER FDA APPROVED INDICATIONS: INITIAL AND RENEWAL: 28 DAYS
Other Criteria	INITIAL: ALL FDA APPROVED INDICATIONS EXCEPT INFANTILE SPASMS: TRIAL OF OR CONTRAINDICATION TO INTRAVENOUS (IV) CORTICOSTEROIDS. ALL FDA APPROVED INDICATIONS EXCEPT INFANTILE SPASMS AND MS: TRIAL OF OR CONTRAINDICATION TO A STANDARD OF CARE THERAPY. RENEWAL: ALL FDA APPROVED INDICATIONS EXCEPT INFANTILE SPASMS AND MS: 1) DEMONSTRATED CLINICAL BENEFIT WHILE ON THERAPY AS INDICATED BY SYMPTOM RESOLUTION AND/OR NORMALIZATION OF LABORATORY TESTS, AND 2) CONTINUES TO POSSESS CONTRAINDICATION TO IV CORTICOSTEROIDS. PART B BEFORE PART D STEP THERAPY, APPLIES ONLY TO BENEFICIARIES IN AN MA- PD PLAN.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

CRIZANLIZUMAB-TMCA

Products Affected

• ADAKVEO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	SICKLE CELL DISEASE (SCD): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST
Coverage Duration	INITIAL: 12 MONTHS. RENEWAL: LIFETIME
Other Criteria	SCD: INITIAL: AGES 18 YEARS OR OLDER: 1) AT LEAST 2 SICKLE CELL CRISES IN THE PAST YEAR, 2) SICKLE-CELL ASSOCIATED SYMPTOMS WHICH ARE INTERFERING WITH ACTIVITIES OF DAILY LIVING, OR 3) HISTORY OF OR HAS RECURRENT ACUTE CHEST SYNDROME. AGES 16 TO 17 YEARS: APPROVED WITHOUT ADDITIONAL CRITERIA. RENEWAL: MAINTAINED OR EXPERIENCED A REDUCTION IN ACUTE COMPLICATIONS OF SCD. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CRIZOTINIB

Products Affected

• XALKORI ORAL CAPSULE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CRIZOTINIB PELLETS

Products Affected

• XALKORI ORAL PELLET 150 MG, 20 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	NON-SMALL CELL LUNG CANCER (NSCLC), ANAPLASTIC LARGE CELL LYMPHOMA (ALCL), INFLAMMATORY MYOFIBROBLASTIC TUMOR (IMT): UNABLE TO SWALLOW CAPSULES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

CYSTEAMINE HYDROCHLORIDE

Products Affected

- CYSTADROPS
- CYSTARAN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DABRAFENIB

Products Affected

• TAFINLAR ORAL CAPSULE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DABRAFENIB SUSPENSION

Products Affected

 TAFINLAR ORAL TABLET FOR SUSPENSION

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	UNABLE TO SWALLOW TAFINILAR CAPSULES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DACOMITINIB

Products Affected

• VIZIMPRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC NON-SMALL CELL LUNG CANCER (NSCLC): NOT ON CONCURRENT THERAPY WITH AN EGFR TYROSINE KINASE-INHIBITOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DALFAMPRIDINE

Products Affected

• dalfampridine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	MULTIPLE SCLEROSIS (MS): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	INITIAL: 3 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	MS: INITIAL: HAS SYMPTOMS OF A WALKING DISABILITY SUCH AS MILD TO MODERATE BILATERAL LOWER EXTREMITY WEAKNESS OR UNILATERAL WEAKNESS PLUS LOWER EXTREMITY OR TRUNCAL ATAXIA. RENEWAL: IMPROVEMENT IN WALKING ABILITY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DARATUMUMAB

Products Affected

• DARZALEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DARATUMUMAB-HYALURONIDASE-FIHJ

Products Affected

• DARZALEX FASPRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DAROLUTAMIDE

Products Affected

• NUBEQA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	INITIAL: NON-METASTATIC CASTRATION-RESISTANT PROSTATE CANCER (NMCRPC): HIGH RISK PROSTATE CANCER (I.E., RAPIDLY INCREASING PROSTATE SPECIFIC ANTIGEN [PSA] LEVELS). NMCRPC, METASTATIC HORMONE-SENSITIVE PROSTATE CANCER (MHSPC): 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG. RENEWAL: NMCRPC, MHSPC: 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GNRH ANALOG.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DASATINIB

Products Affected

• SPRYCEL ORAL TABLET 100 MG, 140 MG, 20 MG, 50 MG, 70 MG, 80 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PREVIOUSLY-TREATED PHILADELPHIA CHROMOSOME- POSITIVE CHRONIC MYELOID LEUKEMIA (Ph+ CML): MUTATIONAL ANALYSIS PRIOR TO INITIATION AND SPRYCEL IS APPROPRIATE PER NCCN GUIDELINE TABLE FOR TREATMENT RECOMMENDATIONS BASED ON BCR- ABL1 MUTATION PROFILE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DECITABINE/CEDAZURIDINE

Products Affected

• INQOVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DEFERASIROX

Products Affected

• deferasirox

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS: SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 1000 MCG/L (AT LEAST TWO LAB VALUES IN THE PREVIOUS THREE MONTHS). NON-TRANSFUSION DEPENDENT THALASSEMIA (NTDT): 1) SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 300 MCG/L (AT LEAST TWO LAB VALUES IN THE PREVIOUS THREE MONTHS) AND 2) LIVER IRON CONCENTRATION (LIC) OF 5 MG FE/G DRY WEIGHT OR GREATER. RENEWAL: CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS: SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 500 MCG/L (AT LEAST TWO LAB VALUES IN THE PREVIOUS THREE MONTHS). NTDT: 1) SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 300 MCG/L (AT LEAST TWO LAB VALUES IN THE PREVIOUS THREE MONTHS) OR 2) LIC OF 3 MG FE/G DRY WEIGHT OR GREATER.
Age Restrictions	
Prescriber Restrictions	INITIAL: CHRONIC IRON OVERLOAD: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR HEMATOLOGIST/ONCOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	INITIAL FOR ALL INDICATIONS: FORMULARY VERSION OF DEFERASIROX SPRINKLE: TRIAL OF OR CONTRAINDICATION TO GENERIC DEFERASIROX TABLET OR TABLET FOR ORAL SUSPENSION.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

DEFERIPRONE

Products Affected

- deferiprone
- FERRIPROX (2 TIMES A DAY)
- FERRIPROX ORAL SOLUTION
- FERRIPROX ORAL TABLET 1,000

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	TRANSFUSIONAL IRON OVERLOAD: RENEWAL: SERUM FERRITIN LEVELS CONSISTENTLY ABOVE 500 MCG/L (AT LEAST TWO LAB VALUES IN THE PREVIOUS THREE MONTHS).
Age Restrictions	
Prescriber Restrictions	TRANSFUSIONAL IRON OVERLOAD: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR HEMATOLOGIST/ONCOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	INITIAL: TRANSFUSIONAL IRON OVERLOAD DUE TO THALASSEMIA SYNDROMES: 1) TRIAL OF, CONTRAINDICATION, INTOLERABLE TOXICITIES, OR CLINICALLY SIGNIFICANT ADVERSE EFFECTS TO A FORMULARY VERSION OF DEFERASIROX OR DEFEROXAMINE, OR 2) CURRENT CHELATION THERAPY (I.E., FORMULARY VERSION OF DEFERASIROX OR DEFEROXAMINE) IS INADEQUATE. TRANSFUSIONAL IRON OVERLOAD DUE TO SICKLE CELL DISEASE OR OTHER ANEMIAS: TRIAL OF OR CONTRAINDICATION TO A FORMULARY VERSION OF DEFERASIROX OR DEFEROXAMINE.
Indications	All FDA-approved Indications.
Off Label Uses	

MG

PA Criteria	Criteria Details
Part B Prerequisite	No

DEFEROXAMINE

Products Affected

• deferoxamine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CHRONIC IRON OVERLOAD: INITIAL: SERUM FERRITIN LEVELS CONSISTENTLY ABOVE 1000 MCG/L (AT LEAST TWO LAB VALUES IN THE PREVIOUS THREE MONTHS). RENEWAL: SERUM FERRITIN LEVELS CONSISTENTLY ABOVE 500 MCG/L (AT LEAST TWO LAB VALUES IN THE PREVIOUS THREE MONTHS).
Age Restrictions	CHRONIC IRON OVERLOAD: INITIAL: 3 YEARS OR OLDER
Prescriber Restrictions	CHRONIC IRON OVERLOAD: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR HEMATOLOGIST/ONCOLOGIST
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D. THIS DRUG MAY BE EITHER BUNDLED WITH AND COVERED UNDER END STAGE RENAL DISEASE DIALYSIS RELATED SERVICES OR COVERED UNDER MEDICARE D DEPENDING UPON THE CIRCUMSTANCES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DENOSUMAB-XGEVA

Products Affected

• XGEVA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DEUTETRABENAZINE

Products Affected

- AUSTEDO ORAL TABLET 12 MG, 6 MG, 9 MG
- AUSTEDO XR ORAL TABLET EXTENDED RELEASE 24 HR 12 MG,

18 MG, 24 MG, 30 MG, 36 MG, 42 MG, 48 MG, 6 MG

 AUSTEDO XR TITRATION KT(WK1-4)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	HUNTINGTON DISEASE: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST OR MOVEMENT DISORDER SPECIALIST. TARDIVE DYSKINESIA: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST, PSYCHIATRIST, OR MOVEMENT DISORDER SPECIALIST.
Coverage Duration	12 MONTHS
Other Criteria	TARDIVE DYSKINESIA: HISTORY OF USING AGENTS THAT CAUSE TARDIVE DYSKINESIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DICLOFENAC TOPICAL GEL

Products Affected

• diclofenac sodium topical gel 3 %

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DICLOFENAC TOPICAL SOLUTION

Products Affected

• *diclofenac sodium topical solution in metered-dose pump*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 MONTHS
Other Criteria	OSTEOARTHRITIS OF THE KNEE: TRIAL OF OR CONTRAINDICATION TO A FORMULARY VERSION OF DICLOFENAC SODIUM 1% TOPICAL GEL AND A FORMULARY VERSION OF DICLOFENAC SODIUM 1.5% TOPICAL DROPS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DIMETHYL FUMARATE

Products Affected

 dimethyl fumarate oral capsule, delayed release(drlec) 120 mg, 120 mg (14)-240 mg (46), 240 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DIROXIMEL FUMARATE

Products Affected

• VUMERITY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DOSTARLIMAB-GXLY

Products Affected

• JEMPERLI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DRONABINOL CAPSULE

Products Affected

• dronabinol

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 MONTHS
Other Criteria	NAUSEA AND VOMITING ASSOCIATED WITH CANCER CHEMOTHERAPY: TRIAL OF OR CONTRAINDICATION TO ONE ANTIEMETIC THERAPY. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D FOR THE INDICATION OF NAUSEA AND VOMITING ASSOCIATED WITH CANCER CHEMOTHERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DROXIDOPA

Products Affected

• droxidopa

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	NEUROGENIC ORTHOSTATIC HYPOTENSION (NOH): INITIAL: 1) BASELINE BLOOD PRESSURE READINGS WHILE THE PATIENT IS SITTING AND ALSO WITHIN 3 MINUTES OF STANDING FROM A SUPINE POSITION. 2) A DECREASE OF AT LEAST 20 MMHG IN SYSTOLIC BLOOD PRESSURE OR 10 MMHG DIASTOLIC BLOOD PRESSURE WITHIN THREE MINUTES AFTER STANDING FROM A SITTING POSITION.
Age Restrictions	
Prescriber Restrictions	NOH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST OR CARDIOLOGIST.
Coverage Duration	INITIAL: 3 MONTHS RENEWAL: 12 MONTHS
Other Criteria	NOH: RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DUPILUMAB

Products Affected

- DUPIXENT PEN
- DUPIXENT SYRINGE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: EOSINOPHILIC ASTHMA: BLOOD EOSINOPHIL LEVEL OF 150 TO 1500 CELLS/MCL WITHIN THE PAST 12 MONTHS. EOSINOPHILIC ESOPHAGITIS (EOE): DIAGNOSIS CONFIRMED BY ESOPHAGOGASTRODUODENOSCOPY (EGD) WITH BIOPSY.
Age Restrictions	
Prescriber Restrictions	INITIAL: AD, PN: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST, ALLERGIST OR IMMUNOLOGIST. ASTHMA: PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN ALLERGY OR PULMONARY MEDICINE. CRSWNP: PRESCRIBED BY OR IN CONSULTATION WITH AN OTOLARYNGOLOGIST, ALLERGIST OR IMMUNOLOGIST. EOE: PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST, ALLERGIST, OR IMMUNOLOGIST.
Coverage Duration	INITIAL: AD, CRSWNP, EOE, PN: 6 MOS, ASTHMA: 4 MOS. RENEWAL: ALL INDICATIONS: 12 MOS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: AD: 1) AD COVERING AT LEAST 10 PERCENT OF
	BODY SURFACE AREA OR AD AFFECTING THE FACE,
	HEAD, NECK, HANDS, FEET, GROIN, OR INTERTRIGINOUS
	AREAS, 2) INTRACTABLE PRURITUS OR
	CRACKING/OOZING/BLEEDING OF AFFECTED SKIN, 3)
	TRIAL OF OR CONTRAINDICATION TO ONE TOPICAL
	(CORTICOSTEROID, CALCINEURIN INHIBITOR, PDE4
	INHIBITOR, OR JAK INHIBITOR), AND 4) NO
	CONCURRENT USE WITH OTHER SYSTEMIC BIOLOGICS
	OR JAK INHIBITORS FOR AD. ASTHMA: 1) CONCURRENT
	THERAPY WITH A MEDIUM, HIGH-DOSE OR MAXIMALLY-
	TOLERATED DOSE OF AN INHALED CORTICOSTEROID
	(ICS) AND ONE OTHER MAINTENANCE MEDICATION, 2)
	ONE ASTHMA EXACERBATION REQUIRING SYSTEMIC
	CORTICOSTEROID BURST LASTING 3 OR MORE DAYS
	WITHIN THE PAST 12 MONTHS, OR ONE SERIOUS
	EXACERBATION REQUIRING HOSPITALIZATION OR ER
	VISIT WITHIN THE PAST 12 MONTHS, OR POOR SYMPTOM
	CONTROL DESPITE CURRENT THERAPY AS EVIDENCED
	BY AT LEAST THREE OF THE FOLLOWING WITHIN THE
	PAST 4 WEEKS: DAYTIME ASTHMA SYMPTOMS MORE
	THAN TWICE/WEEK, ANY NIGHT WAKING DUE TO
	ASTHMA, SABA RELIEVER FOR SYMPTOMS MORE THAN
	TWICE/WEEK, ANY ACTIVITY LIMITATION DUE TO
	ASTHMA, AND 3) NO CONCURRENT USE WITH XOLAIR OR
	OTHER ANTI-IL5 BIOLOGICS WHEN USED FOR ASTHMA.
	CRSWNP: 1) EVIDENCE OF NASAL POLYPS BY DIRECT
	EXAMINATION, ENDOSCOPY OR SINUS CT SCAN, 2)
	INADEQUATELY CONTROLLED DISEASE AS
	DETERMINED BY USE OF SYSTEMIC STEROIDS IN THE
	PAST 2 YEARS OR ENDOSCOPIC SINUS SURGERY, AND 3) A
	56 DAY TRIAL OF ONE TOPICAL NASAL
	CORTICOSTEROID. PN: 1) CHRONIC PRURITIS (ITCH MORE
	THAN 6 WEEKS), MULTIPLE PRURIGINOUS LESIONS, AND

PA Criteria	Criteria Details
	HISTORY OR SIGN OF A PROLONGED SCRATCHING BEHAVIOR, 2) TRIAL OF OR CONTRAINDICATION TO ONE TOPICAL (CORTICOSTEROID OR CALCIPOTRIOL). RENEWAL: AD: 1) IMPROVEMENT WHILE ON THERAPY, AND 2) NO CONCURRENT USE WITH OTHER SYSTEMIC BIOLOGICS OR JAK INHIBITORS FOR AD. CRSWNP, EOE: IMPROVEMENT WHILE ON THERAPY. ASTHMA: 1) NO CONCURRENT USE WITH XOLAIR, OR OTHER ANTI-IL5 BIOLOGICS FOR ASTHMA, 2) CONTINUED USE OF ICS AND ONE OTHER MAINTENANCE MEDICATION, AND 3) CLINICAL RESPONSE AS EVIDENCED BY: (A) REDUCTION IN ASTHMA EXACERBATIONS FROM BASELINE, (B) DECREASED UTILIZATION OF RESCUE MEDICATIONS, (C) INCREASE IN PERCENT PREDICTED FEV1 FROM PRETREATMENT BASELINE, OR (D) REDUCTION IN SEVERITY OR FREQUENCY OF ASTHMA-RELATED SYMPTOMS. PN: IMPROVEMENT OR REDUCTION OF PRURITIS OR PRURIGINOUS LESIONS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

DUVELISIB

Products Affected

• COPIKTRA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

EDARAVONE

Products Affected

- edaravone intravenous solution 30 mg/100 ml
- RADICAVA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	AMYOTROPHIC LATERAL SCLEROSIS (ALS): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST OR ALS SPECIALIST AT AN ALS SPECIALTY CENTER OR CARE CLINIC.
Coverage Duration	ALS: INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	ALS: INITIAL: 1) DURATION OF DISEASE (FROM ONSET OF SYMPTOMS) IS LESS THAN OR EQUAL TO 2 YEARS, 2) NORMAL RESPIRATORY FUNCTION, 3) HAS MILD TO MODERATE ALS WITH A SCORE OF 2 OR HIGHER IN ALL OF THE FOLLOWING 12 ITEMS OF THE AMYOTROPHIC LATERAL SCLEROSIS FUNCTIONAL RATING SCALE REVISED (ALSFRS-R): SPEECH, SALIVATION, SWALLOWING, HANDWRITING, CUTTING FOOD, DRESSING AND HYGIENE, TURNING IN BED, WALKING, CLIMBING STAIRS, DYSPNEA, ORTHOPNEA, RESPIRATORY INSUFFICIENCY, AND 4) TRIAL OF RILUZOLE TABLET OR CURRENTLY TAKING RILUZOLE TABLET. RENEWAL: 1) DOES NOT REQUIRE INVASIVE VENTILATION, AND 2) HAS IMPROVED BASELINE FUNCTIONAL ABILITY OR HAS MAINTAINED A SCORE OF 2 OR HIGHER IN ALL 12 ITEMS OF THE ALSFRS-R.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

EFLAPEGRASTIM-XNST

Products Affected

• ROLVEDON

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	NON MYELOID MALIGNANCY: TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENT: NYVEPRIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

EFLORNITHINE

Products Affected

• IWILFIN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	24 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELACESTRANT

Products Affected

 ORSERDU ORAL TABLET 345 MG, 86 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELAFIBRANOR

Products Affected

• IQIRVO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PRIMARY BILIARY CHOLANGITIS (PBC): INITIAL: DIAGNOSIS CONFIRMED BY TWO OF THE FOLLOWING: 1) ELEVATED ALKALINE PHOSPHATASE LEVEL, 2) PRESENCE OF ANTIMITOCHONDRIAL ANTIBODIES OR OTHER PBC-SPECIFIC AUTOANTIBODIES, INCLUDING SP100 OR GP210, IF AMA IS NEGATIVE, OR 3) HISTOLOGIC EVIDENCE (OBTAINED BY LIVER BIOPSY) OF NON- SUPPURATIVE DESTRUCTIVE CHOLANGITIS AND DESTRUCTION OF INTERLOBULAR BILE DUCTS.
Age Restrictions	
Prescriber Restrictions	PBC: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST OR HEPATOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	PBC: INITIAL: 1) NO CONCURRENT USE WITH ANOTHER SECOND-LINE THERAPY FOR PBC, 2) USED IN COMBINATION WITH URSODIOL IF INADEQUATE RESPONSE AFTER TREATMENT WITH URSODIOL MONOTHERAPY FOR AT LEAST 1 YEAR, OR USED AS MONOTHERAPY IF UNABLE TO TOLERATE URSODIOL, AND 3) DOES NOT HAVE DECOMPENSATED CIRRHOSIS (CHILD-PUGH B OR C), A PRIOR DECOMPENSATION EVENT, OR COMPENSATED CIRRHOSIS WITH EVIDENCE OF PORTAL HYPERTENSION. RENEWAL: 1) CONTINUES TO BENEFIT FROM THE MEDICATION, AND 2) NO CONCURRENT USE WITH ANOTHER SECOND-LINE THERAPY FOR PBC.

PA Criteria	Criteria Details
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELAGOLIX

Products Affected

• ORILISSA ORAL TABLET 150 MG, 200 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	MODERATE TO SEVERE PAIN ASSOCIATED WITH ENDOMETRIOSIS: INITIAL: DIAGNOSIS IS CONFIRMED VIA SURGICAL OR DIRECT VISUALIZATION (E.G., PELVIC ULTRASOUND) OR HISTOPATHOLOGICAL CONFIRMATION (E.G., LAPAROSCOPY OR LAPAROTOMY) IN THE LAST 10 YEARS.
Age Restrictions	MODERATE TO SEVERE PAIN ASSOCIATED WITH ENDOMETRIOSIS: INITIAL: 18 YEARS OF AGE OR OLDER.
Prescriber Restrictions	MODERATE TO SEVERE PAIN ASSOCIATED WITH ENDOMETRIOSIS: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN OBSTETRICIAN/GYNECOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS
Other Criteria	MODERATE TO SEVERE PAIN ASSOCIATED WITH ENDOMETRIOSIS: INITIAL: 1) NO CONCURRENT USE WITH ANOTHER GNRH-MODULATING AGENT, AND 2) TRIAL OF OR CONTRAINDICATION TO AN NSAID AND A PROGESTIN-CONTAINING PREPARATION. RENEWAL: 1) IMPROVEMENT IN PAIN ASSOCIATED WITH ENDOMETRIOSIS WHILE ON THERAPY, AND 2) NO CONCURRENT USE WITH ANOTHER GNRH-MODULATING AGENT
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

ELAPEGADEMASE-LVLR

Products Affected

• REVCOVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	ADENOSINE DEAMINASE SEVERE COMBINED IMMUNE DEFICIENCY (ADA-SCID): INITIAL: ADA-SCID AS MANIFESTED BY: 1) CONFIRMATORY GENETIC TEST, OR 2) SUGGESTIVE LABORATORY FINDINGS (E.G., ELEVATED DEOXYADENOSINE NUCLEOTIDE [DAXP] LEVELS, LYMPHOPENIA) AND HALLMARK SIGNS/SYMPTOMS (E.G., RECURRENT INFECTIONS, FAILURE TO THRIVE, PERSISTENT DIARRHEA).
Age Restrictions	
Prescriber Restrictions	ADA-SCID: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH IMMUNOLOGIST, HEMATOLOGIST/ONCOLOGIST, OR PHYSICIAN SPECIALIZING IN INHERITED METABOLIC DISORDERS.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	ADA-SCID: RENEWAL: 1) IMPROVEMENT OR MAINTENANCE OF IMMUNE FUNCTION FROM BASELINE, AND 2) HAS NOT RECEIVED SUCCESSFUL HCT OR GENE THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELEXACAFTOR-TEZACAFTOR-IVACAFTOR

- TRIKAFTA ORAL GRANULES IN PACKET, SEQUENTIAL
- TRIKAFTA ORAL TABLETS, SEQUENTIAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CYSTIC FIBROSIS (CF): INITIAL: CONFIRMED MUTATION IN CFTR GENE ACCEPTABLE FOR THE TREATMENT OF CYSTIC FIBROSIS.
Age Restrictions	
Prescriber Restrictions	CF: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST OR CYSTIC FIBROSIS EXPERT.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: LIFETIME.
Other Criteria	CF: RENEWAL: 1) MAINTAINED, IMPROVED, OR DEMONSTRATED LESS THAN EXPECTED DECLINE IN FEV1 OR BODY MASS INDEX (BMI), OR 2) REDUCTION IN NUMBER OF PULMONARY EXACERBATIONS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELIGLUSTAT

Products Affected

• CERDELGA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELOSULFASE ALFA

Products Affected

• VIMIZIM

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELRANATAMAB-BCMM

- ELREXFIO 44 MG/1.1 ML VIAL OUTER, SUV, P/F
- ELREXFIO SUBCUTANEOUS SOLUTION 40 MG/ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	RELAPSED OR REFRACTORY MULTIPLE MYELOMA: RENEWAL: 1) HAS RECEIVED AT LEAST 24 WEEKS OF TREATMENT WITH ELREXFIO, AND 2) HAS RESPONDED TO TREATMENT (PARTIAL RESPONSE OR BETTER), AND HAS MAINTAINED THIS RESPONSE FOR AT LEAST 2 MONTHS. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELTROMBOPAG

- PROMACTA ORAL POWDER IN PACKET 12.5 MG, 25 MG
- PROMACTA ORAL TABLET 12.5 MG, 25 MG, 50 MG, 75 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: ITP: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR IMMUNOLOGIST.
Coverage Duration	ITP: INITIAL: 2 MO, RENEWAL: 12 MO. HEPATITIS C, SEVERE APLASTIC ANEMIA: 12 MO.
Other Criteria	INITIAL: PERSISTENT OR CHRONIC IMMUNE THROMBOCYTOPENIA PURPURA (ITP): TRIAL OF OR CONTRAINDICATION TO CORTICOSTEROIDS, IMMUNOGLOBULINS, OR AN INSUFFICIENT RESPONSE TO SPLENECTOMY. ALL INDICATIONS: APPROVAL FOR PROMACTA ORAL SUSPENSION PACKETS REQUIRES A TRIAL OF PROMACTA TABLETS OR PATIENT IS UNABLE TO TAKE TABLET FORMULATION. RENEWAL: ITP: PATIENT HAS SHOWN A CLINICAL RESPONSE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ELTROMBOPAG - ALVAIZ

Products Affected

• ALVAIZ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PERSISTENT OR CHRONIC IMMUNE THROMBOCYTOPENIA (ITP): INITIAL: 1) PLATELET COUNT IS LESS THAN 30 X 10^9/L FROM AT LEAST 2 SEPARATE LABS IN THE LAST 3 MONTHS, OR 2) PLATELET COUNT IS LESS THAN 50 X 10^9/L FROM AT LEAST 2 SEPARATE LABS IN THE LAST 3 MONTHS AND HAD A PRIOR BLEEDING EVENT.
Age Restrictions	
Prescriber Restrictions	INITIAL: ITP: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR IMMUNOLOGIST.
Coverage Duration	ITP: INITIAL: 2 MO, RENEWAL: 12 MO. HEPATITIS C, SEVERE APLASTIC ANEMIA: 12 MO.
Other Criteria	INITIAL: ITP: 1) TRIAL OF OR CONTRAINDICATION TO CORTICOSTEROIDS OR IMMUNOGLOBULINS, OR AN INSUFFICIENT RESPONSE TO SPLENECTOMY, AND 2) NO CONCURRENT USE WITH OTHER THROMBOPOIETIN RECEPTOR AGONISTS (TPO-RAS) OR SPLEEN TYROSINE KINASE (SYK) INHIBITOR. RENEWAL: ITP: 1) IMPROVEMENT IN PLATELET COUNT FROM BASELINE OR REDUCTION IN BLEEDING EVENTS, AND 2) NO CONCURRENT USE WITH OTHER TPO-RAS OR SYK INHIBITOR.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

EMAPALUMAB-LZSG

Products Affected

• GAMIFANT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HEMOPHAGOCYTIC LYMPHOHISTIOCYTOSIS (HLH): INITIAL: 1) A GENETIC TEST IDENTIFYING HLH- ASSOCIATED GENE MUTATION (E.G., PRF1, UNC13D), OR 2) HAS AT LEAST FIVE OF THE FOLLOWING EIGHT DIAGNOSTIC CRITERIA FOR HLH: (A) FEVER, (B) SPLENOMEGALY, (C) CYTOPENIAS (AFFECTING AT LEAST 2 OF 3 CELL LINEAGES), (D) HYPERTRIGLYCERIDEMIA OR HYPOFIBRINOGENEMIA, (E) HEMOPHAGOCYTOSIS IN BONE MARROW OR SPLEEN OR LYMPH NODES AND NO EVIDENCE OF MALIGNANCY, (F) LOW OR ABSENT NATURAL KILLER-CELL ACTIVITY, (G) FERRITIN LEVEL OF 500 MCG/L OR GREATER, (H) SOLUBLE CD25 LEVEL OF 2,400 U/ML OR GREATER.
Age Restrictions	
Prescriber Restrictions	HLH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN IMMUNOLOGIST, HEMATOLOGIST, OR ONCOLOGIST.
Coverage Duration	INITIAL AND RENEWAL: 8 WEEKS.

PA Criteria	Criteria Details
Other Criteria	HLH: INITIAL: 1) CONCURRENT THERAPY WITH DEXAMETHASONE, AND 2) ONE OF THE FOLLOWING: (A) HAS REFRACTORY, RECURRENT, OR PROGRESSIVE DISEASE, OR (B) HAD A TRIAL OF OR INTOLERANCE TO CONVENTIONAL HLH THERAPY (I.E., CHEMOTHERAPY, STEROIDS, IMMUNOTHERAPY). RENEWAL: 1) HAS NOT RECEIVED SUCCESSFUL HEMATOPOIETIC STEM CELL TRANSPLANTATION, AND 2) DEMONSTRATED IMPROVED IMMUNE SYSTEM RESPONSE FROM BASELINE (E.G., RESOLUTION OF FEVER, DECREASED SPLENOMEGALY, IMPROVEMENT IN CNS SYMPTOMS, IMPROVED CBC, INCREASED FIBRINOGEN LEVELS, REDUCED D-DIMER, REDUCED FERRITIN, REDUCED SOLUBLE CD25 LEVELS.)
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ENASIDENIB

Products Affected

• IDHIFA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ENCORAFENIB

Products Affected

• BRAFTOVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ENTRECTINIB

Products Affected

ROZLYTREK ORAL CAPSULE 100
 MG, 200 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ENTRECTINIB PELLETS

Products Affected

• ROZLYTREK ORAL PELLETS IN PACKET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC NON-SMALL CELL LUNG CANCER (NSCLC), SOLID TUMORS: 1) TRIAL OF OR CONTRAINDICATION TO ROZLYTREK CAPSULES MADE INTO AN ORAL SUSPENSION, AND 2) DIFFICULTY OR UNABLE TO SWALLOW CAPSULES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ENZALUTAMIDE

- XTANDI ORAL CAPSULE
- XTANDI ORAL TABLET 40 MG, 80 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	INITIAL: NON-METASTATIC CASTRATION-RESISTANT PROSTATE CANCER (NMCRPC): HIGH RISK PROSTATE CANCER (I.E. RAPIDLY INCREASING PROSTATE SPECIFIC ANTIGEN [PSA] LEVELS). NON-METASTATIC CASTRATION- SENSITIVE PROSTATE CANCER (NMCSPC): HIGH RISK FOR METASTASIS (I.E. PSA DOUBLED OVER 9 MONTHS OR LESS). METASTATIC CRPC (MCRPC), NMCRPC, METASTATIC CSPC (MCSPC), NMCSPC : 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG. RENEWAL: MCRPC, NMCRPC, MCSPC: 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GNRH ANALOG.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

EPCORITAMAB-BYSP

Products Affected

• EPKINLY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

EPOETIN ALFA-EPBX

Products Affected

 RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: CHRONIC KIDNEY DISEASE (CKD), ANEMIA RELATED TO ZIDOVUDINE, OR CANCER CHEMOTHERAPY: HEMOGLOBIN LEVEL OF LESS THAN 10G/DL. ELECTIVE, NON-CARDIAC, NON-VASCULAR SURGERY: HEMOGLOBIN LEVEL LESS THAN 13G/DL. RENEWAL: 1) CKD IN ADULTS NOT ON DIALYSIS: (A) HEMOGLOBIN LEVEL IS LESS THAN 10G/DL, OR (B) HEMOGLOBIN LEVEL HAS REACHED 10G/DL AND THE DOSE IS BEING REDUCED/INTERRUPTED TO DECREASE THE NEED FOR BLOOD TRANSFUSIONS. CKD IN PEDIATRIC PATIENTS: (A) HEMOGLOBIN LEVEL IS LESS THAN 10G/DL, OR (B) HEMOGLOBIN LEVEL HAS APPROACHED OR EXCEEDS 12G/DL AND THE DOSE IS BEING REDUCED/INTERRUPTED TO DECREASE THE NEED FOR BLOOD TRANSFUSIONS. ANEMIA RELATED TO ZIDOVUDINE: HEMOGLOBIN LEVEL BETWEEN 10G/DL AND 12G/DL. CANCER CHEMOTHERAPY: 1) HEMOGLOBIN LEVEL OF LESS THAN 10 G/DL, OR 2) THE HEMOGLOBIN LEVEL DOES NOT EXCEED A LEVEL NEEDED TO AVOID RBC TRANSFUSION.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	ANEMIA FROM CHEMO/CKD WITHOUT DIALYSIS/ZIDOVUDINE: INITIAL/RENEWAL: 12 MONTHS. SURGERY: 1 MONTH.

PA Criteria	Criteria Details
Other Criteria	RENEWAL: CKD: NOT RECEIVING DIALYSIS TREATMENT. THIS DRUG MAY BE EITHER BUNDLED WITH AND COVERED UNDER END STAGE RENAL DISEASE DIALYSIS RELATED SERVICES OR COVERED UNDER MEDICARE D DEPENDING UPON THE CIRCUMSTANCES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

EPOPROSTENOL IV

Products Affected

• epoprostenol

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL AND RENEWAL: 12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ERDAFITINIB

Products Affected

• BALVERSA ORAL TABLET 3 MG, 4 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ERLOTINIB

Products Affected

• erlotinib oral tablet 100 mg, 150 mg, 25 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC NSCLC WITH EGFR MUTATION: NOT ON CONCURRENT THERAPY WITH AN EGFR TYROSINE KINASE-INHIBITOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ESKETAMINE

Products Affected

• SPRAVATO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: TREATMENT-RESISTANT DEPRESSION (TRD), MAJOR DEPRESSIVE DISORDER (MDD): PRESCRIBED BY OR IN CONSULTATION WITH A PSYCHIATRIST.
Coverage Duration	INITIAL: TRD: 3 MONTHS. MDD: 4 WEEKS. RENEWAL: TRD, MDD: 12 MONTHS.
Other Criteria	INITIAL: TRD: 1) NON-PSYCHOTIC, UNIPOLAR DEPRESSION, 2) NO ACTIVE SUBSTANCE ABUSE, AND 3) ADEQUATE TRIAL (AT LEAST 4 WEEKS) OF AT LEAST TWO ANTIDEPRESSANT AGENTS FROM DIFFERENT CLASSES THAT ARE INDICATED FOR DEPRESSION. MDD: 1) NON- PSYCHOTIC, UNIPOLAR DEPRESSION, AND 2) NO ACTIVE SUBSTANCE ABUSE. RENEWAL: TRD, MDD: DEMONSTRATED CLINICAL BENEFIT (IMPROVEMENT IN DEPRESSION) COMPARED TO BASELINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ETANERCEPT

- ENBREL
- ENBREL MINI
- ENBREL SURECLICK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, FACE OR GENITAL AREA.
Age Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), ANKYLOSING SPONDYLITIS (AS): 18 YEARS OR OLDER. PSORIATIC ARTHRITIS (PSA): 2 YEARS OR OLDER.
Prescriber Restrictions	INITIAL: RA, POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA), AS: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSA: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: TRIAL OF OR CONTRAINDICATION TO 3 MONTHS OF TREATMENT WITH ONE DMARD (DISEASE- MODIFYING ANTIRHEUMATIC DRUG) - IF PATIENT TRIED METHOTREXATE, THEN TRIAL AT A DOSE GREATER THAN OR EQUAL TO 20 MG PER WEEK OR MAXIMALLY TOLERATED DOSE IS REQUIRED. PJIA, PSA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD. PSO: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. RENEWAL: RA, PJIA, PSA, AS, PSO: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ETEPLIRSEN

Products Affected

• EXONDYS-51

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	DUCHENNE MUSCULAR DYSTROPHY (DMD): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	DMD: INITIAL: PATIENT IS AMBULATORY AND CURRENTLY RECEIVING TREATMENT WITH OR HAS A CONTRAINDICATION TO CORTICOSTEROIDS. RENEWAL: MAINTAINED OR DEMONSTRATED A LESS THAN EXPECTED DECLINE IN AMBULATORY ABILITY IN MUSCLE FUNCTION ASSESSMENTS OR OTHER MUSCLE FUNCTION (I.E., PULMONARY OR CARDIAC FUNCTION). THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

EVEROLIMUS-AFINITOR

- everolimus (antineoplastic) oral tablet 10 mg, 2.5 mg, 5 mg, 7.5 mg
- torpenz oral tablet 10 mg, 2.5 mg, 5 mg, 7.5 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

EVEROLIMUS-AFINITOR DISPERZ

Products Affected

• *everolimus (antineoplastic) oral tablet for suspension*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FECAL MICROBIOTA CAPSULE

Products Affected

• VOWST

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	30 DAYS
Other Criteria	CLOSTRIDIOIDES DIFFICILE INFECTION (CDI): 1) HAS NOT PREVIOUSLY RECEIVED VOWST: COMPLETION OF ANTIBIOTIC TREATMENT FOR RECURRENT CDI (AT LEAST 3 CDI EPISODES), OR 2) PREVIOUSLY RECEIVED VOWST: (A) TREATMENT FAILURE (DEFINED AS THE PRESENCE OF CDI DIARRHEA WITHIN 8 WEEKS OF FIRST DOSE OF VOWST AND A POSITIVE STOOL TEST FOR C. DIFFICILE), AND (B) HAS NOT RECEIVED MORE THAN ONE TREATMENT COURSE OF VOWST WHICH WAS AT LEAST 12 DAYS AND NOT MORE THAN 8 WEEKS PRIOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FEDRATINIB

Products Affected

• INREBIC

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	MYELOFIBROSIS: INITIAL: TRIAL OF OR CONTRAINDICATION TO JAKAFI (RUXOLITINIB). RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FENFLURAMINE

Products Affected

• FINTEPLA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: DRAVET SYNDROME, LENNOX-GASTAUT SYNDROME (LGS): PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	DRAVET SYNDROME: INITIAL/RENEWAL: 12 MONTHS. LGS: 12 MONTHS.
Other Criteria	INITIAL: LGS: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING ANTIEPILEPTIC MEDICATIONS: RUFINAMIDE, FELBAMATE, CLOBAZAM, TOPIRAMATE, LAMOTRIGINE, CLONAZEPAM. RENEWAL: DRAVET SYNDROME: PATIENT HAS SHOWN CONTINUED CLINICAL BENEFIT (E.G. REDUCTION OF SEIZURES, REDUCED LENGTH OF SEIZURES, SEIZURE CONTROL MAINTAINED).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FENTANYL CITRATE

Products Affected

• fentanyl citrate buccal lozenge on a handle

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	CANCER RELATED PAIN: 1) CURRENTLY ON A MAINTENANCE DOSE OF CONTROLLED-RELEASE OPIOID PAIN MEDICATION, AND 2) TRIAL OF OR CONTRAINDICATION TO AT LEAST ONE IMMEDIATE- RELEASE ORAL OPIOID PAIN AGENT OR PATIENT HAS DIFFICULTY SWALLOWING TABLETS/CAPSULES. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FILGRASTIM-AAFI

Products Affected

• NIVESTYM

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FILGRASTIM-AYOW

Products Affected

• RELEUKO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENT NIVESTYM, WHERE INDICATIONS ALIGN.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FILGRASTIM-SNDZ

Products Affected

• ZARXIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENT: NIVESTYM.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FINASTERIDE/TADALAFIL

Products Affected

• ENTADFI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	BENIGN PROSTATIC HYPERPLASIA (BPH): 18 YEARS OF AGE OR OLDER
Prescriber Restrictions	
Coverage Duration	26 WEEKS
Other Criteria	BPH: 1) TRIAL OF OR CONTRAINDICATION TO ONE 5- ALPHA-REDUCTASE INHIBITOR, AND 2) TRIAL OF OR CONTRAINDICATION TO TADALAFIL 2.5MG OR TADALAFIL 5MG.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FINERENONE

Products Affected

• KERENDIA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FINGOLIMOD

Products Affected

- *fingolimod*GILENYA ORAL CAPSULE 0.25 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FINGOLIMOD LAURYL SULFATE

Products Affected

TASCENSO ODT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	MULTIPLE SCLEROSIS (MS): (1) UNABLE TO SWALLOW FINGOLIMOD CAPSULES, AND (2) TRIAL OF OR CONTRAINDICATION TO FINGOLIMOD CAPSULES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FOSTAMATINIB

Products Affected

• TAVALISSE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	CHRONIC IMMUNE THROMBOCYTOPENIA (ITP): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR IMMUNOLOGIST.
Coverage Duration	INITIAL: 3 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	ITP: RENEWAL: PATIENT HAS SHOWN A CLINICAL RESPONSE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FREMANEZUMAB-VFRM

Products Affected

- AJOVY AUTOINJECTOR
- AJOVY SYRINGE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	MIGRAINE PREVENTION: INITIAL: 1) TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING PREVENTIVE MIGRAINE TREATMENTS: DIVALPROEX SODIUM, TOPIRAMATE, PROPRANOLOL, TIMOLOL, AND 2) NO CONCURRENT USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION. RENEWAL: 1) REDUCTION IN MIGRAINE OR HEADACHE FREQUENCY, MIGRAINE SEVERITY, OR MIGRAINE DURATION WITH THERAPY, AND 2) NO CONCURRENT USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FRUQUINTINIB

Products Affected

 FRUZAQLA ORAL CAPSULE 1 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

FUTIBATINIB

Products Affected

• LYTGOBI ORAL TABLET 12 MG/DAY (4 MG X 3), 16 MG/DAY (4 MG X 4), 20 MG/DAY (4 MG X 5)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INTRAHEPATIC CHOLANGIOCARCINOMA (ICCA): COMPLETE A COMPREHENSIVE OPHTHALMOLOGICAL EXAMINATION, INCLUDING OPTICAL COHERENCE TOMOGRAPHY (OCT), PRIOR TO THE INITIATION OF THERAPY AND AT THE RECOMMENDED SCHEDULED INTERVALS.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GALCANEZUMAB-GNLM

Products Affected

• EMGALITY PEN

EMGALITY SYRINGE SUBCUTANEOUS SYRINGE 120 MG/ML, 300 MG/3 ML (100 MG/ML X

PA Criteria **Criteria Details** Exclusion Criteria **Required Medical** Information Age Restrictions Prescriber Restrictions Coverage **INITIAL: MIGRAINE PREVENTION: 6 MOS. EPISODIC Duration** CLUSTER HEADACHE: 3 MOS. RENEWAL (ALL): 12 MOS. **Other Criteria INITIAL: MIGRAINE PREVENTION: 1) NO CONCURRENT** USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION, AND 2) TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING PREVENTIVE MIGRAINE TREATMENTS: DIVALPROEX SODIUM, TOPIRAMATE, PROPRANOLOL, TIMOLOL. RENEWAL: MIGRAINE PREVENTION: 1) REDUCTION IN MIGRAINE OR HEADACHE FREQUENCY, MIGRAINE SEVERITY, OR MIGRAINE DURATION WITH THERAPY, AND 2) NO CONCURRENT USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION. EPISODIC CLUSTER HEADACHE: IMPROVEMENT IN EPISODIC CLUSTER HEADACHE FREQUENCY AS COMPARED TO BASELINE. Indications All FDA-approved Indications. Off Label Uses Part B No Prerequisite

3)

GANAXOLONE

Products Affected

• ZTALMY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GEFITINIB

Products Affected

• gefitinib

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC NSCLC WITH EGFR MUTATION: NOT ON CONCURRENT THERAPY WITH AN EGFR TYROSINE KINASE-INHIBITOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GILTERITINIB

Products Affected

• XOSPATA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GIVOSIRAN

Products Affected

• GIVLAARI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	ACUTE HEPATIC PORPHYRIA (AHP): INITIAL: GENETIC CONFIRMATION OF MUTATION OR ELEVATED URINARY OR PLASMA PBG (PORPHOBILINOGEN) OR ALA (AMINOLEVULINIC ACID).
Age Restrictions	
Prescriber Restrictions	ACUTE HEPATIC PORPHYRIA (AHP): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A GENETICIST, HEPATOLOGIST, HEMATOLOGIST, GASTROENTEROLOGIST, NEUROLOGIST, DERMATOLOGIST, OR A HEALTHCARE PROVIDER EXPERIENCED IN MANAGING AHP.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS
Other Criteria	AHP: INITIAL: EXPERIENCED TWO OR MORE AHP ATTACKS IN THE PAST 12 MONTHS. RENEWAL: 1) ACHIEVED OR MAINTAINED CLINICAL BENEFIT COMPARED TO BASELINE, AND 2) HAS NOT RECEIVED A LIVER TRANSPLANT. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GLASDEGIB

Products Affected

 DAURISMO ORAL TABLET 100 MG, 25 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GLATIRAMER

Products Affected

- COPAXONE SUBCUTANEOUS SYRINGE 20 MG/ML, 40 MG/ML
- glatiramer subcutaneous syringe 20 mg/ml, 40 mg/ml

• glatopa subcutaneous syringe 20 mg/ml, 40 mg/ml

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GLECAPREVIR/PIBRENTASVIR

Products Affected

• MAVYRET ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HCV RNA LEVEL WITHIN PAST 6 MONTHS
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE.
Other Criteria	1) CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE, 2) TRIAL OF A PREFERRED FORMULARY ALTERNATIVE INCLUDING HARVONI OR EPCLUSA WHEN THESE AGENTS ARE CONSIDERED ACCEPTABLE FOR TREATMENT OF THE SPECIFIC GENOTYPE PER AASLD/IDSA GUIDANCE, 3) NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING MEDICATIONS: CARBAMAZEPINE, RIFAMPIN, ETHINYL ESTRADIOL-CONTAINING MEDICATION, ATAZANAVIR, DARUNAVIR, LOPINAVIR, RITONAVIR, EFAVIRENZ, ATORVASTATIN, LOVASTATIN, SIMVASTATIN, ROSUVASTATIN AT DOSES GREATER THAN 10MG, CYCLOSPORINE AT DOSES GREATER THAN 100MG PER DAY, EPCLUSA, HARVONI, VOSEVI, OR ZEPATIER, AND 4) DOES NOT HAVE MODERATE OR SEVERE HEPATIC IMPAIRMENT (CHILD PUGH B OR C).
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

GLP1-DULAGLUTIDE

Products Affected

• TRULICITY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GLP1-SEMAGLUTIDE

Products Affected

 OZEMPIC SUBCUTANEOUS PEN INJECTOR 0.25 MG OR 0.5 MG (2 MG/3 ML), 0.25 MG OR 0.5 MG(2 MG/1.5 ML), 1 MG/DOSE (4 MG/3 ML), 2 MG/DOSE (8 MG/3 ML)

• RYBELSUS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GLP1-TIRZEPATIDE

Products Affected

• MOUNJARO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GLYCEROL PHENYLBUTYRATE

Products Affected

• RAVICTI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	UREA CYCLE DISORDER (UCD): INITIAL: DIAGNOSIS IS CONFIRMED BY ENZYMATIC, BIOCHEMICAL OR GENETIC TESTING
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	UCD: INITIAL: TRIAL OF OR CONTRAINDICATION TO SODIUM PHENYLBUTYRATE. RENEWAL: PATIENT HAS CLINICAL BENEFIT FROM BASELINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GOSERELIN

Products Affected

• ZOLADEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	ENDOMETRIOSIS: DIAGNOSIS IS CONFIRMED VIA SURGICAL OR DIRECT VISUALIZATION (E.G., PELVIC ULTRASOUND) OR HISTOPATHOLOGICAL CONFIRMATION (E.G., LAPAROSCOPY OR LAPAROTOMY) IN THE LAST 10 YEARS.
Age Restrictions	
Prescriber Restrictions	ENDOMETRIOSIS: PRESCRIBED BY OR IN CONSULTATION WITH AN OBSTETRICIAN/GYNECOLOGIST.
Coverage Duration	STAGE B2-C PROSTATIC CARCINOMA: 4 MOS. ENDOMETRIOSIS: 6 MOS PER LIFETIME. ALL OTHERS: 12 MONTHS.
Other Criteria	ENDOMETRIOSIS: 1) NO CONCURRENT USE WITH ANOTHER GNRH-MODULATING AGENT, 2) TRIAL OF OR CONTRAINDICATION TO NSAID AND PROGESTIN- CONTAINING PREPARATION, AND 3) HAS NOT RECEIVED A TOTAL OF 6 MONTHS OF TREATMENT PER LIFETIME. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

GUSELKUMAB

Products Affected

• TREMFYA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, FACE OR GENITAL AREA.
Age Restrictions	
Prescriber Restrictions	INITIAL: PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR DERMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	INITIAL: PSA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG). PSO: TRIAL OF OR CONTRAINDICATION ONE CONVENTIONAL THERAPY SUCH AS A PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. RENEWAL: PSO, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH CONCENTRATION ORAL OPIOID SOLUTIONS

Products Affected

- morphine concentrate oral solution
- oxycodone oral concentrate

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	OPIOID TOLERANT: 12 MONTHS. HOSPICE, PALLIATIVE CARE OR END OF LIFE CARE: LIFETIME.
Other Criteria	1) OPIOID TOLERANT (I.E. PREVIOUS USE OF 60 MG ORAL MORPHINE PER DAY, 25 MCG TRANSDERMAL FENTANYL PER HOUR, 30 MG ORAL OXYCODONE PER DAY, 8 MG ORAL HYDROMORPHONE PER DAY, 25 MG ORAL OXYMORPHONE PER DAY, 60 MG ORAL HYDROCODONE PER DAY, OR AN EQUIANALGESIC DOSE OF ANOTHER OPIOID) AND HAS TROUBLE SWALLOWING OPIOID TABLETS, CAPSULES, OR LARGE VOLUMES OF LIQUID, OR 2) ENROLLED IN HOSPICE OR PALLIATIVE CARE OR END OF LIFE CARE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY - CARBINOXAMINE

Products Affected

- carbinoxamine maleate oral liquid
- carbinoxamine maleate oral tablet 4 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY - CYPROHEPTADINE

Products Affected

• cyproheptadine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -BUTALBITAL-CONTAINING AGENTS

Products Affected

- ascomp with codeine
- butalbital-acetaminop-caf-cod
- butalbital-acetaminophen oral tablet 50-325 codeine-butalbital-asa-caff mg
- *butalbital-acetaminophen-caff oral capsule* 50-325-40 mg
- butalbital-acetaminophen-caff oral tablet
- butalbital-aspirin-caffeine
- tencon
- *zebutal*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS ARE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -CLEMASTINE

Products Affected

• clemastine oral tablet

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -CONJUGATED ESTROGEN

Products Affected

• PREMARIN ORAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	VULVAR/VAGINAL ATROPHY, OSTEOPOROSIS AND VASOMOTOR SYMPTOMS OF MENOPAUSE: PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HYPOESTROGENISM TREATMENT, PALLIATIVE TREATMENT, AND HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -DIPYRIDAMOLE

Products Affected

• dipyridamole oral

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -DISOPYRAMIDE

Products Affected

• disopyramide phosphate oral capsule

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -ESTRADIOL

Products Affected

• dotti

• lyllana

- estradiol oral
- estradiol transdermal patch semiweekly
- estradiol transdermal patch weekly

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	VULVAR/VAGINAL ATROPHY, OSTEOPOROSIS AND VASOMOTOR SYMPTOMS OF MENOPAUSE: PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HYPOESTROGENISM TREATMENT, PALLIATIVE TREATMENT, AND HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -ESTRADIOL-NORETHINDRONE

- amabelz
- estradiol-norethindrone acet oral tablet 0.5-0.1 mg
- mimvey

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	VULVAR/VAGINAL ATROPHY, OSTEOPOROSIS, AND VASOMOTOR SYMPTOMS OF MENOPAUSE: PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HYPOESTROGENISM TREATMENT AND HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -ESTROGEN-BAZEDOXIFENE

Products Affected

• DUAVEE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -ESTROGEN-MEDROXYPROGESTERONE

- PREMPHASE
- PREMPRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -**GLYBURIDE FORMULATIONS**

- glyburide
- glyburide micronized glyburide-metformin

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	TRIAL OF GLIMEPIRIDE, GLIPIZIDE, OR PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS ARE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -KETOROLAC

- ketorolac injection solution 15 mg/ml, 30 mg/ml, 30 mg/ml (1 ml)
- ketorolac injection syringe 15 mg/ml, 30 mg/ml
- ketorolac intramuscular solution
- *ketorolac intramuscular syringe*
- ketorolac oral

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	30 DAYS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -NORETHINDRONE-ESTRADIOL

- fyavolv
- jinteli
- norethindrone ac-eth estradiol oral tablet 0.5-2.5 mg-mcg, 1-5 mg-mcg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -PHENOBARBITAL

Products Affected

• phenobarbital

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	FOR TREATMENT OF EPILEPSY/SEIZURES IN PATIENTS WHO ARE NEWLY PRESCRIBED PHENOBARBITAL: PATIENT HAS NOT RESPONDED TO AT LEAST ONE ANTICONVULSANT OR PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -PROMETHAZINE

- promethazine injection solution
- promethazine oral
- promethazine rectal
- promethegan

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRURITUS/URTICARIA/SEASONAL/PERENNIAL ALLERGY: 1) TRIAL OF OR CONTRAINDICATION TO A NON- SEDATING ANTIHISTAMINE SUCH AS LEVOCETIRIZINE, OR 2) PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. NAUSEA AND VOMITING: PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH-RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS REQUIRE PHYSICIAN ATTESTATION THAT REQUESTED MEDICATION IS USED TO TREAT A DIAGNOSIS UNRELATED TO THE TERMINAL ILLNESS OR RELATED CONDITION, AND ARE APPROVED WITHOUT TRIAL OF FORMULARY ALTERNATIVES NOR REQUIRING PRESCRIBER ACKNOWLEDGEMENT.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -SCOPOLAMINE

Products Affected

• scopolamine base

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS REQUIRE PHYSICIAN ATTESTATION THAT REQUESTED MEDICATION IS USED TO TREAT A DIAGNOSIS UNRELATED TO THE TERMINAL ILLNESS OR RELATED CONDITION, AND ARE APPROVED WITHOUT REQUIRING PRESCRIBER ACKNOWLEDGEMENT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY -SKELETAL MUSCLE RELAXANTS

- chlorzoxazone oral tablet 250 mg, 500 mg, 750 mg
- cyclobenzaprine oral tablet 10 mg, 5 mg
- methocarbamol oral tablet 500 mg, 750 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED A HIGH RISK MEDICATION FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY-DIPHENHYDRAMINE ELIXIR

Products Affected

• diphenhydramine hcl oral elixir

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRURITUS OR URTICARIA: TRIAL OF OR CONTRAINDICATION TO A NON-SEDATING ANTIHISTAMINE SUCH AS LEVOCETIRIZINE. INSOMNIA: TRIAL OF SILENOR AND BELSOMRA. MOTION SICKNESS AND ANTIPARKINSONISM: PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS AND ANAPHYLACTIC REACTIONS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY-DIPHENOXYLATE-ATROPINE

Products Affected

• diphenoxylate-atropine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY-INDOMETHACIN

- indomethacin oral capsule 25 mg, 50 mg
- indomethacin oral capsule, extended release

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY-MEGESTROL

- megestrol oral suspension 400 mg/10 ml (40 mg/ml)
- megestrol oral tablet

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HIGH RISK DRUGS IN THE ELDERLY-PAROXETINE

- paroxetine hcl oral suspension
- paroxetine hcl oral tablet

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0-64 YEARS.
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS THAT THE DRUG IS CONSIDERED HIGH RISK FOR PATIENTS 65 YEARS AND OLDER. HOSPICE PATIENTS WILL BE APPROVED WITHOUT REQUIRING ADDITIONAL CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

HISTRELIN-SUPPRELIN LA

Products Affected

• SUPPRELIN LA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CENTRAL PRECOCIOUS PUBERTY (CPP): INITIAL: FEMALES: ELEVATED LEVELS OF FOLLICLE- STIMULATING HORMONE (FSH) GREATER THAN 4.0 MIU/ML AND LUTEINIZING HORMONE (LH) LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS. MALES: ELEVATED LEVELS OF FSH GREATER THAN 5.0 MIU/ML AND LH LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS.
Age Restrictions	
Prescriber Restrictions	CPP: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	CPP: INITIAL: FEMALES: 1) YOUNGER THAN 8 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR BREAST DEVELOPMENT AND PUBIC HAIR GROWTH. MALES: 1) YOUNGER THAN 9 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR GENITAL DEVELOPMENT AND PUBIC HAIR GROWTH. RENEWAL: 1) TANNER STAGING AT INITIAL DIAGNOSIS HAS STABILIZED OR REGRESSED DURING THREE SEPARATE MEDICAL VISITS IN THE PREVIOUS YEAR, AND 2) HAS NOT REACHED ACTUAL AGE WHICH CORRESPONDS TO CURRENT PUBERTAL AGE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

IBRUTINIB

Products Affected

• IMBRUVICA ORAL CAPSULE 140 MG, 70 MG 280 MG, 420 MG

- IMBRUVICA ORAL SUSPENSION
- IMBRUVICA ORAL TABLET 140 MG,

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IBUPROFEN-FAMOTIDINE

Products Affected

• ibuprofen-famotidine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	TRIAL OF ONE OF THE FOLLOWING GENERIC, FEDERAL LEGEND HISTAMINE H2-RECEPTOR ANTAGONISTS: FAMOTIDINE, CIMETIDINE, OR NIZATIDINE, AND TRIAL OF GENERIC, FEDERAL LEGEND IBUPROFEN.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ICATIBANT

- icatibant
- sajazir

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HEREDITARY ANGIOEDEMA (HAE): DIAGNOSIS CONFIRMED BY COMPLEMENT TESTING.
Age Restrictions	
Prescriber Restrictions	HAE: PRESCRIBED BY OR IN CONSULTATION WITH AN ALLERGIST, IMMUNOLOGIST, OR HEMATOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	HAE: NO CONCURRENT USE WITH OTHER MEDICATIONS FOR TREATMENT OF ACUTE HAE ATTACKS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IDELALISIB

Products Affected

• ZYDELIG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IMATINIB

Products Affected

• imatinib oral tablet 100 mg, 400 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	ADJUVANT GASTROINTESTINAL STROMAL TUMOR TREATMENT: 36 MONTHS. ALL OTHER DIAGNOSES: 12 MONTHS.
Other Criteria	PHILADELPHIA CHROMOSOME POSITIVE CHRONIC MYELOID LEUKEMIA: PATIENT HAS NOT RECEIVED A PREVIOUS TREATMENT WITH ANOTHER TYROSINE KINASE INHIBITOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INFLIXIMAB

Products Affected

• infliximab

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, GENITAL AREA, OR FACE.
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), ANKYLOSING SPONDYLITIS (AS): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PSA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OTEZLA. PSO: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, SKYRIZI, TREMFYA, OTEZLA. AS: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, COSENTYX, ENBREL, XELJANZ, RINVOQ. CD: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, RINVOQ, SKYRIZI. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, RINVOQ, SKYRIZI. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI. RENEWAL: RA, AS, PSO, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INFLIXIMAB-ABDA

Products Affected

• RENFLEXIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, GENITAL AREA, OR FACE.
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), ANKYLOSING SPONDYLITIS (AS): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PSA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OTEZLA. PSO: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, SKYRIZI, TREMFYA, OTEZLA. AS: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, COSENTYX, ENBREL, XELJANZ, RINVOQ. CD: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ, SKYRIZI, RENEWAL: RA, AS, PSO, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INFLIXIMAB-AXXQ

Products Affected

• AVSOLA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, GENITAL AREA, OR FACE.
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), ANKYLOSING SPONDYLITIS (AS): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PSA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OTEZLA. PSO: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, SKYRIZI, TREMFYA, OTEZLA. AS: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, COSENTYX, ENBREL, XELJANZ, RINVOQ. CD: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ, SKYRIZI. RENEWAL: RA, AS, PSO, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INFLIXIMAB-DYYB

Products Affected

• INFLECTRA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, GENITAL AREA, OR FACE.
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), ANKYLOSING SPONDYLITIS (AS): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PSA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OTEZLA. PSO: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, SKYRIZI, TREMFYA, OTEZLA. AS: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, COSENTYX, ENBREL, XELJANZ, RINVOQ. CD: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ. UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: HUMIRA, STELARA, SKYRIZI, RINVOQ, SKYRIZI. RENEWAL: RA, AS, PSO, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INFLIXIMAB-DYYB - SQ

Products Affected

• ZYMFENTRA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: ULCERATIVE COLITIS (UC), CROHNS DISEASE (CD): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: UC: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI. CD: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, STELARA, SKYRIZI, RINVOQ.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INTERFERON FOR MS-AVONEX

- AVONEX INTRAMUSCULAR PEN INJECTOR KIT
- AVONEX INTRAMUSCULAR SYRINGE KIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INTERFERON FOR MS-BETASERON

Products Affected

• BETASERON SUBCUTANEOUS KIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INTERFERON FOR MS-PLEGRIDY

Products Affected

 PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML
 PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INTERFERON GAMMA-1B

Products Affected

• ACTIMMUNE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: CHRONIC GRANULOMATOUS DISEASE (CGD): PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST, INFECTIOUS DISEASE SPECIALIST, OR IMMUNOLOGIST. SEVERE MALIGNANT OSTEOPETROSIS (SMO): PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	RENEWAL: CGD, SMO: 1) DEMONSTRATED CLINICAL BENEFIT COMPARED TO BASELINE, AND 2) HAS NOT RECEIVED HEMATOPOIETIC CELL TRANSPLANTATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IPILIMUMAB

Products Affected

• YERVOY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: UNRESECT/MET MEL: 4MO, RCC/CRC/HCC: 3MO, ALL OTHERS: 12MO. INITIAL/RENEWAL: CUTAN MEL: 6MO
Other Criteria	RENEWAL: ADJUVANT CUTANEOUS MELANOMA: NO EVIDENCE OF DISEASE RECURRENCE (DEFINED AS THE APPEARANCE OF ONE OR MORE NEW MELANOMA LESIONS: LOCAL, REGIONAL OR DISTANT METASTASIS). THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ITRACONAZOLE SOLUTION

Products Affected

• itraconazole oral solution

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 MONTHS
Other Criteria	ESOPHAGEAL CANDIDIASIS AND OROPHARYNGEAL CANDIDIASIS: TRIAL OF OR CONTRAINDICATION TO FLUCONAZOLE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IVACAFTOR

Products Affected

• KALYDECO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CYSTIC FIBROSIS (CF): INITIAL: CONFIRMED MUTATION IN CFTR GENE ACCEPTABLE FOR THE TREATMENT OF CYSTIC FIBROSIS
Age Restrictions	
Prescriber Restrictions	CF: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST OR CYSTIC FIBROSIS EXPERT
Coverage Duration	INITIAL: 12 MONTHS. RENEWAL: LIFETIME
Other Criteria	CF: INITIAL: NOT HOMOZYGOUS FOR F508DEL MUTATION IN CFTR GENE. RENEWAL: 1) MAINTAINED, IMPROVED, OR DEMONSTRATED LESS THAN EXPECTED DECLINE IN FEV1 OR BODY MASS INDEX (BMI), OR 2) REDUCTION IN NUMBER OF PULMONARY EXACERBATIONS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IVOSIDENIB

Products Affected

• TIBSOVO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IXAZOMIB

Products Affected

• NINLARO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

IXEKIZUMAB

- TALTZ AUTOINJECTOR
- TALTZ SYRINGE SUBCUTANEOUS SYRINGE 80 MG/ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, FACE, OR GENITAL AREA. NON-RADIOGRAPHIC AXIAL SPONDYLOARTHRITIS (NR-AXSPA): 1) C-REACTIVE PROTEIN (CRP) LEVELS ABOVE THE UPPER LIMIT OF NORMAL, OR 2) SACROILIITIS ON MAGNETIC RESONANCE IMAGING (MRI).
Age Restrictions	
Prescriber Restrictions	INITIAL: PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR DERMATOLOGIST. ANKYLOSING SPONDYLITIS (AS), NR-AXSPA: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: PSO: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS, WHERE AGES ALIGN: COSENTYX, ENBREL, HUMIRA, STELARA, SKYRIZI, TREMFYA, OTEZLA. PSA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, XELJANZ, RINVOQ, SKYRIZI, TREMFYA, OTEZLA. AS: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: ENBREL, HUMIRA, COSENTYX, XELJANZ, RINVOQ. NR-AXSPA: 1) TRIAL OF OR CONTRAINDICATION TO BOTH OF THE PREFERRED AGENTS: COSENTYX, RINVOQ, OR 2) TRIAL OF COSENTYX AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. RENEWAL: PSO, PSA, AS, NR- AXSPA: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LANADELUMAB-FLYO

Products Affected

 TAKHZYRO SUBCUTANEOUS SOLUTION (150 MG/ML)

• TAKHZYRO SUBCUTANEOUS SYRINGE 150 MG/ML, 300 MG/2 ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HEREDITARY ANGIOEDEMA (HAE): INITIAL: DIAGNOSIS CONFIRMED BY COMPLEMENT TESTING.
Age Restrictions	
Prescriber Restrictions	HAE: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ALLERGIST, IMMUNOLOGIST, OR HEMATOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	HAE: INITIAL: NOT ON CONCURRENT TREATMENT WITH ALTERNATIVE PROPHYLACTIC AGENT FOR HAE ATTACKS. RENEWAL: 1) IMPROVEMENT COMPARED TO BASELINE IN HAE ATTACKS (I.E., REDUCTIONS IN ATTACK FREQUENCY OR ATTACK SEVERITY), AND 2) NOT ON CONCURRENT TREATMENT WITH ALTERNATIVE PROPHYLACTIC AGENT FOR HAE ATTACKS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LANREOTIDE

Products Affected

- *lanreotide subcutaneous syringe 120 mg/0.5 ml*
- SOMATULINE DEPOT SUBCUTANEOUS SYRINGE 120

MG/0.5 ML, 60 MG/0.2 ML, 90 MG/0.3 ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	ACROMEGALY: INITIAL: THERAPY IS PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	ACROMEGALY: INITIAL: 3 MOS, RENEWAL: 12 MOS.GEP- NETS, CARCINOID SYNDROME: 12 MOS.
Other Criteria	ACROMEGALY: INITIAL: TRIAL OF OR CONTRAINDICATION TO ONE GENERIC OCTREOTIDE INJECTION. RENEWAL: 1) REDUCTION, NORMALIZATION, OR MAINTENANCE OF IGF-1 LEVELS BASED ON AGE AND GENDER, AND 2) IMPROVEMENT OR SUSTAINED REMISSION OF CLINICAL SYMPTOMS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LAPATINIB

Products Affected

• lapatinib

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LAROTRECTINIB

- VITRAKVI ORAL CAPSULE 100 MG, 25 MG
- VITRAKVI ORAL SOLUTION

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	APPROVAL FOR VITRAKVI ORAL SOLUTION: TRIAL OF VITRAKVI CAPSULES OR PATIENT IS UNABLE TO TAKE CAPSULE FORMULATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LEDIPASVIR-SOFOSBUVIR

- HARVONI ORAL PELLETS IN PACKET 33.75-150 MG, 45-200 MG
- HARVONI ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HCV RNA LEVEL WITHIN PAST 6 MONTHS.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE.
Other Criteria	1) CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE, AND 2) NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING: CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, OXCARBAZEPINE, RIFAMPIN, RIFABUTIN, RIFAPENTINE, ROSUVASTATIN, TIPRANAVIR/RITONAVIR, SOFOSBUVIR (AS A SINGLE AGENT), EPCLUSA, ZEPATIER, MAVYRET, OR VOSEVI. REQUESTS FOR HARVONI 45MG-200MG PELLETS: PATIENT IS UNABLE TO SWALLOW TABLETS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LENALIDOMIDE

Products Affected

• lenalidomide

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LENVATINIB

Products Affected

• LENVIMA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LETERMOVIR

- PREVYMIS INTRAVENOUS SOLUTION 240 MG/12 ML, 480 MG/24 ML
- PREVYMIS ORAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	HSCT: NOT AT RISK FOR LATE CMV: 4 MOS, AT RISK FOR LATE CMV: 7 MOS. KIDNEY TRANSPLANT: 7 MOS.
Other Criteria	HEMATOPOIETIC STEM CELL TRANSPLANT (HSCT): 1) THERAPY WILL BE INITIATED BETWEEN DAY 0 AND DAY 28 POST TRANSPLANT, AND 2) WILL NOT RECEIVE THE MEDICATION BEYOND 100 DAYS POST TRANSPLANT IF NOT AT RISK FOR LATE CYTOMEGALOVIRUS (CMV) INFECTION AND DISEASE, OR BEYOND 200 DAYS POST TRANSPLANT IF AT RISK FOR LATE CMV INFECTION AND DISEASE. KIDNEY TRANSPLANT: 1) THERAPY WILL BE INITIATED BETWEEN DAY 0 AND DAY 7 POST TRANSPLANT, AND 2) WILL NOT RECEIVE THE MEDICATION BEYOND 200 DAYS POST TRANSPLANT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LEUPROLIDE

Products Affected

• leuprolide subcutaneous kit

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	PROSTATE CANCER: 12 MONTHS.
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LEUPROLIDE DEPOT

Products Affected

• *leuprolide (3 month)*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LEUPROLIDE-ELIGARD

- ELIGARD
- ELIGARD (3 MONTH)
- ELIGARD (4 MONTH)
- ELIGARD (6 MONTH)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS.
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LEUPROLIDE-LUPRON DEPOT

- LUPRON DEPOT
- LUPRON DEPOT (3 MONTH)
- LUPRON DEPOT (4 MONTH)
- LUPRON DEPOT (6 MONTH)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: ENDOMETRIOSIS: DIAGNOSIS IS CONFIRMED VIA SURGICAL OR DIRECT VISUALIZATION (E.G., PELVIC ULTRASOUND) OR HISTOPATHOLOGICAL CONFIRMATION (E.G., LAPAROSCOPY OR LAPAROTOMY) IN THE LAST 10 YEARS.
Age Restrictions	
Prescriber Restrictions	INITIAL: ENDOMETRIOSIS: PRESCRIBED BY OR IN CONSULTATION WITH AN OBSTETRICIAN/GYNECOLOGIST.
Coverage Duration	PROSTATE CA: 12 MOS. UTERINE FIBROIDS: 3 MOS. ENDOMETRIOSIS: INITIAL/RENEWAL: 6 MOS.
Other Criteria	INITIAL: ENDOMETRIOSIS: 1) NO CONCURRENT USE WITH ANOTHER GNRH-MODULATING AGENT, 2) TRIAL OF OR CONTRAINDICATION TO NSAID AND PROGESTIN- CONTAINING PREPARATION, AND 3) HAS NOT RECEIVED A TOTAL OF 12 MONTHS OF TREATMENT PER LIFETIME. RENEWAL: ENDOMETRIOSIS: 1) IMPROVEMENT OF PAIN RELATED TO ENDOMETRIOSIS WHILE ON THERAPY, 2) RECEIVING CONCOMITANT ADD-BACK THERAPY (I.E., COMBINATION ESTROGEN-PROGESTIN OR PROGESTIN- ONLY CONTRACEPTIVE PREPARATION), 3) NO CONCURRENT USE WITH ANOTHER GNRH-MODULATING AGENT, AND 4) HAS NOT RECEIVED A TOTAL OF 12 MONTHS OF TREATMENT PER LIFETIME. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

LEUPROLIDE-LUPRON DEPOT-PED

- LUPRON DEPOT-PED
- LUPRON DEPOT-PED (3 MONTH)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CENTRAL PRECOCIOUS PUBERTY (CPP): INITIAL: FEMALES: ELEVATED LEVELS OF FOLLICLE- STIMULATING HORMONE (FSH) GREATER THAN 4.0 MIU/ML AND LUTEINIZING HORMONE (LH) LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS. MALES: ELEVATED LEVELS OF FSH GREATER THAN 5.0 MIU/ML AND LH LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS.
Age Restrictions	
Prescriber Restrictions	CPP: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	CPP: INITIAL: FEMALES: 1) YOUNGER THAN 8 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR BREAST DEVELOPMENT AND PUBIC HAIR GROWTH. MALES: 1) YOUNGER THAN 9 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR GENITAL DEVELOPMENT AND PUBIC HAIR GROWTH. RENEWAL: 1) TANNER STAGING AT INITIAL DIAGNOSIS HAS STABILIZED OR REGRESSED DURING THREE SEPARATE MEDICAL VISITS IN THE PREVIOUS YEAR, AND 2) HAS NOT REACHED ACTUAL AGE WHICH CORRESPONDS TO CURRENT PUBERTAL AGE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

LEVODOPA

Products Affected

• INBRIJA INHALATION CAPSULE, W/INHALATION DEVICE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	PARKINSONS DISEASE (PD): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	PD: INITIAL: 1) NOT CURRENTLY TAKING MORE THAN 1600MG OF LEVODOPA PER DAY, AND 2) PHYSICIAN HAS OPTIMIZED DRUG THERAPY FOR PARKINSONS DISEASE. RENEWAL: IMPROVEMENT WITH MOTOR FLUCTUATIONS DURING OFF EPISODES WITH THE USE OF INBRIJA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

L-GLUTAMINE

- ENDARI
- glutamine (sickle cell)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	SICKLE CELL DISEASE(SCD): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST
Coverage Duration	INITIAL: 12 MONTHS. RENEWAL: LIFETIME.
Other Criteria	SCD: INITIAL: AGES 18 YEARS OR OLDER: 1) AT LEAST 2 SICKLE CELL CRISES IN THE PAST YEAR, 2) SICKLE-CELL ASSOCIATED SYMPTOMS WHICH ARE INTERFERING WITH ACTIVITIES OF DAILY LIVING, OR 3) HISTORY OF OR HAS RECURRENT ACUTE CHEST SYNDROME. AGES 5 TO 17 YEARS: APPROVED WITHOUT ADDITIONAL CRITERIA. RENEWAL: MAINTAINED OR EXPERIENCED A REDUCTION IN ACUTE COMPLICATIONS OF SCD.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LIDOCAINE OINTMENT

Products Affected

• lidocaine topical ointment

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. THIS DRUG MAY BE EITHER BUNDLED WITH AND COVERED UNDER END STAGE RENAL DISEASE DIALYSIS RELATED SERVICES OR COVERED UNDER MEDICARE D DEPENDING UPON THE CIRCUMSTANCES.
Indications	All Medically-accepted Indications.
Off Label Uses	
Part B Prerequisite	No

LIDOCAINE PATCH

- lidocaine topical adhesive patch, medicated 5 %
- tridacaine
- ZTLIDO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	1) PAIN ASSOCIATED WITH POST-HERPETIC NEURALGIA, 2) NEUROPATHY DUE TO DIABETES MELLITUS, 3) CHRONIC BACK PAIN, OR 4) OSTEOARTHRITIS OF THE KNEE OR HIP.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All Medically-accepted Indications.
Off Label Uses	
Part B Prerequisite	No

LIDOCAINE PRILOCAINE

Products Affected

• lidocaine-prilocaine topical cream

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG MAY BE EITHER BUNDLED WITH AND COVERED UNDER END STAGE RENAL DISEASE DIALYSIS RELATED SERVICES OR COVERED UNDER MEDICARE D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All Medically-accepted Indications.
Off Label Uses	
Part B Prerequisite	No

LIDOCAINE SOLUTION

Products Affected

 lidocaine hcl mucous membrane solution 4 % (40 mg/ml)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All Medically-accepted Indications.
Off Label Uses	
Part B Prerequisite	No

LOMITAPIDE

Products Affected

• JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST, ENDOCRINOLOGIST OR LIPIDOLOGIST.
Coverage Duration	12 MONTHS

PA Criteria	Criteria Details
Other Criteria	1) DIAGNOSIS DETERMINED BY A) DEFINITE SIMON BROOME DIAGNOSTIC CRITERIA, OR B) DUTCH LIPID NETWORK CRITERIA SCORE OF 8 OR GREATER, OR C) CLINICAL DIAGNOSIS BASED ON A HISTORY OF AN UNTREATED LDL-C CONCENTRATION GREATER THAN 500 MG/DL TOGETHER WITH EITHER XANTHOMA BEFORE 10 YEARS OF AGE, OR EVIDENCE OF HEFH IN BOTH PARENTS. 2) LDL-C LEVEL GREATER THAN OR EQUAL TO 70MG/DL WHILE ON MAXIMAL DRUG TREATMENT. 3) TRIAL OF EVOLOCUMAB UNLESS THE PATIENT HAS NON- FUNCTIONING LDL RECEPTORS. 4) MEETS ONE OF THE FOLLOWING: A) TAKING A HIGH-INTENSITY STATIN (I.E., ATOR VASTATIN 40-80MG DAILY, ROSUVASTATIN 20-40MG DAILY) FOR A DURATION OF AT LEAST 8 WEEKS, B) TAKING A MAXIMALLY TOLERATED DOSE OF ANY STATIN FOR A DURATION OF AT LEAST 8 WEEKS GIVEN THAT THE PATIENT CANNOT TOLERATE A HIGH- INTENSITY STATIN, C) ABSOLUTE CONTRAINDICATION TO STATIN THERAPY (E.G., ACTIVE DECOMPENSATED LIVER DISEASE, NURSING FEMALE, PREGNANCY OR PLANS TO BECOME PREGNANT, HYPERSENSITIVITY REACTIONS), D) STATIN INTOLERANCE, OR E) TRIAL OF ROSUVASTATIN, ATORVASTATIN, OR STATIN THERAPY AT ANY DOSE AND HAS EXPERIENCED SKELETAL- MUSCLE RELATED SYMPTOMS (E.G., MYOPATHY).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LONCASTUXIMAB TESIRINE-LPYL

Products Affected

• ZYNLONTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LORLATINIB

Products Affected

 LORBRENA ORAL TABLET 100 MG, 25 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LUMACAFTOR-IVACAFTOR

- ORKAMBI ORAL GRANULES IN PACKET
- ORKAMBI ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: CYSTIC FIBROSIS (CF): CONFIRMED MUTATION IN CFTR GENE ACCEPTABLE FOR THE TREATMENT OF CF.
Age Restrictions	
Prescriber Restrictions	CF: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST OR CF EXPERT.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: LIFETIME.
Other Criteria	CF: RENEWAL: MAINTAINED, IMPROVED, OR DEMONSTRATED LESS THAN EXPECTED DECLINE IN FEV1 OR BODY MASS INDEX (BMI), OR REDUCTION IN NUMBER OF PULMONARY EXACERBATIONS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

LUMASIRAN

Products Affected

• OXLUMO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MACITENTAN

Products Affected

• OPSUMIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MARGETUXIMAB-CMKB

Products Affected

• MARGENZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MECHLORETHAMINE

Products Affected

• VALCHLOR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MEPOLIZUMAB

Products Affected

- NUCALA SUBCUTANEOUS AUTO-INJECTOR
- NUCALA SUBCUTANEOUS RECON SOLN
- NUCALA SUBCUTANEOUS SYRINGE 100 MG/ML, 40 MG/0.4 ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: ASTHMA: BLOOD EOSINOPHIL LEVEL GREATER THAN OR EQUAL TO 150 CELLS/MCL WITHIN THE PAST 12 MONTHS.
Age Restrictions	
Prescriber Restrictions	INITIAL: ASTHMA: PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN PULMONARY OR ALLERGY MEDICINE. NASAL POLYPS: PRESCRIBED BY OR IN CONSULTATION WITH AN OTOLARYNGOLOGIST, ALLERGIST OR IMMUNOLOGIST.
Coverage Duration	INITIAL: ASTHMA: 4 MO. NASAL POLYPS: 6 MO. OTHERS: 12 MO. RENEWAL: NASAL POLYPS, ASTHMA: 12 MO.

PA Criteria	Criteria Details
Other Criteria	INITIAL: ASTHMA: 1) CONCURRENT THERAPY WITH A MEDIUM, HIGH-DOSE OR MAXIMALLY TOLERATED DOSE OF AN INHALED CORTICOSTEROID AND AT LEAST ONE OTHER MAINTENANCE MEDICATION, AND 2) ONE OF THE FOLLOWING: (A) AT LEAST ONE ASTHMA EXACERBATION REQUIRING SYSTEMIC CORTICOSTEROID BURST LASTING 3 OR MORE DAYS WITHIN THE PAST 12 MONTHS OR AT LEAST ONE SERIOUS EXACERBATION REQUIRING HOSPITALIZATION OR ER VISIT WITHIN THE PAST 12 MONTHS, OR (B) POOR SYMPTOM CONTROL DESPITE CURRENT THERAPY AS EVIDENCED BY AT LEAST THREE OF THE FOLLOWING WITHIN THE PAST 4 WEEKS: DAYTIME ASTHMA SYMPTOMS MORE THAN TWICE/WEEK, ANY NIGHT WAKING DUE TO ASTHMA, SABA RELIEVER FOR SYMPTOMS MORE THAN TWICE/WEEK, ANY ACTIVITY LIMITATION DUE TO ASTHMA, AND 3) NOT CONCURRENTLY RECEIVING XOLAIR, DUPIXENT OR OTHER ANTI-IL5 BIOLOGICS WHEN THESE ARE USED FOR THE TREATMENT OF ASTHMA. NASAL POLYPS: PREVIOUS 56 DAY TRIAL OF ONE TOPICAL NASAL CORTICOSTEROID. RENEWAL: ASTHMA: 1) NOT CONCURRENTLY RECEIVING XOLAIR, DUPIXENT OR OTHER ANTI-IL5 BIOLOGICS WHEN THESE ARE USED FOR THE TREATMENT OF ASTHMA. (DONTINUED USE OF ICS AND AT LEAST ONE OTHER MAINTENANCE MEDICATION, AND 3) CLINICAL RESPONSE AS EVIDENCED BY ONE OF THE FOLLOWING: (A) REDUCTION IN ASTHMA EXACERBATIONS FROM BASELINE, (B) DECREASED UTILIZATION OF RESCUE MEDICATIONS, (C) REDUCTION IN SEVERITY OR FREQUENCY OF ASTHMA-RELATED SYMPTOMS, OR (D) INCREASE IN PERCENT PREDICTED FEVI FROM
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

METHYLNALTREXONE INJECTABLE

Products Affected

- RELISTOR SUBCUTANEOUS
 SOLUTION
- RELISTOR SUBCUTANEOUS SYRINGE 12 MG/0.6 ML, 8 MG/0.4 ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	ADVANCED ILLNESS: 6 MONTHS. CHRONIC NON-CANCER PAIN: 12 MONTHS.
Other Criteria	CHRONIC NON-CANCER PAIN: 1) HAS BEEN TAKING OPIOIDS FOR AT LEAST 4 WEEKS, AND 2) TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENTS: NALOXEGOL (MOVANTIK) AND LUBIPROSTONE (AMITIZA)
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

METHYLNALTREXONE ORAL

Products Affected

RELISTOR ORAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	OPIOID INDUCED CONSTIPATION WITH CHRONIC NON- CANCER PAIN: 1) HAS BEEN TAKING OPIOIDS FOR AT LEAST 4 WEEKS, AND 2) TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENTS: NALOXEGOL (MOVANTIK) AND LUBIPROSTONE (AMITIZA)
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MIDOSTAURIN

Products Affected

• RYDAPT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	ACUTE MYELOID LEUKEMIA: 6 MONTHS. ADVANCED SYSTEMIC MASTOCYTOSIS: 12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MIFEPRISTONE

Products Affected

• mifepristone oral tablet 300 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CUSHINGS SYNDROME (CS): INITIAL: DIAGNOSIS CONFIRMED BY ONE OF THE FOLLOWING: 1) 24-HR URINE FREE CORTISOL (2 OR MORE TESTS TO CONFIRM), 2) OVERNIGHT 1MG DEXAMETHASONE TEST, OR 3) LATE NIGHT SALIVARY CORTISOL (2 OR MORE TESTS TO CONFIRM).
Age Restrictions	
Prescriber Restrictions	CS: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	INITIAL AND RENEWAL: 12 MONTHS.
Other Criteria	CS: INITIAL: HYPERCORTISOLISM IS NOT A RESULT OF CHRONIC GLUCOCORTICOIDS. RENEWAL: 1) CONTINUES TO HAVE IMPROVEMENT OF GLUCOSE TOLERANCE OR STABLE GLUCOSE TOLERANCE (E.G., REDUCED A1C, IMPROVED FASTING GLUCOSE, ETC.), 2) CONTINUES TO HAVE TOLERABILITY TO THERAPY, AND 3) CONTINUES TO NOT BE A CANDIDATE FOR SURGICAL TREATMENT OR HAS FAILED SURGERY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MIGALASTAT

Products Affected

• GALAFOLD

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	FABRY DISEASE: INITIAL SYMPTOMATIC OR EVIDENCE OF INJURY FROM GL-3 TO THE KIDNEY, HEART, OR CENTRAL NERVOUS SYSTEM RECOGNIZED BY LABORATORY, HISTOLOGICAL, OR IMAGING FINDINGS.
Age Restrictions	
Prescriber Restrictions	FABRY DISEASE: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEPHROLOGIST, CARDIOLOGIST, OR SPECIALIST IN GENETICS OR INHERITED METABOLIC DISORDERS.
Coverage Duration	INITIAL: 6 MOS. RENEWAL: 12 MOS.
Other Criteria	FABRY DISEASE: INITIAL: NOT CONCURRENTLY USING ENZYME REPLACEMENT THERAPY (I.E. FABRAZYME). RENEWAL: 1) IMPROVEMENT OR STABILIZATION, AND 2) NOT CONCURRENTLY USING ENZYME REPLACEMENT THERAPY (I.E. FABRAZYME).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MIGLUSTAT

Products Affected

- miglustat
- yargesa

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MILTEFOSINE

Products Affected

• IMPAVIDO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MOBOCERTINIB

Products Affected

• EXKIVITY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MOMELOTINIB

Products Affected

• OJJAARA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

MOSUNETUZUMAB-AXGB

Products Affected

• LUNSUMIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	RELAPSED OR REFRACTORY FOLLICULAR LYMPHOMA: INITIAL: 6 MONTHS. RENEWAL: 7 MONTHS.
Other Criteria	RELAPSED OR REFRACTORY FOLLICULAR LYMPHOMA: RENEWAL: 1) HAS ACHIEVED A PARTIAL RESPONSE TO TREATMENT, AND 2) HAS NOT PREVIOUSLY RECEIVED MORE THAN 17 CYCLES OF TREATMENT. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NAFARELIN

Products Affected

• SYNAREL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: ENDOMETRIOSIS: DIAGNOSIS IS CONFIRMED VIA SURGICAL OR DIRECT VISUALIZATION (E.G., PELVIC ULTRASOUND) OR HISTOPATHOLOGICAL CONFIRMATION (E.G., LAPAROSCOPY OR LAPAROTOMY) IN THE LAST 10 YEARS. CENTRAL PRECOCIOUS PUBERTY (CPP): FEMALES: ELEVATED LEVELS OF FOLLICLE- STIMULATING HORMONE (FSH) GREATER THAN 4.0 MIU/ML AND LUTEINIZING HORMONE (LH) LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS. MALES: ELEVATED LEVELS OF FSH GREATER THAN 5.0 MIU/ML AND LH LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS.
Age Restrictions	
Prescriber Restrictions	INITIAL: ENDOMETRIOSIS: PRESCRIBED BY OR IN CONSULTATION WITH AN OBSTETRICIAN/GYNECOLOGIST. CPP: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	ENDOMETRIOSIS: 6 MONTHS. CPP: INITIAL/RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: ENDOMETRIOSIS: 1) NO CONCURRENT USE WITH ANOTHER GNRH-MODULATING AGENT, 2) TRIAL OF OR CONTRAINDICATION TO NSAID AND PROGESTIN- CONTAINING PREPARATION, AND 3) HAS NOT RECEIVED A TOTAL OF 6 MONTHS OF TREATMENT PER LIFETIME. CPP: FEMALES: 1) YOUNGER THAN 8 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR BREAST DEVELOPMENT AND PUBIC HAIR GROWTH. MALES: 1) YOUNGER THAN 9 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR GENITAL DEVELOPMENT AND PUBIC HAIR GROWTH. RENEWAL: CPP: 1) TANNER STAGE 2 OR ABOVE FOR GENITAL DEVELOPMENT AND PUBIC HAIR GROWTH. RENEWAL: CPP: 1) TANNER STAGING AT INITIAL DIAGNOSIS HAS STABILIZED OR REGRESSED DURING THREE SEPARATE MEDICAL VISITS IN THE PREVIOUS YEAR, AND 2) HAS NOT REACHED ACTUAL AGE WHICH CORRESPONDS TO CURRENT PUBERTAL AGE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NARCOLEPSY AGENTS

Products Affected

- armodafinil
- modafinil oral tablet 100 mg, 200 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All Medically-accepted Indications.
Off Label Uses	
Part B Prerequisite	No

NATALIZUMAB

Products Affected

• TYSABRI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: CROHNS DISEASE (CD): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	MULTIPLE SCLEROSIS (MS): 12 MOS. CD: INITIAL: 6 MOS, RENEWAL: 12 MOS.
Other Criteria	INITIAL: MS: TRIAL OF TWO AGENTS INDICATED FOR THE TREATMENT OF MS. CD: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, STELARA, SKYRIZI, RINVOQ. RENEWAL: CD: 1) RECEIVED AT LEAST 12 MONTHS OF THERAPY WITH TYSABRI AND HAS NOT REQUIRED MORE THAN 3 MONTHS OF CORTICOSTEROID USE WITHIN THE PAST 12 MONTHS TO CONTROL CROHNS DISEASE WHILE ON TYSABRI, OR 2) HAS ONLY RECEIVED 6 MONTHS OF THERAPY WITH TYSABRI AND HAS TAPERED OFF CORTICOSTEROIDS DURING THE FIRST 24 WEEKS OF TYSABRI THERAPY. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NAXITAMAB-GQGK

Products Affected

• DANYELZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NEDOSIRAN

Products Affected

• RIVFLOZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NERATINIB

Products Affected

• NERLYNX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	EARLY-STAGE (STAGE I-III) BREAST CANCER: MEDICATION IS BEING REQUESTED WITHIN 2 YEARS OF COMPLETING THE LAST TRASTUZUMAB DOSE. ALL OTHER FDA APPROVED INDICATIONS ARE COVERED WITHOUT ADDITIONAL CRITERIA, EXCEPT THOSE CRITERIA IN THE FDA APPROVED LABEL.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NILOTINIB

Products Affected

• TASIGNA ORAL CAPSULE 150 MG, 200 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	PREVIOUSLY TREATED CML: MUTATIONAL ANALYSIS PRIOR TO INITIATION AND TASIGNA IS APPROPRIATE PER NCCN GUIDELINE TABLE FOR TREATMENT RECOMMENDATIONS BASED ON BCR-ABL1 MUTATION PROFILE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NINTEDANIB

Products Affected

• OFEV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: IDIOPATHIC PULMONARY FIBROSIS (IPF): 1) A USUAL INTERSTITIAL PNEUMONIA (UIP) PATTERN AS EVIDENCED BY HIGH-RESOLUTION COMPUTED TOMOGRAPHY (HRCT) ALONE OR VIA A COMBINATION OF SURGICAL LUNG BIOPSY AND HRCT, AND 2) BASELINE FORCED VITAL CAPACITY (FVC) AT LEAST 50% OF PREDICTED VALUE. SYSTEMIC SCLEROSIS- ASSOCIATED INTERSTITIAL LUNG DISEASE (SSC-ILD): 1) AT LEAST 10% FIBROSIS ON A CHEST HRCT, AND 2) BASELINE FVC AT LEAST 40% OF PREDICTED VALUE. CHRONIC FIBROSING INTERSTITIAL LUNG DISEASE WITH A PROGRESSIVE PHENOTYPE (PF-ILD): 1) AT LEAST 10% FIBROSIS ON A CHEST HRCT, AND 2) BASELINE FVC AT LEAST 45% OF PREDICTED VALUE.
Age Restrictions	
Prescriber Restrictions	INITIAL: IPF: PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST. SSC-ILD, PF-ILD: PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST OR RHEUMATOLOGIST.
Coverage Duration	INITIAL: SSC-ILD: 6 MOS. IPF, PF-ILD: 12 MOS. RENEWAL (ALL INDICATIONS): 12 MOS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: IPF: 1) DOES NOT HAVE OTHER KNOWN CAUSES OF INTERSTITIAL LUNG DISEASE (E.G., CONNECTIVE TISSUE DISEASE, DRUG TOXICITY, ASBESTOS OR BERYLLIUM EXPOSURE, HYPERSENSITIVITY PNEUMONITIS), AND 2) TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENT: ESBRIET (PIRFENIDONE). SSC-ILD: 1) DOES NOT HAVE OTHER KNOWN CAUSES OF INTERSTITIAL LUNG DISEASE (E.G., HEART FAILURE/FLUID OVERLOAD, DRUG- INDUCED LUNG TOXICITY, RECURRENT ASPIRATION), AND 2) TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENT: ACTEMRA SUBQ. PF-ILD: LUNG FUNCTION AND RESPIRATORY SYMPTOMS OR CHEST IMAGING HAVE WORSENED/PROGRESSED DESPITE TREATMENT WITH MEDICATIONS USED IN CLINICAL PRACTICE FOR ILD (NOT ATTRIBUTABLE TO COMORBIDITIES SUCH AS INFECTION, HEART FAILURE). RENEWAL: IPF, SSC-ILD, PF-ILD: CLINICAL MEANINGFUL IMPROVEMENT OR MAINTENANCE IN ANNUAL RATE OF DECLINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NIRAPARIB

Products Affected

- ZEJULA ORAL CAPSULE
- ZEJULA ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	RECURRENT EPITHELIAL OVARIAN, FALLOPIAN TUBE, OR PRIMARY PERITONEAL CANCER: 1) ZEJULA WILL BE USED AS MONOTHERAPY, AND 2) ZEJULA IS STARTED NO LATER THAN 8 WEEKS AFTER THE MOST RECENT PLATINUM-CONTAINING REGIMEN.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NIRAPARIB/ABIRATERONE

Products Affected

• AKEEGA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC CASTRATION-RESISTANT PROSTATE CANCER (MCRPC): 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NIROGACESTAT

Products Affected

 OGSIVEO ORAL TABLET 100 MG, 150 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NIRSEVIMAB-ALIP

Products Affected

• BEYFORTUS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 MONTHS
Other Criteria	RESPIRATORY SYNCYTIAL VIRUS (RSV) PREVENTION: 1) HAS NOT COMPLETED A COURSE OF SYNAGIS OR HAS NOT RECEIVED A DOSE OF SYNAGIS IN THE PAST 30 DAYS, AND WILL NOT RECEIVE FURTHER DOSES OF SYNAGIS WITHIN THE SAME RSV SEASON, AND 2) HAS NOT RECEIVED MORE THAN 2 DOSES OF BEYFORTUS PER LIFETIME, UNLESS HAS UNDERGONE OR WILL UNDERGO A CARDIAC SURGERY WITH CARDIOPULMONARY BYPASS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NITISINONE

Products Affected

- nitisinone
- ORFADIN ORAL SUSPENSION

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HEREDITARY TYROSINEMIA TYPE 1 (HT-1): INITIAL: DIAGNOSIS CONFIRMED BY ELEVATED URINARY OR PLASMA SUCCINYLACETONE LEVELS OR A MUTATION IN THE FUMARYLACETOACETATE HYDROLASE GENE. RENEWAL: URINARY OR PLASMA SUCCINYLACETONE LEVELS HAVE DECREASED FROM BASELINE WHILE ON TREATMENT WITH NITISINONE.
Age Restrictions	
Prescriber Restrictions	HT-1: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PRESCRIBER SPECIALIZING IN INHERITED METABOLIC DISEASES.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	HT-1: INITIAL: ORFADIN SUSPENSION: TRIAL OF OR CONTRAINDICATION TO PREFERRED NITISINONE TABLETS OR CAPSULES.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NIVOLUMAB

Products Affected

• OPDIVO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	UNRESECTABLE OR METASTATIC MELANOMA: NO CONCURRENT USE WITH TARGETED THERAPY (I.E., BRAF INHIBITORS, MEK INHIBITORS, AND NTRK INHIBITORS).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NIVOLUMAB-RELATLIMAB-RMBW

Products Affected

• OPDUALAG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

NOGAPENDEKIN ALFA

Products Affected

• ANKTIVA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	40 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OBETICHOLIC ACID

Products Affected

• OCALIVA

PA Criteria	Criteria Details
Exclusion	PRIMARY BILIARY CHOLANGITIS (PBC):
Criteria	INITIAL/RENEWAL: COMPLETE BILIARY OBSTRUCTION.
Required Medical	PBC: INITIAL: DIAGNOSIS CONFIRMED BY TWO OF THE
Information	FOLLOWING: 1) ALKALINE PHOSPHATASE LEVEL OF AT
	LEAST 1.5 TIMES THE UPPER LIMIT OF NORMAL, 2)
	PRESENCE OF ANTIMITOCHONDRIAL ANTIBODIES AT A
	TITER OF 1:40 OR HIGHER, OR 3) HISTOLOGIC EVIDENCE OF NON-SUPPURATIVE DESTRUCTIVE CHOLANGITIS AND
	DESTRUCTION OF INTERLOBULAR BILE DUCTS.
	DESTRUCTION OF INTERLOBULAR BILL DUCTS.
Age Restrictions	
Prescriber	PBC: INITIAL: PRESCRIBED BY OR IN CONSULTATION
Restrictions	WITH A GASTROENTEROLOGIST OR HEPATOLOGIST.
Coverage	INITIAL/RENEWAL: 12 MONTHS
Duration	
Other Criteria	PBC: INITIAL: USED IN COMBINATION WITH
	URSODEOXYCHOLIC ACID IN A PATIENT WITH AN
	INADEQUATE RESPONSE TO URSODEOXYCHOLIC ACID,
	OR AS MONOTHERAPY IN A PATIENT WHO IS UNABLE TO
	TOLERATE URSODEOXYCHOLIC ACID. RENEWAL:
	CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B	No
Prerequisite	

OCRELIZUMAB

Products Affected

• OCREVUS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	RELAPSING FORM OF MULTIPLE SCLEROSIS (MS): TRIAL OF TWO AGENTS INDICATED FOR THE TREATMENT OF RELAPSING FORMS OF MS. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OFATUMUMAB-SQ

Products Affected

• KESIMPTA PEN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OLANZAPINE/SAMIDORPHAN

Products Affected

• LYBALVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	SCHIZOPHRENIA/BIPOLAR I: PRESCRIBED BY OR IN CONSULTATION WITH A PSYCHIATRIST
Coverage Duration	12 MONTHS
Other Criteria	SCHIZOPHRENIA: (1) PATIENT IS AT HIGH RISK OF WEIGHT GAIN AND (2) TRIAL OF OR CONTRAINDICATION TO LATUDA OR ONE OF THE FOLLOWING ORAL ANTIPSYCHOTICS: RISPERIDONE, CLOZAPINE TABLET, OLANZAPINE, IMMEDIATE RELEASE QUETIAPINE FUMARATE, ZIPRASIDONE, ARIPIPRAZOLE. BIPOLAR I: (1) PATIENT IS AT HIGH RISK OF WEIGHT GAIN AND (2) TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING ORAL ANTIPSYCHOTICS: RISPERIDONE, OLANZAPINE, IMMEDIATE RELEASE QUETIAPINE FUMARATE, ZIPRASIDONE, ARIPIPRAZOLE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OLAPARIB

Products Affected

• LYNPARZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	RECURRENT EPITHELIAL OVARIAN, FALLOPIAN TUBE OR PRIMARY PERITONEAL CANCER: MEDICATION WILL BE USED AS MONOTHERAPY. METASTATIC CASTRATION- RESISTANT PROSTATE CANCER: 1) PREVIOUSLY RECEIVED A BILATERAL ORCHIECTOMY, OR 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG. ALL OTHER FDA APPROVED INDICATIONS ARE COVERED WITHOUT ADDITIONAL CRITERIA, EXCEPT THOSE CRITERIA IN THE FDA APPROVED LABEL.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OLUTASIDENIB

Products Affected

• REZLIDHIA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OMACETAXINE

Products Affected

• SYNRIBO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OMALIZUMAB

Products Affected

• XOLAIR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: ASTHMA: POSITIVE SKIN PRICK OR BLOOD TEST (E.G., ELISA, FEIA) TO A PERENNIAL AEROALLERGEN AND A BASELINE IGE SERUM LEVEL GREATER THAN OR EQUAL TO 30 IU/ML. FOOD ALLERGY: 1) IGE SERUM LEVEL OF AT LEAST 30 IU/ML, AND 2) ALLERGEN SPECIFIC IGE SERUM LEVEL OF AT LEAST 6 KUA/L TO AT LEAST ONE FOOD, OR POSITIVE SKIN PRICK TEST TO AT LEAST ONE FOOD, OR POSITIVE MEDICALLY MONITORED FOOD CHALLENGE TO AT LEAST ONE FOOD.
Age Restrictions	
Prescriber Restrictions	INITIAL AND RENEWAL: CHRONIC SPONTANEOUS URTICARIA (CSU): PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN ALLERGY OR PULMONARY MEDICINE, DERMATOLOGY OR IMMUNOLOGY. INITIAL: NASAL POLYPS: PRESCRIBED BY OR IN CONSULTATION WITH AN OTOLARYNGOLOGIST, ALLERGIST OR IMMUNOLOGIST. ASTHMA: PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN ALLERGY OR PULMONARY MEDICINE. FOOD ALLERGY: PRESCRIBED BY OR IN CONSULTATION WITH ALLERGIST OR IMMUNOLOGIST.
Coverage Duration	INITIAL/RENEWAL: ASTHMA 4 MO/12 MO, CSU 6 MO, NASAL POLYPS 6 MO/12 MO, FOOD ALLERGY 12 MO/24 MO

PA Criteria	Criteria Details
Other Criteria	INITIAL: CSU: TRIAL OF OR CONTRAINDICATION TO A
	MAXIMALLY TOLERATED DOSE OF AN H1 ANTI-
	HISTAMINE AND STILL EXPERIENCES HIVES ON MOST
	DAYS OF THE WEEK. NASAL POLYPS: 1) A 56 DAY TRIAL
	OF ONE TOPICAL NASAL CORTICOSTEROID, AND 2) TRIAL
	OF OR CONTRAINDICATION TO ONE PREFERRED AGENT:
	NUCALA, DUPIXENT. ASTHMA: 1) CONCURRENT
	THERAPY WITH A MEDIUM, HIGH-DOSE OR MAXIMALLY
	TOLERATED DOSE OF AN INHALED CORTICOSTEROID
	AND AT LEAST ONE OTHER MAINTENANCE
	MEDICATION, 2) ONE OF THE FOLLOWING: (A) PATIENT
	EXPERIENCED AT LEAST ONE ASTHMA EXACERBATION
	REQUIRING SYSTEMIC CORTICOSTEROID BURST
	LASTING 3 OR MORE DAYS WITHIN THE PAST 12 MONTHS
	OR AT LEAST ONE SERIOUS EXACERBATION REQUIRING
	HOSPITALIZATION OR ER VISIT WITHIN THE PAST 12
	MONTHS, OR (B) PATIENT HAS POOR SYMPTOM CONTROL
	DESPITE CURRENT THERAPY AS EVIDENCED BY AT
	LEAST THREE OF THE FOLLOWING WITHIN THE PAST 4
	WEEKS: DAYTIME ASTHMA SYMPTOMS MORE THAN
	TWICE/WEEK, ANY NIGHT WAKING DUE TO ASTHMA,
	SABA RELIEVER FOR SYMPTOMS MORE THAN
	TWICE/WEEK, ANY ACTIVITY LIMITATION DUE TO
	ASTHMA, AND 3) NOT CONCURRENTLY RECEIVING
	DUPIXENT OR ANTI-IL5 BIOLOGICS WHEN THESE ARE
	USED FOR THE TREATMENT OF ASTHMA. FOOD
	ALLERGY: 1) CONCURRENT USE WITH AN ACTIVE
	PRESCRIPTION FOR EPINEPHRINE AUTO-
	INJECTOR/INJECTION, AND 2) NO CONCURRENT USE
	WITH PEANUT-SPECIFIC IMMUNOTHERAPY. RENEWAL:
	CSU: DIAGNOSIS OF CSU. NASAL POLYPS: CLINICAL
	BENEFIT COMPARED TO BASELINE. ASTHMA: 1) NOT
	CONCURRENTLY RECEIVING DUPIXENT OR ANTI-IL5
	BIOLOGICS WHEN THESE ARE USED FOR THE

PA Criteria	Criteria Details
	TREATMENT OF ASTHMA, 2) CONTINUED USE OF ICS AND AT LEAST ONE OTHER MAINTENANCE MEDICATION, AND 3) CLINICAL RESPONSE AS EVIDENCED BY ONE OF THE FOLLOWING: (A) REDUCTION IN ASTHMA EXACERBATIONS FROM BASELINE, (B) DECREASED UTILIZATION OF RESCUE MEDICATIONS, (C) REDUCTION IN SEVERITY OR FREQUENCY OF ASTHMA-RELATED SYMPTOMS, OR (D) INCREASE IN PERCENT PREDICTED FEV1 FROM PRETREATMENT BASELINE. FOOD ALLERGY: 1) PERSISTENT IGE-MEDIATED FOOD ALLERGY, 2) CONCURRENT USE WITH AN ACTIVE PRESCRIPTION FOR EPINEPHRINE AUTO-INJECTOR/INJECTION, AND 3) NO CONCURRENT USE WITH PEANUT-SPECIFIC IMMUNOTHERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OPICAPONE

Products Affected

• ONGENTYS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	PARKINSONS DISEASE: 18 YEARS OF AGE OR OLDER
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OSIMERTINIB

Products Affected

• TAGRISSO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	EGFR EXON 19 DELETIONS OR EXON 21 L858R MUTATIONS NON-SMALL CELL LUNG CANCER (NSCLC) AND METASTATIC NSCLC WITH EGFR T790M MUTATION: NOT ON CONCURRENT THERAPY WITH AN EGFR TYROSINE KINASE-INHIBITOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

OXANDROLONE

Products Affected

• oxandrolone

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 MONTHS
Other Criteria	PROTEIN CATABOLISM, BONE PAIN: 1) MONITORED FOR PELIOSIS HEPATIS, LIVER CELL TUMORS, AND BLOOD LIPID CHANGES, 2) DOES NOT HAVE KNOWN OR SUSPECTED: CARCINOMA OF THE PROSTATE OR BREAST IN MALE PATIENTS, CARCINOMA OF THE BREAST IN FEMALES WITH HYPERCALCEMIA, NEPHROSIS (THE NEPHROTIC PHASE OF NEPHRITIS), OR HYPERCALCEMIA, AND 3) DOES NOT HAVE SEVERE HEPATIC DYSFUNCTION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PACRITINIB

Products Affected

• VONJO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	MYELOFIBROSIS: RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PALBOCICLIB

Products Affected

• IBRANCE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	ADVANCED OR METASTATIC BREAST CANCER: TRIAL OF OR CONTRAINDICATION TO ONE OF THE PREFERRED AGENTS, WHERE INDICATIONS ALIGN: KISQALI, VERZENIO.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PALIVIZUMAB

Products Affected

• SYNAGIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: GESTATIONAL AGE
Age Restrictions	INITIAL AND RENEWAL: LESS THAN 24 MONTHS OF AGE.
Prescriber Restrictions	
Coverage Duration	CRITERIA CONSISTENT WITH THE CDC RSV CENSUS REGIONAL TREND. SEE OTHER CRITERIA.
Other Criteria	CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT RECOMMENDATIONS FROM THE AMERICAN ACADEMY OF PEDIATRICS FOR PALIVIZUMAB PROPHYLAXIS FOR RESPIRATORY SYNCYTIAL VIRUS (RSV) INFECTIONS AND THE CENTERS FOR DISEASE CONTROL AND PREVENTION RSV CENSUS REGIONAL TREND. APPROVAL WILL BE FOR UP TO A MAXIMUM OF 5 MONTHS DEPENDENT UPON REMAINING LENGTH OF RESPIRATORY SYNCYTIAL VIRUS SEASON. RENEWAL: ADDITIONAL 1 MONTH OF TREATMENT FOR CARDIOPULMONARY BYPASS SURGERY DURING RSV PROPHYLAXIS SEASON.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PARATHYROID HORMONE

Products Affected

• NATPARA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	HYPOCALCEMIA SECONDARY TO HYPOPARATHYROIDISM: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	HYPOCALCEMIA SECONDARY TO HYPOPARATHYROIDISM: 1) TRIAL OF OR CONTRAINDICATION TO CALCITRIOL, 2) HYPOPARATHYROIDISM IS NOT DUE TO A CALCIUM SENSING RECEPTOR (CSR) MUTATION, AND 3) HYPOPARATHYROIDISM IS NOT CONSIDERED ACUTE POST-SURGICAL HYPOPARATHYROIDISM.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PASIREOTIDE DIASPARTATE

Products Affected

• SIGNIFOR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	CUSHINGS DISEASE (CD): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	CD: RENEWAL: 1) CONTINUED IMPROVEMENT OF CUSHINGS DISEASE, AND 2) MAINTAINED TOLERABILITY TO SIGNIFOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PAZOPANIB

Products Affected

• pazopanib

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	ADVANCED SOFT TISSUE SARCOMA (STS): NOT USED FOR ADIPOCYTIC STS OR GASTROINTESTINAL STROMAL TUMORS (GIST)
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGFILGRASTIM - APGF

Products Affected

• NYVEPRIA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGFILGRASTIM - CBQV

Products Affected

- UDENYCA
- UDENYCA AUTOINJECTOR
- UDENYCA ONBODY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY, ACUTE RADIATION EXPOSURE: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	NON MYELOID MALIGNANCY: 1) TRIAL OF OR CONTRAINDICATION TO NYVEPRIA, OR 2) BARRIER TO ACCESS (E.G., TRAVEL BARRIERS OR UNABLE TO RETURN TO CLINIC FOR INJECTIONS).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGFILGRASTIM - JMDB

Products Affected

• FULPHILA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	NON MYELOID MALIGNANCY: TRIAL OF OR CONTRAINDICATION TO NYVEPRIA
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGFILGRASTIM-BMEZ

Products Affected

• ZIEXTENZO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY, ACUTE RADIATION EXPOSURE: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	NON MYELOID MALIGNANCY: TRIAL OF OR CONTRAINDICATION TO NYVEPRIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGFILGRASTIM-FPGK

Products Affected

• STIMUFEND

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY, ACUTE RADIATION EXPOSURE: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	NON MYELOID MALIGNANCY: TRIAL OF OR CONTRAINDICATION TO NYVEPRIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGFILGRASTIM-NEULASTA ONPRO

Products Affected

• NEULASTA ONPRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY, ACUTE RADIATION EXPOSURE: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGFILGRASTIM-PBBK

Products Affected

• FYLNETRA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON MYELOID MALIGNANCY: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	NON MYELOID MALIGNANCY: TRIAL OF OR CONTRAINDICATION TO PREFERRED AGENT: NYVEPRIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGINTERFERON ALFA-2A

Products Affected

• PEGASYS

PA Criteria	Criteria Details
Exclusion Criteria	CHRONIC HEPATITIS C VIRUS INFECTION. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	HEPATITIS B: PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, OR PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (E.G., HEPATOLOGIST).
Coverage Duration	CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE.
Other Criteria	CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGUNIGALSIDASE ALFA-IWXJ

Products Affected

• ELFABRIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	FABRY DISEASE: INITIAL: 1) SYMPTOMATIC OR EVIDENCE OF INJURY TO THE KIDNEY, HEART, OR CENTRAL NERVOUS SYSTEM RECOGNIZED BY LABORATORY, HISTOLOGICAL, OR IMAGING FINDINGS, AND 2) ONE OF THE FOLLOWING: (A) FEMALES: GALACTOSIDASE ALPHA (GLA) GENE MUTATION VIA GENETIC TESTING, OR (B) MALES: ENZYME ASSAY INDICATING ALPHA GALACTOSIDASE A DEFICIENCY OR GLA GENE MUTATION VIA GENETIC TESTING.
Age Restrictions	
Prescriber Restrictions	FABRY DISEASE: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH NEPHROLOGIST, CARDIOLOGIST, OR SPECIALIST IN GENETICS OR INHERITED METABOLIC DISORDERS.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	FABRY DISEASE: INITIAL: NO CONCURRENT USE WITH ANOTHER FABRY DISEASE THERAPY. RENEWAL: 1) DEMONSTRATED IMPROVEMENT OR STABILIZATION, AND 2) NO CONCURRENT USE WITH ANOTHER FABRY DISEASE THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGVALIASE-PQPZ

Products Affected

• PALYNZIQ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	PHENYLKETONURIA (PKU): INITIAL: NOT ON CONCURRENT TREATMENT WITH KUVAN. RENEWAL: 1) CONTINUES TO BENEFIT FROM THE MEDICATION, AND 2) NOT ON CONCURRENT TREATMENT WITH KUVAN.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEGVISOMANT

Products Affected

• SOMAVERT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEMBROLIZUMAB

Products Affected

• KEYTRUDA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	UNRESECTABLE OR METASTATIC MELANOMA: NO CONCURRENT USE WITH TARGETED THERAPY (I.E., BRAF INHIBITORS, MEK INHIBITORS, AND NTRK INHIBITORS).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEMIGATINIB

Products Affected

• PEMAZYRE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	CHOLANGIOCARCINOMA, MYELOID/LYMPHOID NEOPLASMS: COMPREHENSIVE OPHTHALMOLOGICAL EXAMINATION, INCLUDING OPTICAL COHERENCE TOMOGRAPHY (OCT), WILL BE COMPLETED PRIOR TO INITIATION OF THERAPY AND AT THE RECOMMENDED SCHEDULED INTERVALS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PENICILLAMINE TABLET

Products Affected

• penicillamine oral tablet

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: CYSTINURIA: HAS NEPHROLITHIASIS AND ONE OF THE FOLLOWING: 1) STONE ANALYSIS SHOWING PRESENCE OF CYSTINE, 2) PRESENCE OF PATHOGNOMONIC HEXAGONAL CYSTINE CRYSTALS ON URINALYSIS, OR 3) FAMILY HISTORY OF CYSTINURIA AND POSITIVE CYANIDE-NITROPRUSSIDE SCREENING.
Age Restrictions	
Prescriber Restrictions	INITIAL: WILSONS DISEASE: PRESCRIBED BY OR IN CONSULTATION WITH A HEPATOLOGIST OR GASTROENTEROLOGIST. CYSTINURIA: PRESCRIBED BY OR IN CONSULTATION WITH A NEPHROLOGIST. RHEUMATOID ARTHRITIS (RA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage Duration	INITIAL: 12 MONTHS, RENEWAL: LIFETIME.

PA Criteria	Criteria Details
Other Criteria	INITIAL: WILSONS DISEASE: 1) LEIPZIG SCORE OF 4 OR GREATER. RA: 1) NO HISTORY OR OTHER EVIDENCE OF RENAL INSUFFICIENCY, AND 2) TRIAL OF OR CONTRAINDICATION TO 3 MONTHS OF TREATMENT WITH ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) - IF PATIENT TRIED METHOTREXATE, THEN TRIAL AT A DOSE GREATER THAN OR EQUAL TO 20 MG PER WEEK OR MAXIMALLY TOLERATED DOSE IS REQUIRED. RENEWAL: RA: 1) NO HISTORY OR OTHER EVIDENCE OF RENAL INSUFFICIENCY, AND 2) EXPERIENCED OR MAINTAINED IMPROVEMENT IN TENDER JOINT COUNT OR SWOLLEN JOINT COUNT COMPARED TO BASELINE. WILSONS DISEASE, CYSTINURIA: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PEXIDARTINIB

Products Affected

• TURALIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PIMAVANSERIN

Products Affected

• NUPLAZID

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	PSYCHOSIS IN PARKINSONS DISEASE (PD): INITIAL: 18 YEARS OR OLDER
Prescriber Restrictions	PSYCHOSIS IN PD: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST, GERIATRICIAN, OR A BEHAVIORAL HEALTH SPECIALIST (E.G., PSYCHIATRIST).
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	PSYCHOSIS IN PD: RENEWAL: IMPROVEMENT IN PSYCHOSIS SYMPTOMS FROM BASELINE AND DEMONSTRATES A CONTINUED NEED FOR TREATMENT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PIRFENIDONE

Products Affected

- pirfenidone oral capsule
- pirfenidone oral tablet 267 mg, 534 mg, 801 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	IDIOPATHIC PULMONARY FIBROSIS (IPF): INITIAL: 1) A USUAL INTERSTITIAL PNEUMONIA (UIP) PATTERN AS EVIDENCED BY HIGH-RESOLUTION COMPUTED TOMOGRAPHY (HRCT) ALONE OR VIA A COMBINATION OF SURGICAL LUNG BIOPSY AND HRCT, AND 2) PREDICTED FORCED VITAL CAPACITY (FVC) OF AT LEAST 50% AT BASELINE.
Age Restrictions	IPF: INITIAL: 18 YEARS OR OLDER.
Prescriber Restrictions	IPF: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	IPF: INITIAL: 1) DOES NOT HAVE OTHER KNOWN CAUSES OF INTERSTITIAL LUNG DISEASE (E.G., CONNECTIVE TISSUE DISEASE, DRUG TOXICITY, ASBESTOS OR BERYLLIUM EXPOSURE, HYPERSENSITIVITY PNEUMONITIS, SYSTEMIC SCLEROSIS, RHEUMATOID ARTHRITIS, RADIATION, SARCOIDOSIS, BRONCHIOLITIS OBLITERANS ORGANIZING PNEUMONIA, HUMAN IMMUNODEFICIENCY VIRUS (HIV) INFECTION, VIRAL HEPATITIS, OR CANCER). RENEWAL: CLINICAL MEANINGFUL IMPROVEMENT OR MAINTENANCE IN ANNUAL RATE OF DECLINE.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

PIRTOBRUTINIB

Products Affected

 JAYPIRCA ORAL TABLET 100 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

POMALIDOMIDE

Products Affected

• POMALYST

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PONATINIB

Products Affected

• ICLUSIG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	CML: MUTATIONAL ANALYSIS PRIOR TO INITIATION AND ICLUSIG IS APPROPRIATE PER NCCN GUIDELINE TABLE FOR TREATMENT RECOMMENDATIONS BASED ON BCR-ABL1 MUTATION PROFILE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

POSACONAZOLE

Products Affected

• posaconazole oral

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	OROPHARYNGEAL CANDIDIASIS (OPC): 3 MONTHS. PROPHYLAXIS: 6 MONTHS. TREATMENT: 12 WEEKS.
Other Criteria	POSACONAZOLE SUSPENSION ONLY: 1) OPC: TRIAL OF OR CONTRAINDICATION TO FLUCONAZOLE OR ITRACONAZOLE. 2) PROPHYLAXIS OF INVASIVE ASPERGILLUS AND CANDIDA INFECTION: INABILITY TO SWALLOW TABLETS. POSACONZOLE TABLETS ONLY: 1) TREATMENT OF INVASIVE ASPERGILLOSIS, 2) PROPHYLAXIS OF INVASIVE ASPERGILLUS AND CANDIDA INFECTION: AT HIGH RISK OF DEVELOPING THESE INFECTIONS DUE TO BEING SEVERELY IMMUNOCOMPROMISED. CONTINUATION OF THERAPY AFTER HOSPITAL DISCHARGE REQUIRES NO EXTRA CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

POSACONAZOLE-POWDERMIX

Products Affected

• NOXAFIL ORAL SUSP, DELAYED RELEASE FOR RECON

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 MONTHS
Other Criteria	PROPHYLAXIS OF INVASIVE ASPERGILLUS AND CANDIDA INFECTION: INABILITY TO SWALLOW TABLETS. CONTINUATION OF THERAPY AFTER HOSPITAL DISCHARGE REQUIRES NO EXTRA CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PRALSETINIB

Products Affected

• GAVRETO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PRAMLINTIDE

Products Affected

- SYMLINPEN 120
- SYMLINPEN 60

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	TYPE I OR TYPE II DIABETES: REQUIRING INSULIN OR CONTINUOUS INSULIN INFUSION (INSULIN PUMP) FOR GLYCEMIC CONTROL
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

PYRIMETHAMINE

Products Affected

• pyrimethamine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	TOXOPLASMOSIS: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN INFECTIOUS DISEASE SPECIALIST.
Coverage Duration	TOXOPLASMOSIS: INITIAL: 8 WEEKS, RENEWAL: 6 MOS.
Other Criteria	TOXOPLASMOSIS: RENEWAL: ONE OF THE FOLLOWING: (1) PERSISTENT CLINICAL DISEASE (HEADACHE, NEUROLOGICAL SYMPTOMS, OR FEVER) AND PERSISTENT RADIOGRAPHIC DISEASE (ONE OR MORE MASS LESIONS ON BRAIN IMAGING), OR (2) CD4 COUNT LESS THAN 200 CELLS/MM3 AND CURRENTLY TAKING AN ANTI-RETROVIRAL THERAPY IF HIV POSITIVE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

QUININE

Products Affected

• quinine sulfate

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

QUIZARTINIB

Products Affected

• VANFLYTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RAMUCIRUMAB

Products Affected

• CYRAMZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

REGORAFENIB

Products Affected

• STIVARGA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RELUGOLIX

Products Affected

• ORGOVYX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

REPOTRECTINIB

Products Affected

• AUGTYRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RESLIZUMAB

Products Affected

• CINQAIR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	ASTHMA: INITIAL: BLOOD EOSINOPHIL LEVEL GREATER THAN OR EQUAL TO 150 CELLS/MCL WITHIN THE PAST 12 MONTHS.
Age Restrictions	
Prescriber Restrictions	ASTHMA: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN ALLERGY OR PULMONARY MEDICINE.
Coverage Duration	ASTHMA: INITIAL: 4 MONTHS. RENEWAL: 12 MONTHS

PA Criteria	Criteria Details
Other Criteria	ASTHMA: INITIAL: 1) CONCURRENT THERAPY WITH A MEDIUM, HIGH-DOSE OR MAXIMALLY TOLERATED DOSE OF AN INHALED CORTICOSTEROID (ICS) AND ONE OTHER MAINTENANCE MEDICATION, AND 2) ONE ASTHMA EXACERBATION REQUIRING SYSTEMIC CORTICOSTEROID BURST LASTING 3 OR MORE DAYS WITHIN THE PAST 12 MONTHS, OR AT LEAST ONE SERIOUS EXACERBATION REQUIRING HOSPITALIZATION OR ER VISIT WITHIN THE PAST 12 MONTHS, OR POOR SYMPTOM CONTROL DESPITE CURRENT THERAPY AS EVIDENCED BY AT LEAST THREE OF THE FOLLOWING WITHIN THE PAST 4 WEEKS: DAYTIME ASTHMA SYMPTOMS MORE THAN TWICE/WEEK, ANY NIGHT WAKING DUE TO ASTHMA, SABA RELIEVER FOR SYMPTOMS MORE THAN TWICE/WEEK, ANY ACTIVITY LIMITATION DUE TO ASTHMA, 3) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: FASENRA, NUCALA, DUPIXENT, AND 4) NO CONCURRENT USE WITH XOLAIR, DUPIXENT OR OTHER ANTI-IL5 BIOLOGICS WHEN USED FOR ASTHMA. RENEWAL: 1) NO CONCURRENT USE WITH XOLAIR, DUPIXENT OR OTHER ANTI-IL5 BIOLOGICS WHEN USED FOR ASTHMA, 2) CONTINUED USE OF ICS AND ONE OTHER MAINTENANCE MEDICATION, AND 3) CLINICAL RESPONSE AS EVIDENCED BY: (A) REDUCTION IN ASTHMA EXACERBATIONS FROM BASELINE, (B) DECREASED UTILIZATION OF RESCUE MEDICATIONS, (C) REDUCTION IN SEVERITY OR FREQUENCY OF ASTHMA- RELATED SYMPTOMS, OR (D) INCREASE IN PERCENT PREDICTED FEVI FROM PRETREATMENT BASELINE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RETIFANLIMAB-DLWR

Products Affected

• ZYNYZ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RIBOCICLIB

Products Affected

 KISQALI ORAL TABLET 200 MG/DAY (200 MG X 1), 400 MG/DAY (200 MG X 2), 600 MG/DAY (200 MG X 3)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RIBOCICLIB-LETROZOLE

Products Affected

 KISQALI FEMARA CO-PACK ORAL TABLET 200 MG/DAY(200 MG X 1)-2.5 MG, 400 MG/DAY(200 MG X 2)-2.5 MG, 600 MG/DAY(200 MG X 3)-2.5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RIFAXIMIN

Products Affected

• XIFAXAN ORAL TABLET 200 MG, 550 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	TRAVELERS DIARRHEA/HE: 12 MOS. IBS-D: 8 WKS.
Other Criteria	RIFAXIMIN 550 MG TABLETS: HE: TRIAL OF OR CONTRAINDICATION TO LACTULOSE OR CONCURRENT LACTULOSE THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RIMEGEPANT

Products Affected

• NURTEC ODT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: ACUTE MIGRAINE TREATMENT: TRIAL OF OR CONTRAINDICATION TO ONE TRIPTAN (E.G., SUMATRIPTAN, RIZATRIPTAN). EPISODIC MIGRAINE PREVENTION: 1) NO CONCURRENT USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION, AND 2) TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING PREVENTIVE MIGRAINE TREATMENTS: DIVALPROEX SODIUM, TOPIRAMATE, PROPRANOLOL, TIMOLOL. RENEWAL: ACUTE MIGRAINE TREATMENT: 1) IMPROVEMENT FROM BASELINE IN A VALIDATED ACUTE TREATMENT PATIENT-REPORTED OUTCOME QUESTIONNAIRE, OR 2) THERAPY WORKS CONSISTENTLY IN MAJORITY OF MIGRAINE ATTACKS. EPISODIC MIGRAINE PREVENTION: 1) NO CONCURRENT USE WITH OTHER CGRP INHIBITORS FOR MIGRAINE PREVENTION, AND 2) REDUCTION IN MIGRAINE OR HEADACHE FREQUENCY, MIGRAINE SEVERITY, OR MIGRAINE DURATION WITH THERAPY.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

RIOCIGUAT

Products Affected

• ADEMPAS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PULMONARY ARTERIAL HYPERTENSION (PAH): DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS. PERSISTENT/RECURRENT CHRONIC THROMBOEMBOLIC PULMONARY HYPERTENSION (CTEPH) (WHO GROUP 4): WHO FUNCTIONAL CLASS II-IV SYMPTOMS.
Age Restrictions	
Prescriber Restrictions	INITIAL: PAH, CTEPH: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: PAH: NOT CONCURRENTLY TAKING NITRATES, NITRIC OXIDE DONORS, PHOSPHODIESTERASE INHIBITORS, OR NON-SPECIFIC PDE INHIBITORS. CTEPH: 1) NOT CONCURRENTLY TAKING NITRATES, NITRIC OXIDE DONORS, PHOSPHODIESTERASE INHIBITORS, OR NON-SPECIFIC PDE INHIBITORS, AND 2) NOT A CANDIDATE FOR SURGERY OR HAS INOPERABLE CTEPH OR HAS PERSISTENT OR RECURRENT DISEASE AFTER SURGICAL TREATMENT. RENEWAL: PAH, CTEPH: NOT CONCURRENTLY TAKING NITRATES, NITRIC OXIDE DONORS, PHOSPHODIESTERASE INHIBITORS, OR NON- SPECIFIC PDE INHIBITORS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RIPRETINIB

Products Affected

• QINLOCK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RISANKIZUMAB-RZAA

Products Affected

• SKYRIZI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PLAQUE PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, FACE OR GENITAL AREA.
Age Restrictions	
Prescriber Restrictions	INITIAL: PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR DERMATOLOGIST. CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	INITIAL: PSO: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. PSA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG). CD, UC: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. RENEWAL: PSO, PSA: CONTINUES TO BENEFIT FROM THE MEDICATION.

PA Criteria	Criteria Details
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RISDIPLAM

Products Affected

• EVRYSDI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	SPINAL MUSCULAR ATROPHY (SMA): INITIAL: GENE MUTATION ANALYSIS INDICATING MUTATIONS OR DELETIONS OF BOTH ALLELES OF THE SURVIVAL MOTOR NEURON 1 (SMN1) GENE. FOR PRESYMPTOMATIC PATIENTS: UP TO THREE COPIES OF SURVIVAL MOTOR NEURON 2 (SMN2) BASED ON NEWBORN SCREENING.
Age Restrictions	
Prescriber Restrictions	SMA: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROMUSCULAR SPECIALIST OR SMA SPECIALIST AT A SMA SPECIALTY CENTER.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	SMA: INITIAL: FOR SYMPTOMATIC PATIENTS: (1) BASELINE MOTOR FUNCTION ASSESSMENT BY A NEUROMUSCULAR SPECIALIST OR SMA SPECIALIST, AND (2) IF PATIENT RECEIVED GENE THERAPY, THE PATIENT HAD LESS THAN EXPECTED CLINICAL BENEFIT. RENEWAL: IMPROVED, MAINTAINED, OR DEMONSTRATED LESS THAN EXPECTED DECLINE IN: (1) MOTOR FUNCTION ASSESSMENTS COMPARED TO BASELINE, OR (2) OTHER MUSCLE FUNCTION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RITUXIMAB AND HYALURONIDASE HUMAN-SQ

Products Affected

• RITUXAN HYCELA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	FOLLICULAR LYMPHOMA (FL), DIFFUSE LARGE B-CELL LYMPHOMA (DLBCL), CHRONIC LYMPHOCYTIC LEUKEMIA (CLL): HAS RECEIVED OR WILL RECEIVE AT LEAST ONE FULL DOSE OF A RITUXIMAB PRODUCT BY INTRAVENOUS INFUSION PRIOR TO INITIATION OF RITUXIMAB AND HYALURONIDASE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RITUXIMAB-ABBS

Products Affected

• TRUXIMA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. NHL, CLL: PRESCRIBED BY OR IN CONSULTATION WITH AN ONCOLOGIST.
Coverage Duration	RA: INITIAL: 6 MO, RENEWAL: 12 MO. NHL, GPA, MPA: 12 MO. CLL: 6 MO.
Other Criteria	RA: INITIAL: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RITUXIMAB-ARRX

Products Affected

• RIABNI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	RHEUMATOID ARTHRITIS (RA): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. NHL, CLL: PRESCRIBED BY OR IN CONSULTATION WITH AN ONCOLOGIST.
Coverage Duration	RA: INITIAL: 6 MO, RENEWAL: 12 MO. NHL, WG, MPA: 12 MO. CLL: 6 MO.
Other Criteria	RA: INITIAL: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RITUXIMAB-PVVR

Products Affected

• RUXIENCE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	RA: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. NHL, CLL: PRESCRIBED BY OR IN CONSULTATION WITH AN ONCOLOGIST.
Coverage Duration	NHL, GPA, MPA: 12 MONTHS. CLL: 6 MONTHS. RA: INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	RA: INITIAL: 1) TRIAL OF OR CONTRAINDICATION TO TWO PREFERRED AGENTS: HUMIRA, ENBREL, RINVOQ, XELJANZ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ROMIPLOSTIM

Products Affected

• NPLATE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	IMMUNE THROMBOCYTOPENIA (ITP): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR IMMUNOLOGIST.
Coverage Duration	ITP: INITIAL: 4 MO, RENEWAL: 12 MO. HEMATOPOIETIC SYNDROME OF ACUTE RADIATION SYNDROME: 12 MO.
Other Criteria	ITP: INITIAL: TRIAL OF OR CONTRAINDICATION TO CORTICOSTEROIDS, IMMUNOGLOBULINS, OR AN INSUFFICIENT RESPONSE TO SPLENECTOMY. RENEWAL: CLINICAL RESPONSE TO THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ROPEGINTERFERON ALFA-2B-NJFT

Products Affected

• BESREMI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RUCAPARIB

Products Affected

• RUBRACA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	METASTATIC CASTRATION-RESISTANT PROSTATE CANCER: ONE OF THE FOLLOWING: 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

RUXOLITINIB

Products Affected

• JAKAFI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	MYELOFIBROSIS: INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS. POLYCYTHEMIA VERA, GVHD: 12 MONTHS.
Other Criteria	MYELOFIBROSIS: RENEWAL: PATIENT CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SAFINAMIDE

Products Affected

• XADAGO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SAPROPTERIN

Products Affected

- javygtor oral tablet, soluble sapropterin oral tablet, soluble

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 1 MONTH, RENEWAL 12 MONTHS.
Other Criteria	HYPERPHENYLALANINEMIA (HPA): INITIAL: NOT CONCURRENTLY USING PALYNZIQ. RENEWAL: 1) CONTINUES TO BENEFIT FROM TREATMENT, AND 2) NOT CONCURRENTLY USING PALYNZIQ.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SARILUMAB

Products Affected

• KEVZARA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	RHEUMATOID ARTHRITIS (RA), POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage Duration	RA, PJIA: INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS. POLYMYALGIA RHEUMATICA (PMR): 12 MONTHS.
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PJIA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ IR, RINVOQ. RENEWAL: RA, PJIA: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SATRALIZUMAB-MWGE

Products Affected

• ENSPRYNG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NEUROMYELITIS OPTICA SPECTRUM DISORDER (NMOSD): INITIAL: PRESCRIBED BY AN OPHTHALMOLOGIST OR PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	NMOSD: INITIAL: 1) ONE OF THE FOLLOWING CORE CLINICAL CHARACTERISTIC: (A) OPTIC NEURITIS, (B) ACUTE MYELITIS, (C) AREA POSTREMA SYNDROME, (D) ACUTE BRAINSTEM SYNDROME, (E) SYMPTOMATIC NARCOLEPSY OR ACUTE DIENCEPHALIC CLINICAL SYNDROME WITH NMOSD-TYPICAL DIENCEPHALIC MRI LESIONS, OR (F) SYMPTOMATIC CEREBRAL SYNDROME WITH NMOSD-TYPICAL BRAIN LESIONS, AND 2) NO CONCURRENT USE WITH RITUXIMAB, INEBILIZUMAB, OR ECULIZUMAB. RENEWAL: 1) REDUCTION IN RELAPSE FREQUENCY FROM BASELINE, AND 2) NO CONCURRENT USE WITH RITUXIMAB, INEBILIZUMAB, OR ECULIZUMAB.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SEBELIPASE ALFA

Products Affected

• KANUMA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	LYSOSOMAL ACID LIPASE (LAL) DEFICIENCY: INITIAL: DIAGNOSIS CONFIRMED BY 1) PRESENCE OF CLINICAL FEATURES (E.G., HEPATOMEGALY, ELEVATED SERUM TRANSAMINASES, DYSLIPIDEMIA, SPLENOMEGALY), AND 2) ONE OF THE FOLLOWING: (A) BLOOD TEST INDICATING LOW OR ABSENT LEVELS OF LAL ENZYME ACTIVITY, (B) DRIED BLOOD SPOT TEST INDICATING LOW OR ABSENT LAL ENZYME ACTIVITY, OR (C) GENETIC TEST INDICATING THE BI-ALLELIC PRESENCE OF ALTERED LIPA GENE(S).
Age Restrictions	
Prescriber Restrictions	LAL DEFICIENCY: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST, HEPATOLOGIST, GASTROENTEROLOGIST, MEDICAL GENETICIST, LIPIDOLOGIST, OR METABOLIC SPECIALIST.
Coverage Duration	LAL DEFICIENCY: INITIAL: 6 OR 12 MONTHS, SEE OTHER CRITERIA. RENEWAL: 12 MONTHS

PA Criteria	Criteria Details
Other Criteria	LAL DEFICIENCY: INITIAL: RAPIDLY PROGRESSIVE LAL DEFICIENCY PRESENTING WITHIN THE FIRST 6 MONTHS OF LIFE: 12 MONTHS. LAL DEFICIENCY PRESENTING AFTER THE FIRST 6 MONTHS OF LIFE AND NOT CONSIDERED RAPIDLY PROGRESSIVE: 6 MONTHS. RENEWAL: LAL DEFICIENCY PRESENTING AFTER THE FIRST 6 MONTHS OF LIFE AND NOT CONSIDERED RAPIDLY PROGRESSIVE: IMPROVEMENT IN ONE OF THE FOLLOWING CLINICAL PARAMETERS DURING THE PAST 6 MONTHS: (1) A RELATIVE REDUCTION FROM BASELINE IN LDL-C, NON-HDL-C, OR TRIGLYCERIDES, (2) NORMALIZATION OF ASPARTATE AMINOTRANSFERASE (AST) BASED ON AGE- AND GENDER-SPECIFIC NORMAL RANGES, OR (3) DECREASE IN LIVER FAT CONTENT COMPARED TO BASELINE ASSESSED BY ABDOMINAL IMAGING.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SECUKINUMAB

Products Affected

• COSENTYX (2 SYRINGES)

GES) • COSENTYX UNOREADY PEN

COSENTYX PEN (2 PENS)
COSENTYX SUBCUTANEOUS SYRINGE 75 MG/0.5 ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, GENITAL AREA, OR FACE. NON-RADIOGRAPHIC AXIAL SPONDYLOARTHRITIS (NR-AXSPA): 1) C-REACTIVE PROTEIN (CRP) LEVELS ABOVE THE UPPER LIMIT OF NORMAL, OR 2) SACROILLITIS ON MAGNETIC RESONANCE IMAGING (MRI).
Age Restrictions	
Prescriber Restrictions	INITIAL: PSO, HIDRADENITIS SUPPURATIVA (HS): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR A DERMATOLOGIST. ANKYLOSING SPONDYLITIS (AS), NR-AXSPA, ENTHESITIS- RELATED ARTHRITIS (ERA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage Duration	INITIAL: HS: 4 MONTHS, ALL OTHER INDICATIONS: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: PSO: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. PSA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE-MODIFYING ANTI- RHEUMATIC DRUG). NR-AXSPA: TRIAL OF OR CONTRAINDICATION TO AN NSAID (NON-STEROIDAL ANTI-INFLAMMATORY DRUG). ERA: TRIAL OF OR CONTRAINDICATION TO ONE NSAID, SULFASALAZINE, OR METHOTREXATE. HS: NO CONCURRENT USE WITH OTHER SYSTEMIC BIOLOGICS FOR HS OR OTHER IL-17 INHIBITORS FOR ANY INDICATION. RENEWAL: PSO, PSA, AS, NR-AXSPA, ERA: CONTINUES TO BENEFIT FROM THE MEDICATION. HS: 1) NO CONCURRENT USE WITH OTHER SYSTEMIC BIOLOGICS FOR HS OR OTHER IL-17 INHIBITORS FOR ANY INDICATION, AND 2) CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SECUKINUMAB IV

Products Affected

• COSENTYX INTRAVENOUS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: NON-RADIOGRAPHIC AXIAL SPONDYLOARTHRITIS (NR-AXSPA): 1) C-REACTIVE PROTEIN (CRP) LEVELS ABOVE THE UPPER LIMIT OF NORMAL, OR 2) SACROILIITIS ON MAGNETIC RESONANCE IMAGING (MRI).
Age Restrictions	
Prescriber Restrictions	INITIAL: PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR A DERMATOLOGIST. ANKYLOSING SPONDYLITIS (AS), NR- AXSPA: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	INITIAL: PSA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE-MODIFYING ANTI-RHEUMATIC DRUG). NR-AXSPA: TRIAL OF OR CONTRAINDICATION TO AN NSAID (NON-STEROIDAL ANTI-INFLAMMATORY DRUG). RENEWAL: PSA, AS, NR-AXSPA: CONTINUES TO BENEFIT FROM THE MEDICATION. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SELEXIPAG

Products Affected

- UPTRAVI INTRAVENOUS
- UPTRAVI ORAL TABLET 1,000 MCG, 1,200 MCG, 1,400 MCG, 1,600 MCG, 200 MCG, 400 MCG, 600 MCG, 800 MCG
- UPTRAVI ORAL TABLETS, DOSE PACK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	PAH: INITIAL: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING AGENTS FROM DIFFERENT DRUG CLASSES: 1) FORMULARY VERSION OF AN ORAL ENDOTHELIN RECEPTOR ANTAGONIST, 2) FORMULARY VERSION OF AN ORAL PHOSPHODIESTERASE TYPE-5 INHIBITOR, OR 3) FORMULARY VERSION OF AN ORAL CGMP STIMULATOR.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SELINEXOR

Products Affected

 XPOVIO ORAL TABLET 100 MG/WEEK (50 MG X 2), 40 MG/WEEK (40 MG X 1), 40MG TWICE WEEK (40 MG X 2), 60 MG/WEEK (60 MG X 1), 60MG TWICE WEEK (120 MG/WEEK), 80 MG/WEEK (40 MG X 2), 80MG TWICE WEEK (160 MG/WEEK)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SELPERCATINIB

Products Affected

• RETEVMO ORAL CAPSULE 40 MG, 80 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SELUMETINIB

Products Affected

 KOSELUGO ORAL CAPSULE 10 MG, 25 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SILDENAFIL IV

Products Affected

• sildenafil (pulm.hypertension) intravenous

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	PAH: INITIAL/RENEWAL: NOT CONCURRENTLY OR INTERMITTENTLY TAKING ORAL ERECTILE DYSFUNCTION AGENTS (E.G. CIALIS, VIAGRA), ANY ORGANIC NITRATES IN ANY FORM, OR GUANYLATE CYCLASE STIMULATORS.
Indications	All Medically-accepted Indications.
Off Label Uses	
Part B Prerequisite	No

SILDENAFIL TABLET

Products Affected

• sildenafil (pulm.hypertension) oral tablet

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: AGES 18 YEARS OR OLDER: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS. AGES 1 TO 17 YEARS: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PAP GREATER THAN 20 MMHG, 2) PCWP OF 15 MMHG OR LESS, AND 3) PVR OF 3 WOOD UNITS OR GREATER.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	PAH: INITIAL/RENEWAL: NOT CONCURRENTLY OR INTERMITTENTLY TAKING ORAL ERECTILE DYSFUNCTION AGENTS (E.G. CIALIS, VIAGRA), ANY ORGANIC NITRATES IN ANY FORM, OR GUANYLATE CYCLASE STIMULATORS. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All Medically-accepted Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

SIPONIMOD

Products Affected

• MAYZENT ORAL TABLET 0.25 MG, 1 • MAYZENT STARTER(FOR 2MG MG, 2 MG

MAINT)

• MAYZENT STARTER(FOR 1MG MAINT)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	MULTIPLE SCLEROSIS: RENEWAL: DEMONSTRATION OF CLINICAL BENEFIT COMPARED TO PRE-TREATMENT BASELINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SIROLIMUS PROTEIN-BOUND

Products Affected

• FYARRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SODIUM OXYBATE-XYREM

Products Affected

• sodium oxybate

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: ALL INDICATIONS: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST, PSYCHIATRIST, OR SPECIALIST IN SLEEP MEDICINE
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS
Other Criteria	INITIAL: EXCESSIVE DAYTIME SLEEPINESS (EDS) IN NARCOLEPSY: 1) NOT CURRENTLY TAKING A SEDATIVE HYPNOTIC AGENT, 2) FOR PATIENTS 18 YEARS OR OLDER: TRIAL OF OR CONTRAINDICATION TO A FORMULARY VERSION OF MODAFINIL, ARMODAFINIL, OR SOLRIAMFETOL AND ONE OTHER GENERIC STIMULANT INDICATED FOR EDS IN NARCOLEPSY, AND 3) FOR PATIENTS 7 TO 17 YEARS OF AGE: TRIAL OF OR CONTRAINDICATION TO ONE GENERIC STIMULANT INDICATED FOR EDS IN NARCOLEPSY. CATAPLEXY IN NARCOLEPSY: NOT CURRENTLY TAKING A SEDATIVE HYPNOTIC AGENT. RENEWAL (ALL INDICATIONS): 1) SUSTAINED IMPROVEMENT OF SYMPTOMS COMPARED TO BASELINE, AND 2) NOT CURRENTLY TAKING A SEDATIVE HYPNOTIC AGENT.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

SODIUM PHENYLBUTYRATE TABLETS

Products Affected

• sodium phenylbutyrate oral tablet

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	UREA CYCLE DISORDER (UCD): INITIAL: UCD IS CONFIRMED VIA ENZYMATIC, BIOCHEMICAL OR GENETIC TESTING.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	UCD: RENEWAL: CLINICAL BENEFIT FROM BASELINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SOFOSBUVIR/VELPATASVIR

Products Affected

- EPCLUSA ORAL PELLETS IN PACKET 150-37.5 MG, 200-50 MG
- EPCLUSA ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HCV RNA LEVEL WITHIN PAST 6 MONTHS.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE.
Other Criteria	1) CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE, 2) NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING MEDICATIONS: AMIODARONE, CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, OXCARBAZEPINE, RIFAMPIN, RIFABUTIN, RIFAPENTINE, HIV REGIMEN THAT CONTAINS EFAVIRENZ, ROSUVASTATIN AT DOSES ABOVE 10MG, TIPRANAVIR/RITONAVIR, TOPOTECAN, SOVALDI (AS A SINGLE AGENT), HARVONI, ZEPATIER, MAVYRET, OR VOSEVI, AND 3) PATIENTS WITH DECOMPENSATED CIRRHOSIS REQUIRE CONCURRENT RIBAVIRIN UNLESS RIBAVIRIN INELIGIBLE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SOFOSBUVIR/VELPATASVIR/VOXILAPREVIR

Products Affected

• VOSEVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HCV RNA LEVEL WITHIN PAST 6 MONTHS
Age Restrictions	
Prescriber Restrictions	
Coverage	CRITERIA WILL BE APPLIED CONSISTENT WITH
Duration	CURRENT AASLD/IDSA GUIDANCE.
Other Criteria	1) CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE, 2) NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING MEDICATIONS: AMIODARONE, CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, OXCARBAZEPINE, RIFAMPIN, RIFABUTIN, RIFAPENTINE, CYCLOSPORINE, PITAVASTATIN, PRAVASTATIN (DOSES ABOVE 40MG), ROSUVASTATIN, METHOTREXATE, MITOXANTRONE, IMATINIB, IRINOTECAN, LAPATINIB, SULFASALAZINE, TOPOTECAN, OR HIV REGIMEN THAT CONTAINS EFAVIRENZ, ATAZANAVIR, LOPINAVIR, TIPRANAVIR/RITONAVIR, SOVALDI (AS A SINGLE AGENT), EPCLUSA, HARVONI, ZEPATIER, OR MAVYRET, AND 3) DOES NOT HAVE MODERATE OR SEVERE HEPATIC IMPAIRMENT (CHILD-PUGH B OR C).
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

SOLRIAMFETOL

Products Affected

• SUNOSI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: EXCESSIVE DAYTIME SLEEPINESS (EDS) IN NARCOLEPSY: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST, PSYCHIATRIST, OR SPECIALIST IN SLEEP MEDICINE.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: EDS IN NARCOLEPSY: TRIED THE FORMULARY VERSION OF MODAFINIL OR ARMODAFINIL, AND ONE OTHER GENERIC STIMULANT INDICATED FOR EDS IN NARCOLEPSY. EDS IN OBSTRUCTIVE SLEEP APNEA (OSA): TRIED THE FORMULARY VERSION OF MODAFINIL OR ARMODAFINIL. RENEWAL: EDS IN NARCOLEPSY OR OSA: SUSTAINED IMPROVEMENT OF SYMPTOMS COMPARED TO BASELINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SOMATROPIN - NORDITROPIN

Products Affected

• NORDITROPIN FLEXPRO

PA Criteria	Criteria Details
Exclusion Criteria	INITIAL/RENEWAL: ATHLETIC ENHANCEMENT, ANTI- AGING PURPOSES.
Required Medical Information	INITIAL: PEDIATRIC GROWTH HORMONE DEFICIENCY (GHD), IDIOPATHIC SHORT STATURE (ISS), SMALL FOR GESTATIONAL AGE (SGA), TURNER SYNDROME (TS), NOONAN SYNDROME: HEIGHT AT LEAST 2 STANDARD DEVIATIONS (SD) BELOW THE MEAN HEIGHT FOR NORMAL CHILDREN OF THE SAME AGE AND GENDER. PRADER WILLI SYNDROME (PWS): CONFIRMED GENETIC DIAGNOSIS.
Age Restrictions	
Prescriber Restrictions	INITIAL/RENEWAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: ADULT GHD: GROWTH HORMONE DEFICIENCY ALONE OR ASSOCIATED WITH MULTIPLE HORMONE DEFICIENCIES (HYPOPITUITARISM), AS A RESULT OF PITUITARY DISEASES, HYPOTHALAMIC DISEASE, SURGERY, RADIATION THERAPY, TRAUMA, OR CONTINUATION OF THERAPY FROM CHILDHOOD ONSET GROWTH HORMONE DEFICIENCY. PEDIATRIC GHD, ISS, SGA, TS, NOONAN SYNDROME: OPEN EPIPHYSES AS CONFIRMED BY RADIOGRAPH OF THE WRIST AND HAND. RENEWAL: PEDIATRIC GHD: 1) IMPROVEMENT WHILE ON THERAPY (I.E., INCREASED HEIGHT OR INCREASED GROWTH VELOCITY), AND 2) OPEN EPIPHYSES AS CONFIRMED BY RADIOGRAPH OF THE WRIST AND HAND OR PATIENT HAS NOT COMPLETED PREPUBERTALGROWTH. ISS, SGA, TS, NOONAN SYNDROME: 1) IMPROVEMENT WHILE ON THERAPY (I.E., INCREASED HEIGHT OR INCREASED GROWTH VELOCITY), AND 2) OPEN EPIPHYSES AS CONFIRMED BY RADIOGRAPH OF THE WRIST AND HAND. PWS: IMPROVEMENT IN BODY COMPOSITION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SOMATROPIN - SEROSTIM

Products Affected

• SEROSTIM SUBCUTANEOUS RECON SOLN 4 MG, 5 MG, 6 MG

PA Criteria	Criteria Details
Exclusion Criteria	INITIAL/RENEWAL: ATHLETIC ENHANCEMENT, ANTI- AGING PURPOSES
Required Medical Information	INITIAL: HIV/WASTING: MEETS ONE OF THE FOLLOWING CRITERIA FOR WEIGHT LOSS: 1) 10% UNINTENTIONAL WEIGHT LOSS OVER 12 MONTHS, 2) 7.5% UNINTENTIONAL WEIGHT LOSS OVER 6 MONTHS, 3) 5% BODY CELL MASS (BCM) LOSS WITHIN 6 MONTHS, 4) BODY CELL MASS (BCM) LESS THAN 35% (MEN) OF TOTAL BODY WEIGHT AND BODY MASS INDEX (BMI) LESS THAN 27 KG PER METER SQUARED, 5) BCM LESS THAN 23% (WOMEN) OF TOTAL BODY WEIGHT AND A BODY MASS INDEX (BMI) LESS THAN 27 KG PER METER SQUARED, OR 6) BMI LESS THAN 18.5 KG PER METER SQUARED.
Age Restrictions	
Prescriber Restrictions	HIV/WASTING: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST, NUTRITIONAL SUPPORT SPECIALIST, OR INFECTIOUS DISEASE SPECIALIST.
Coverage Duration	INITIAL/RENEWAL: 3 MONTHS.
Other Criteria	HIV/WASTING: INITIAL: 1) CURRENTLY ON HIV ANTIRETROVIRAL THERAPY, AND 2) INADEQUATE RESPONSE TO ONE PREVIOUS THERAPY (E.G., MEGACE, APPETITE STIMULANTS, ANABOLIC STEROIDS). RENEWAL: 1) CURRENTLY ON HIV ANTIRETROVIRAL THERAPY, AND 2) CLINICAL BENEFIT IN MUSCLE MASS AND WEIGHT.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

SONIDEGIB

Products Affected

• ODOMZO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	LOCALLY ADVANCED BASAL CELL CARCINOMA (BCC): BASELINE SERUM CREATINE KINASE (CK) AND SERUM CREATININE LEVELS
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SORAFENIB

Products Affected

• sorafenib

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SOTATERCEPT-CSRK

Products Affected

• WINREVAIR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	PAH: INITIAL: 1) ON BACKGROUND PAH THERAPY (FOR AT LEAST 3 MONTHS) WITH AT LEAST TWO OF THE FOLLOWING AGENTS FROM DIFFERENT DRUG CLASSES: A) ORAL ENDOTHELIN RECEPTOR ANTAGONIST, B) ORAL PHOSPHODIESTERASE TYPE-5 INHIBITOR FOR PAH, C) ORAL CGMP STIMULATOR, D) IV/SQ PROSTACYCLIN, OR 2) ON ONE AGENT FROM ONE OF THE ABOVE DRUG CLASSES, AND HAS A CONTRAINDICATION OR INTOLERANCE TO ALL OF THE OTHER DRUG CLASSES.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

SOTORASIB

Products Affected

• LUMAKRAS ORAL TABLET 120 MG, 320 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

STIRIPENTOL

Products Affected

- DIACOMIT ORAL CAPSULE 250 MG, 500 MG
- DIACOMIT ORAL POWDER IN PACKET 250 MG, 500 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	DRAVET SYNDROME: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

SUNITINIB

Products Affected

• *sunitinib malate*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	GASTROINTESTINAL STROMAL TUMORS (GIST): TRIAL OF OR CONTRAINDICATION TO IMATINIB (GLEEVEC).
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TADALAFIL - ADCIRCA, ALYQ

Products Affected

- alyq
- tadalafil (pulm. hypertension)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	PAH: INITIAL/RENEWAL: NOT CONCURRENTLY OR INTERMITTENTLY TAKING ORAL ERECTILE DYSFUNCTION AGENTS (E.G. CIALIS, VIAGRA), ANY ORGANIC NITRATES IN ANY FORM, OR GUANYLATE CYCLASE STIMULATORS. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All Medically-accepted Indications.
Off Label Uses	
Part B Prerequisite	No

TADALAFIL-CIALIS

Products Affected

• tadalafil oral tablet 2.5 mg, 5 mg

PA Criteria	Criteria Details
Exclusion Criteria	ERECTILE DYSFUNCTION WITHOUT DIAGNOSIS OF BENIGN PROSTATIC HYPERPLASIA (BPH).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	BPH: 1) TRIAL OF ONE ALPHA BLOCKER (E.G., DOXAZOSIN, TERAZOSIN, TAMSULOSIN, ALFUZOSIN), AND 2) TRIAL OF ONE 5-ALPHA-REDUCTASE INHIBITOR (E.G., FINASTERIDE, DUTASTERIDE). APPLIES TO 2.5MG AND 5MG STRENGTHS ONLY. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TALAZOPARIB

Products Affected

• TALZENNA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	ADVANCED OR METASTATIC BREAST CANCER: PATIENT HAS BEEN TREATED WITH CHEMOTHERAPY IN THE NEOADJUVANT, ADJUVANT, OR METASTATIC SETTING. PATIENTS WITH HORMONE RECEPTOR (HR)-POSITIVE BREAST CANCER MUST HAVE ADDITIONAL PRIOR TREATMENT WITH ENDOCRINE THERAPY OR BE CONSIDERED INAPPROPRIATE FOR ENDOCRINE THERAPY. METASTATIC CASTRATION-RESISTANT PROSTATE CANCER: 1) RECEIVED A BILATERAL ORCHIECTOMY, 2) CASTRATE LEVEL OF TESTOSTERONE (I.E., LESS THAN 50 NG/DL), OR 3) CONCURRENT USE WITH A GONADOTROPIN RELEASING HORMONE (GNRH) ANALOG.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TALIMOGENE

Products Affected

• IMLYGIC INJECTION SUSPENSION 10EXP6 (1 MILLION) PFU/ML

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	UNRESECTABLE MELANOMA: 1) IMLYGIC TO BE INJECTED INTO CUTANEOUS, SUBCUTANEOUS, AND/OR NODAL LESIONS THAT ARE VISIBLE, PALPABLE, OR DETECTABLE BY ULTRASOUND GUIDANCE, 2) NOT CURRENTLY RECEIVING IMMUNOSUPPRESSIVE THERAPY, 3) NO HISTORY OF PRIMARY OR ACQUIRED IMMUNODEFICIENT STATES, LEUKEMIA, LYMPHOMA, OR AIDS, AND 4) NO CONCURRENT USE WITH PEMBROLIZUMAB, NIVOLUMAB, IPILIMUMAB, DABRAFENIB, TRAMETINIB, VEMURAFENIB, INTERLEUKIN-2, INTERFERON, DACARBAZINE, TEMOZOLOMIDE, PACLITAXEL, CARBOPLATIN, IMATINIB, MELPHALAN, IMIQUIMOD, OR RADIATION THERAPY.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TALQUETAMAB-TGVS

Products Affected

• TALVEY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TASIMELTEON

- HETLIOZ LQ
- tasimelteon

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	LIFETIME
Other Criteria	NON-24 HOUR SLEEP-WAKE DISORDER: PATIENT IS LIGHT-INSENSITIVE OR HAS TOTAL BLINDNESS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TAZEMETOSTAT

Products Affected

• TAZVERIK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TBO-FILGRASTIM

Products Affected

• GRANIX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	NON-MYELOID MALIGNANCIES: PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST OR ONCOLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	NON-MYELOID MALIGNANCIES: TRIAL OF OR CONTRAINDICATION TO THE PREFERRED AGENT: NIVESTYM.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TEBENTAFUSP-TEBN

Products Affected

• KIMMTRAK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TECLISTAMAB-CQYV

Products Affected

• TECVAYLI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TEDUGLUTIDE

Products Affected

• GATTEX 30-VIAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	SHORT BOWEL SYNDROME (SBS): INITIAL/RENEWAL: PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS
Other Criteria	SBS: INITIAL: PATIENT IS DEPENDENT ON INTRAVENOUS PARENTERAL NUTRITION DEFINED AS REQUIRING PARENTERAL NUTRITION AT LEAST THREE TIMES PER WEEK. RENEWAL: ACHIEVED OR MAINTAINED A DECREASED NEED FOR PARENTERAL SUPPORT COMPARED TO BASELINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TELOTRISTAT

Products Affected

• XERMELO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	CARCINOID SYNDROME DIARRHEA: PRESCRIBED BY OR IN CONSULTATION WITH AN ONCOLOGIST OR GASTROENTEROLOGIST
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TEPOTINIB

Products Affected

• ТЕРМЕТКО

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TEPROTUMUMAB-TRBW

Products Affected

• TEPEZZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TERIFLUNOMIDE

Products Affected

• teriflunomide

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TESAMORELIN

Products Affected

• EGRIFTA SV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	3 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TESTOSTERONE

Products Affected

- testosterone transdermal gel in metereddose pump 12.5 mgl 1.25 gram (1%), 20.25
 mgl1.25 gram (1.62%)
- testosterone transdermal gel in packet 1 %

(25 mg/2.5gram), 1 % (50 mg/5 gram) testosterone transdermal solution in metered pump wlapp

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	MALE HYPOGONADISM: INITIAL: CONFIRMED BY: 1) AT LEAST TWO TOTAL SERUM TESTOSTERONE LEVELS OF LESS THAN 300 NG/DL TAKEN ON SEPARATE OCCASIONS, OR 2) FREE SERUM TESTOSTERONE LEVEL OF LESS THAN 5 NG/DL.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	MALE HYPOGONADISM: INITIAL: 1) 40 YEARS OR OLDER: PROSTATE SPECIFIC ANTIGEN HAS BEEN EVALUATED FOR PROSTATE CANCER SCREENING. RENEWAL: 1) 40 YEARS OR OLDER: PROSTATE SPECIFIC ANTIGEN HAS BEEN EVALUATED FOR PROSTATE CANCER SCREENING, AND 2) IMPROVED SYMPTOMS COMPARED TO BASELINE AND TOLERANCE TO TREATMENT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TESTOSTERONE CYPIONATE

Products Affected

• testosterone cypionate

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	MALE HYPOGONADISM: INITIAL: CONFIRMED BY: 1) AT LEAST TWO TOTAL SERUM TESTOSTERONE LEVELS OF LESS THAN 300 NG/DL TAKEN ON SEPARATE OCCASIONS, OR 2) FREE SERUM TESTOSTERONE LEVEL OF LESS THAN 5 NG/DL.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	MALE HYPOGONADISM: INITIAL: 1) 40 YEARS OR OLDER: PROSTATE SPECIFIC ANTIGEN HAS BEEN EVALUATED FOR PROSTATE CANCER SCREENING. RENEWAL: 1) 40 YEARS OR OLDER: PROSTATE SPECIFIC ANTIGEN HAS BEEN EVALUATED FOR PROSTATE CANCER SCREENING, AND 2) IMPROVED SYMPTOMS COMPARED TO BASELINE AND TOLERANCE TO TREATMENT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TESTOSTERONE ENANTHATE

- *testosterone enanthate*
- XYOSTED

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	MALE HYPOGONADISM: INITIAL: CONFIRMED BY: 1) AT LEAST TWO TOTAL SERUM TESTOSTERONE LEVELS OF LESS THAN 300 NG/DL TAKEN ON SEPARATE OCCASIONS, OR 2) FREE SERUM TESTOSTERONE LEVEL OF LESS THAN 5 NG/DL.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	MALE HYPOGONADISM: INITIAL/RENEWAL: 12 MO. ALL OTHER INDICATIONS: LIFETIME OF MEMBERSHIP IN PLAN.
Other Criteria	MALE HYPOGONADISM: INITIAL: 1) 40 YEARS OR OLDER: PROSTATE SPECIFIC ANTIGEN HAS BEEN EVALUATED FOR PROSTATE CANCER SCREENING. RENEWAL: 1) 40 YEARS OR OLDER: PROSTATE SPECIFIC ANTIGEN HAS BEEN EVALUATED FOR PROSTATE CANCER SCREENING, AND 2) IMPROVED SYMPTOMS COMPARED TO BASELINE AND TOLERANCE TO TREATMENT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TETRABENAZINE

Products Affected

• tetrabenazine

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	HUNTINGTONS DISEASE: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST OR MOVEMENT DISORDER SPECIALIST
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TEZACAFTOR/IVACAFTOR

Products Affected

• SYMDEKO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CYSTIC FIBROSIS (CF): INITIAL: CONFIRMED MUTATION IN CFTR GENE ACCEPTABLE FOR THE TREATMENT OF CYSTIC FIBROSIS.
Age Restrictions	
Prescriber Restrictions	CF: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST OR CYSTIC FIBROSIS EXPERT
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: LIFETIME
Other Criteria	CF: RENEWAL: MAINTAINED, IMPROVED, OR DEMONSTRATED LESS THAN EXPECTED DECLINE IN FEV1 OR BODY MASS INDEX (BMI), OR REDUCTION IN NUMBER OF PULMONARY EXACERBATIONS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

THALIDOMIDE

Products Affected

• THALOMID

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TILDRAKIZUMAB-ASMN

Products Affected

• ILUMYA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PLAQUE PSORIASIS (PSO): INITIAL: PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, GENITAL AREA, OR FACE.
Age Restrictions	
Prescriber Restrictions	PSO: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.
Other Criteria	PSO: INITIAL: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: COSENTYX, ENBREL, HUMIRA, STELARA, SKYRIZI, TREMFYA, OTEZLA. RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TISOTUMAB VEDOTIN-TFTV

Products Affected

• TIVDAK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TIVOZANIB

Products Affected

• FOTIVDA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TOCILIZUMAB IV

Products Affected

• ACTEMRA

PA Criteria	Criteria Details
Exclusion Criteria	CORONAVIRUS DISEASE 2019 (COVID-19) IN HOSPITALIZED ADULTS
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. SYSTEMIC JUVENILE IDIOPATHIC ARTHRITIS (SJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST, DERMATOLOGIST, OR IMMUNOLOGIST.
Coverage Duration	INITIAL: RA, PJIA, SJIA, GCA: 6 MONTHS. CRS: 1 MONTH. RENEWAL: RA, PJIA, SJIA, GCA: 12 MONTHS.
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PJIA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ IR, RINVOQ. SJIA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG). RENEWAL: RA, PJIA, SJIA: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

TOCILIZUMAB SQ

- ACTEMRA
- ACTEMRA ACTPEN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. SYSTEMIC JUVENILE IDIOPATHIC ARTHRITIS (SJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST, DERMATOLOGIST, OR IMMUNOLOGIST. SYSTEMIC SCLEROSIS-ASSOCIATED INTERSTITIAL LUNG DISEASE (SSC-ILD): PRESCRIBED BY OR IN CONSULTATION WITH A PULMONOLOGIST OR RHEUMATOLOGIST.
Coverage Duration	INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: 1) TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ, RINVOQ, OR 2) TRIAL OF A TNF INHIBITOR AND PHYSICIAN HAS INDICATED THE PATIENT CANNOT USE A JAK INHIBITOR DUE TO THE BLACK BOX WARNING FOR INCREASED RISK OF MORTALITY, MALIGNANCIES, AND SERIOUS CARDIOVASCULAR EVENTS. PJIA: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING PREFERRED AGENTS: HUMIRA, ENBREL, XELJANZ IR, RINVOQ. SJIA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG). SSC-ILD: DOES NOT HAVE OTHER KNOWN CAUSES OF INTERSTITIAL LUNG DISEASE (E.G., CONNECTIVE TISSUE DISEASE, DRUG TOXICITY, ASBESTOS OR BERYLLIUM EXPOSURE, HYPERSENSITIVITY PNEUMONITIS). RENEWAL: RA, PJIA, SJIA: CONTINUES TO BENEFIT FROM THE MEDICATION. SSC-ILD: CLINICAL MEANINGFUL IMPROVEMENT OR MAINTENANCE IN ANNUAL RATE OF DECLINE.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TOFACITINIB

- XELJANZ
- XELJANZ XR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	INITIAL: RHEUMATOID ARTHRITIS (RA), ANKYLOSING SPONDYLITIS (AS), POLYARTICULAR COURSE JUVENILE IDIOPATHIC ARTHRITIS (PCJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR DERMATOLOGIST. ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: RA: TRIAL OF OR CONTRAINDICATION TO 3 MONTHS OF TREATMENT WITH ONE DMARD (DISEASE- MODIFYING ANTIRHEUMATIC DRUG) - IF A PATIENT TRIED METHOTREXATE, THEN TRIAL AT A DOSE GREATER THAN OR EQUAL TO 20MG PER WEEK OR MAXIMALLY TOLERATED DOSE IS REQUIRED. PSA, PCJIA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD. UC: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. RENEWAL: RA, PSA, AS, PCJIA: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

TOLVAPTAN

- JYNARQUE ORAL TABLET
- JYNARQUE ORAL TABLETS, SEQUENTIAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	AUTOSOMAL DOMINANT POLYCYSTIC KIDNEY DISEASE (ADPKD): INITIAL: 1) CONFIRMED POLYCYSTIC KIDNEY DISEASE VIA CT, MRI, OR ULTRASOUND, AND 2) GENETIC TESTING FOR CAUSATIVE MUTATIONS OR FAMILY HISTORY OF CONFIRMED POLYCYSTIC KIDNEY DISEASE IN ONE OR BOTH PARENTS.
Age Restrictions	
Prescriber Restrictions	ADPKD: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A NEPHROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	ADPKD: INITIAL: DOES NOT HAVE ESRD (I.E., RECEIVING DIALYSIS OR HAS UNDERGONE RENAL TRANSPLANT). RENEWAL: HAS NOT PROGRESSED TO ESRD/DIALYSIS OR TRANSPLANT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TOPICAL TRETINOIN

- ALTRENO
- tretinoin

PA Criteria	Criteria Details
Exclusion Criteria	COSMETIC INDICATIONS SUCH AS WRINKLES, PHOTOAGING, MELASMA.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	ACNE VULGARIS: BRAND TOPICAL TRETINOIN REQUIRES TRIAL OF OR CONTRAINDICATION TO A GENERIC TOPICAL TRETINOIN PRODUCT.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TORIPALIMAB-TPZI

Products Affected

• LOQTORZI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	NASOPHARYNGEAL CARCINOMA (NPC): FIRST LINE TREATMENT: 24 MOS, PREVIOUSLY TREATED: LIFETIME.
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TOVORAFENIB

Products Affected

• OJEMDA 100 MG TAB (400 MG DOSE)

MG/WEEK (100 MG X 5)

- OJEMDA ORAL SUSPENSION FOR RECONSTITUTION
- OJEMDA ORAL TABLET 500

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRAMETINIB

Products Affected

• MEKINIST ORAL TABLET 0.5 MG, 2 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRAMETINIB SOLUTION

Products Affected

• MEKINIST ORAL RECON SOLN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	UNABLE TO SWALLOW MEKINIST TABLETS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRASTUZUMAB HYALURONIDASE

Products Affected

• HERCEPTIN HYLECTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	ADJUVANT BREAST CANCER, METASTATIC BREAST CANCER: TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING PREFERRED AGENTS: HERZUMA, OGIVRI, ONTRUZANT, TRAZIMERA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRASTUZUMAB-ANNS

Products Affected

• KANJINTI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	ADJUVANT BREAST CANCER, METASTATIC BREAST CANCER: TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING PREFERRED AGENTS: HERZUMA, OGIVRI, ONTRUZANT, TRAZIMERA. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRASTUZUMAB-DKST

Products Affected

• OGIVRI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRASTUZUMAB-DTTB

Products Affected

• ONTRUZANT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRASTUZUMAB-PKRB

Products Affected

• HERZUMA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRASTUZUMAB-QYYP

Products Affected

• TRAZIMERA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TREMELIMUMAB-ACTL

Products Affected

• IMJUDO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	UHCC: 30 DAYS. METASTATIC NON-SMALL CELL LUNG CANCER (NSCLC): 5 MONTHS.
Other Criteria	UNRESECTABLE HEPATOCELLULAR CARCINOMA (UHCC): HAS NOT RECEIVED PRIOR TREATMENT WITH IMJUDO.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TREPROSTINIL INHALED

Products Affected

• TYVASO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PULMONARY ARTERIAL HYPERTENSION (PAH), PULMONARY HYPERTENSION-INTERSTITIAL LUNG DISEASE (PH-ILD): DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	INITIAL: PAH, PH-ILD: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL: PAH: 12 MONTHS, PH-ILD: 6 MONTHS. RENEWAL: PAH, PH-ILD: 12 MONTHS.
Other Criteria	INITIAL: PAH: TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING AGENTS FROM DIFFERENT DRUG CLASSES: 1) FORMULARY VERSION OF AN ORAL ENDOTHELIN RECEPTOR ANTAGONIST, 2) FORMULARY VERSION OF AN ORAL PHOSPHODIESTERASE TYPE-5 INHIBITOR, 3) FORMULARY VERSION OF AN ORAL CGMP STIMULATOR, 4) FORMULARY VERSION OF AN IV/SQ PROSTACYCLIN. THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

TREPROSTINIL INJECTABLE

Products Affected

• treprostinil sodium

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	PULMONARY ARTERIAL HYPERTENSION (PAH): INITIAL: DIAGNOSIS CONFIRMED BY RIGHT HEART CATHETERIZATION WITH THE FOLLOWING PARAMETERS: 1) MEAN PULMONARY ARTERY PRESSURE (PAP) GREATER THAN 20 MMHG, 2) PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, AND 3) PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 2 WOOD UNITS.
Age Restrictions	
Prescriber Restrictions	PAH: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS
Other Criteria	PAH: INITIAL: 1) CONTINUATION OF THERAPY FROM HOSPITAL DISCHARGE, 2) NEW START AND PHYSICIAN INDICATED PATIENT IS INTERMEDIATE OR HIGH RISK, OR 3) NEW START AND TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING AGENTS FROM DIFFERENT DRUG CLASSES: (A) FORMULARY VERSION OF AN ORAL ENDOTHELIN RECEPTOR ANTAGONIST, (B) FORMULARY VERSION OF AN ORAL PHOSPHODIESTERASE TYPE-5 INHIBITOR, OR (C) FORMULARY VERSION OF AN ORAL CGMP STIMULATOR. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

TRIENTINE CAPSULE

Products Affected

• trientine oral capsule 250 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	WILSONS DISEASE: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A HEPATOLOGIST OR GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 12 MONTHS, RENEWAL: LIFETIME.
Other Criteria	WILSONS DISEASE: INITIAL: 1) LEIPZIG SCORE OF 4 OR GREATER, AND 2) TRIAL OF OR CONTRAINDICATION TO FORMULARY VERSION OF PENICILLAMINE TABLET. RENEWAL: CONTINUES TO BENEFIT FROM THE MEDICATION.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRIFLURIDINE/TIPIRACIL

Products Affected

• LONSURF ORAL TABLET 15-6.14 MG, 20-8.19 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRIPTORELIN-TRELSTAR

Products Affected

 TRELSTAR INTRAMUSCULAR SUSPENSION FOR RECONSTITUTION

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS.
Other Criteria	THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

TRIPTORELIN-TRIPTODUR

Products Affected

• TRIPTODUR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	CENTRAL PRECOCIOUS PUBERTY (CPP): INITIAL: FEMALES: ELEVATED LEVELS OF FOLLICLE- STIMULATING HORMONE (FSH) GREATER THAN 4.0 MIU/ML AND LUTEINIZING HORMONE (LH) LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS. MALES: ELEVATED LEVELS OF FSH GREATER THAN 5.0 MIU/ML AND LH LEVEL GREATER THAN 0.2 TO 0.3 MIU/ML AT DIAGNOSIS.
Age Restrictions	
Prescriber Restrictions	CPP: INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH AN ENDOCRINOLOGIST.
Coverage Duration	INITIAL/RENEWAL: 12 MONTHS.
Other Criteria	CPP: INITIAL: FEMALES: 1) YOUNGER THAN 8 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR BREAST DEVELOPMENT AND PUBIC HAIR GROWTH. MALES: 1) YOUNGER THAN 9 YEARS OF AGE AT ONSET OF CPP, AND 2) AT TANNER STAGE 2 OR ABOVE FOR GENITAL DEVELOPMENT AND PUBIC HAIR GROWTH. RENEWAL: 1) TANNER STAGING AT INITIAL DIAGNOSIS HAS STABILIZED OR REGRESSED DURING THREE SEPARATE MEDICAL VISITS IN THE PREVIOUS YEAR, AND 2) HAS NOT REACHED ACTUAL AGE WHICH CORRESPONDS TO CURRENT PUBERTAL AGE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

TUCATINIB

Products Affected

 TUKYSA ORAL TABLET 150 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

UBROGEPANT

Products Affected

• UBRELVY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	ACUTE MIGRAINE TREATMENT: INITIAL: TRIAL OF OR CONTRAINDICATION TO ONE TRIPTAN (E.G., SUMATRIPTAN, RIZATRIPTAN). RENEWAL: 1) IMPROVEMENT FROM BASELINE IN A VALIDATED ACUTE TREATMENT PATIENT-REPORTED OUTCOME QUESTIONNAIRE, OR 2) THERAPY WORKS CONSISTENTLY IN MAJORITY OF MIGRAINE ATTACKS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

UPADACITINIB

Products Affected

• RINVOQ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	NON-RADIOGRAPHIC AXIAL SPONDYLOARTHRITIS (NR- AXSPA): 1) C-REACTIVE PROTEIN LEVELS ABOVE THE UPPER LIMIT OF NORMAL, OR 2) SACROILIITIS ON MAGNETIC RESONANCE IMAGING (MRI).
Age Restrictions	
Prescriber Restrictions	INITIAL: RA, AS, NR-AXSPA, POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSA: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST OR DERMATOLOGIST. AD: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST, ALLERGIST, OR IMMUNOLOGIST. UC, CD: PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.

PA Criteria	Criteria Details
Other Criteria	INITIAL: RA: TRIAL OF OR CONTRAINDICATION TO 3 MONTHS OF TREATMENT WITH ONE DMARD (DISEASE- MODIFYING ANTIRHEUMATIC DRUG) - IF A PATIENT TRIED METHOTREXATE, THEN TRIAL AT A DOSE GREATER THAN OR EQUAL TO 20MG PER WEEK OR MAXIMALLY TOLERATED DOSE IS REQUIRED. PSA, PJIA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD. ATOPIC DERMATITIS: 1) ATOPIC DERMATITIS COVERING AT LEAST 10 PERCENT OF BODY SURFACE AREA OR ATOPIC DERMATITIS AFFECTING THE FACE, HEAD, NECK, HANDS, FEET, GROIN, OR INTERTRIGINOUS AREAS, 2) INTRACTABLE PRURITUS OR CRACKING/OOZING/BLEEDING OF AFFECTED SKIN, 3) TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING: TOPICAL CORTICOSTEROID, TOPICAL CALCINEURIN INHIBITOR, TOPICAL PDE4 INHIBITOR, OR TOPICAL JAK INHIBITOR, AND 4) NO CONCURRENT USE WITH OTHER SYSTEMIC BIOLOGIC/JAK INHIBITOR FOR THE TREATMENT OF ATOPIC DERMATITIS. UC: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. NR-AXSPA: TRIAL OF OR CONTRAINDICATION TO AN NSAID (NON-STEROIDAL ANTI-INFLAMMATORY DRUG). CD: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. NSAID (NON-STEROIDAL ANTI-INFLAMMATORY DRUG). CD: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. RENEWAL: RA, PSA, AS, NR-AXSPA, PJIA: CONTINUES TO BENEFIT FROM THE MEDICATION. ATOPIC DERMATITIS: 1) IMPROVEMENT WHILE ON THERAPY, AND 2) NO CONCURRENT USE WITH OTHER
	SYSTEMIC BIOLOGIC/JAK INHIBITOR FOR THE TREATMENT OF ATOPIC DERMATITIS.
Indications	All FDA-approved Indications.
Off Label Uses	

PA Criteria	Criteria Details
Part B Prerequisite	No

USTEKINUMAB

Products Affected

• STELARA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	INITIAL: PLAQUE PSORIASIS (PSO): PSORIASIS COVERING 3 PERCENT OR MORE OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, GENITAL AREA, OR FACE.
Age Restrictions	
Prescriber Restrictions	INITIAL: PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSO: PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	INITIAL: PSA: TRIAL OF OR CONTRAINDICATION TO ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG). PSO: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. CD, UC: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. RENEWAL: PSA, PSO: CONTINUES TO BENEFIT FROM THE MEDICATION.

PA Criteria	Criteria Details
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

USTEKINUMAB IV

Products Affected

• STELARA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	CROHNS DISEASE (CD), ULCERATIVE COLITIS (UC): PRESCRIBED BY OR IN CONSULTATION WITH A GASTROENTEROLOGIST.
Coverage Duration	2 MONTHS
Other Criteria	CD, UC: TRIAL OF OR CONTRAINDICATION TO ONE CONVENTIONAL THERAPY SUCH AS A CORTICOSTEROID (E.G., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VALBENAZINE

Products Affected

- INGREZZA
- INGREZZA INITIATION PK(TARDIV)
- INGREZZA SPRINKLE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	TARDIVE DYSKINESIA (TD): PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST, PSYCHIATRIST, OR MOVEMENT DISORDER SPECIALIST.
Coverage Duration	12 MONTHS
Other Criteria	TD: PRIOR HISTORY OF USING AGENTS THAT CAUSE TARDIVE DYSKINESIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VANDETANIB

Products Affected

• CAPRELSA ORAL TABLET 100 MG, 300 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	CURRENTLY STABLE ON CAPRELSA REQUIRES NO EXTRA CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VEMURAFENIB

Products Affected

• ZELBORAF

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	MELANOMA: ZELBORAF WILL BE USED ALONE OR IN COMBINATION WITH COTELLIC
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VENETOCLAX

Products Affected

- VENCLEXTA ORAL TABLET 10 MG, 100 MG, 50 MG
- VENCLEXTA STARTING PACK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VERICIGUAT

Products Affected

• VERQUVO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	INITIAL/RENEWAL:12 MONTHS.
Other Criteria	HEART FAILURE (HF): INITIAL: TRIAL OF OR CONTRAINDICATION TO 1) ONE AGENT FROM ANY OF THE FOLLOWING STANDARD OF CARE CLASSES: A) ACE INHIBITOR, ARB, OR ARNI, B) BETA BLOCKER (I.E., BISOPROLOL, CARVEDILOL, METOPROLOL SUCCINATE), OR C) ALDOSTERONE ANTAGONIST (I.E., SPIRONOLACTONE, EPLERENONE), AND 2) ONE PREFERRED SGLT-2 INHIBITOR (E.G., FARXIGA, XIGDUO XR, JARDIANCE). INITIAL/RENEWAL: NOT CONCURRENTLY TAKING LONG-ACTING NITRATES OR NITRIC OXIDE DONORS, RIOCIGUAT, OR PDE-5 INHIBITORS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VESTRONIDASE ALFA VJBK

Products Affected

• MEPSEVII

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	MUCOPOLYSACCHARIDOSIS VII (MPS VII): INITIAL: PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN GENETIC OR METABOLIC DISORDERS.
Coverage Duration	INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS.
Other Criteria	MPS VII: INITIAL: 1) HAS NOT UNDERGONE SUCCESSFUL BONE MARROW OR STEM CELL TREATMENT FOR MPS VII, 2) LIMITATION IN MOBILITY, BUT REMAINS SUFFICIENTLY AMBULATORY, AND 3) DIAGNOSIS CONFIRMED BY: (A) URINARY GAG (GLYCOSAMINOGLYCAN) LEVEL IS GREATER THAN THREE TIMES THE UPPER LEVEL OF NORMAL BASED ON THE LABORATORY ASSAY, (B) BETA-GLUCURONIDASE ENZYME ACTIVITY DEFICIENCY OR GENETIC TESTING, AND (C) ONE OF THE FOLLOWING CLINICAL SIGNS OF MPS VII: ENLARGED LIVER AND SPLEEN, JOINT LIMITATIONS, AIRWAY OBSTRUCTIONS OR PULMONARY DYSFUNCTION. RENEWAL: IMPROVED, MAINTAINED, OR DEMONSTRATED A LESS THAN EXPECTED DECLINE IN AMBULATORY ABILITY FROM BASELINE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D.
Indications	All FDA-approved Indications.

PA Criteria	Criteria Details
Off Label Uses	
Part B Prerequisite	No

VIGABATRIN

Products Affected

- vigabatrin
- vigadrone
- vigpoder

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	REFRACTORY COMPLEX PARTIAL SEIZURES (CPS), INFANTILE SPASMS: PRESCRIBED BY OR IN CONSULTATION WITH A NEUROLOGIST.
Coverage Duration	12 MONTHS
Other Criteria	CPS: TRIAL OF OR CONTRAINDICATION TO TWO ANTIEPILEPTIC AGENTS.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VISMODEGIB

Products Affected

• ERIVEDGE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

VORICONAZOLE SUSPENSION

Products Affected

• voriconazole oral suspension for reconstitution

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	CANDIDA INFECTIONS: 3 MOS. ALL OTHER INDICATIONS: 6 MOS.
Other Criteria	CANDIDA INFECTIONS: TRIAL OF OR CONTRAINDICATION TO FLUCONAZOLE. ALL INDICATIONS: INABILITY TO SWALLOW TABLETS OR AN INDICATION FOR ESOPHAGEAL CANDIDIASIS. CONTINUATION OF THERAPY AFTER HOSPITAL DISCHARGE REQUIRES NO EXTRA CRITERIA.
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ZANUBRUTINIB

Products Affected

• BRUKINSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 MONTHS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

ZURANOLONE

Products Affected

• ZURZUVAE ORAL CAPSULE 20 MG, 25 MG, 30 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	14 DAYS
Other Criteria	
Indications	All FDA-approved Indications.
Off Label Uses	
Part B Prerequisite	No

INDEX

abiraterone	
ACTEMRA	
ACTEMRA ACTPEN	422
ACTHAR	72
ACTHAR SELFJECT	
SUBCUTANEOUS PEN INJECTO)R
40 UNIT/0.5 ML, 80 UNIT/ML	72
ACTIMMUNE	
ADAKVEO	74
ADEMPAS	
AJOVY AUTOINJECTOR	
AJOVY SYRINGE	154
AKEEGA	
ALECENSA	15
ALTRENO	
ALUNBRIG ORAL TABLET 180	MG,
30 MG, 90 MG	
ALUNBRIG ORAL TABLETS, DO	DSE
PACK	55
ALVAIZ	120
<i>alyq</i>	395
amabelz	181
ambrisentan	17
ANKTIVA	284
apomorphine	21
armodafinil	269
ascomp with codeine	175
AUGTYRO	
AUSTEDO ORAL TABLET 12 MO	Э , 6
MG, 9 MG	93
AUSTEDO XR ORAL TABLET	
EXTENDED RELEASE 24 HR 12	
MG, 18 MG, 24 MG, 30 MG, 36 MG	G,
42 MG, 48 MG, 6 MG	93
AUSTEDO XR TITRATION	
KT(WK1-4) AVONEX INTRAMUSCULAR PI	93
AVONEX INTRAMUSCULAR PI	EN
INJECTOR KIT	213
AVONEX INTRAMUSCULAR	
SYRINGE KIT	213
AVSOLA	208
AYVAKIT	

BALVERSA ORAL TABLET 3 MG, 4	
MG, 5 MG	.134
bendamustine intravenous recon soln	43
BENDAMUSTINE INTRAVENOUS	
SOLUTION	43
BENDEKA	43
BENLYSTA INTRAVENOUS	40
BENLYSTA SUBCUTANEOUS	40
BESREMI	356
betaine	46
BETASERON SUBCUTANEOUS KIT	Γ
	214
bexarotene	
BEYFORTUS	280
bortezomib injection	52
bosentan	
BOSULIF ORAL CAPSULE 100 MG,	
50 MG	54
BOSULIF ORAL TABLET 100 MG,	
400 MG, 500 MG	54
BRAFTOVI	
BRUKINSA	
butalbital-acetaminop-caf-cod	
butalbital-acetaminophen oral tablet 50-	
	.175
butalbital-acetaminophen-caff oral	
capsule 50-325-40 mg	.175
butalbital-acetaminophen-caff oral tablet	
butalbital-aspirin-caffeine	
CABLIVI INJECTION KIT	
CABOMETYX ORAL TABLET 20	
MG, 40 MG, 60 MG	59
CALQUENCE (ACALABRUTINIB	
MAL)	8
CAPRELSA ORAL TABLET 100 MG.	
300 MG	
carbinoxamine maleate oral liquid	
carbinoxamine maleate oral tablet 4 mg	
carglumic acid	
CAYSTON	
CERDELGA	
chlorzoxazone oral tablet 250 mg, 500	
mg, 750 mg	191
<i>b</i> ,	

CINQAIR	337
CINRYZE	56
clemastine oral tablet	176
codeine-butalbital-asa-caff	
COMETRIQ ORAL CAPSULE 100	
MG/DAY(80 MG X1-20 MG X1), 140	
MG/DAY(80 MG X1-20 MG X3), 60	=0
MG/DAY (20 MG X 3/DAY)	58
COPAXONE SUBCUTANEOUS	
SYRINGE 20 MG/ML, 40 MG/ML	
COPIKTRA	
CORTROPHIN GEL	72
COSENTYX (2 SYRINGES)	365
COSENTYX INTRAVENOUS	
COSENTYX PEN (2 PENS)	
COSENTYX SUBCUTANEOUS	
SYRINGE 75 MG/0.5 ML	365
COSENTYX UNOREADY PEN	
COTELLIC	
cyclobenzaprine oral tablet 10 mg, 5 mg	
cyproheptadine	
CYRAMZA	
CYSTADROPS	
CYSTARAN	
dalfampridine	
DANYELZA	271
DARZALEX	82
DARZALEX FASPRO	83
DAURISMO ORAL TABLET 100	
MG, 25 MG	162
deferasirox	
deferiprone	
deferoxamine	
DIACOMIT ORAL CAPSULE 250	91
	202
MG, 500 MG	393
DIACOMIT ORAL POWDER IN	202
PACKET 250 MG, 500 MG	
diclofenac sodium topical gel 3 %	94
diclofenac sodium topical solution in	
metered-dose pump	95
dimethyl fumarate oral capsule, delayed	
release(dr/ec) 120 mg, 120 mg (14)-24	0
mg (46), 240 mg	96
diphenhydramine hcl oral elixir	192
diphenoxylate-atropine	

dipyridamole oral	178
disopyramide phosphate oral capsule	.179
DOPTELET (10 TAB PACK)	32
DOPTELET (15 TAB PACK)	32
DOPTELET (30 TAB PACK)	32
dotti	180
dronabinol	99
droxidopa	. 100
DUAVEE	
DUPIXENT PEN	101
DUPIXENT SYRINGE	101
edaravone intravenous solution 30 mg/10	0
<i>ml</i>	
EGRIFTA SV	410
ELFABRIO	
ELIGARD	234
ELIGARD (3 MONTH)	. 234
ELIGARD (4 MONTH)	. 234
ELIGARD (6 MONTH)	
ELREXFIO 44 MG/1.1 ML VIAL	
OUTER, SUV, P/F	
ELREXFIO SUBCUTANEOUS	
SOLUTION 40 MG/ML	. 118
EMGALITY PEN	
EMGALITY SYRINGE	
SUBCUTANEOUS SYRINGE 120	
MG/ML, 300 MG/3 ML (100 MG/ML	
X 3)	.157
ENBREL	
ENBREL MINI	
ENBREL SURECLICK	
ENDARI	
ENSPRYNG	
ENTADFI	
EPCLUSA ORAL PELLETS IN	. 177
PACKET 150-37.5 MG, 200-50 MG	380
EPCLUSA ORAL TABLET	
EPIDIOLEX	
EPKINLY	
epoprostenol	
ERBITUX	
ERIVEDGE	
ERLEADA ORAL TABLET 240 MG,	
60 MG	20

erlotinib oral tablet 100 mg, 150 mg, 25	
mg	135
estradiol oral	180
estradiol transdermal patch semiweekly.	180
estradiol transdermal patch weekly	
estradiol-norethindrone acet oral tablet	
0.5-0.1 mg	181
everolimus (antineoplastic) oral tablet 1	
mg, 2.5 mg, 5 mg, 7.5 mg	
everolimus (antineoplastic) oral tablet for	or
suspension	
EVRYSDI	. 350
EXKIVITY	264
EXONDYS-51	139
FABRAZYME	
FASENRA	44
FASENRA PEN	
fentanyl citrate buccal lozenge on a	
handle	145
FERRIPROX (2 TIMES A DAY)	89
FERRIPROX ORAL SOLUTION	
FERRIPROX ORAL TABLET 1,000	
MG	89
fingolimod	
FINTEPLA	
FOTIVDA	
FRUZAQLA ORAL CAPSULE 1 MC	
5 MG	
FULPHILA	
FYARRO	
fyavolv	
FYLNETRA	
GALAFOLD	
GAMIFANT	
GATTEX 30-VIAL	
GAVRETO	
gefitinib	
GILENYA ORAL CAPSULE 0.25 M	
GILOTRIF	
GIVLAARI	
glatiramer subcutaneous syringe 20	
mg/ml, 40 mg/ml	163
glatopa subcutaneous syringe 20 mg/ml,	105
40 mg/ml	163
10 111611111	

glutamine (sickle cell)24	0
<i>glyburide</i> 18	34
glyburide micronized18	
glyburide-metformin18	34
GRANIX40	
HAEGARDA SUBCUTANEOUS	
RECON SOLN 2,000 UNIT, 3,000	
UNIT5	57
HARVONI ORAL PELLETS IN	
PACKET 33.75-150 MG, 45-200 MG 22	28
HARVONI ORAL TABLET22	28
HERCEPTIN HYLECTA 43	52
HERZUMA 43	66
HETLIOZ LQ 40	00
HUMIRA PEN 1	
HUMIRA PEN CROHNS-UC-HS	
START 1	0
HUMIRA PEN PSOR-UVEITS-ADOL	
HS1	0
HUMIRA SUBCUTANEOUS	
SYRINGE KIT 40 MG/0.8 ML 1	0
HUMIRA(CF) 1	0
HUMIRA(CF) PEDI CROHNS	
STARTER1	0
HUMIRA(CF) PEN1	0
HUMIRA(CF) PEN CROHNS-UC-HS.1	0
HUMIRA(CF) PEN PEDIATRIC UC1	0
HUMIRA(CF) PEN PSOR-UV-ADOL	
HS1	
IBRANCE	9
ibuprofen-famotidine20	0
<i>icatibant</i>)1
ICLUSIG32	25
IDHIFA12	24
ILARIS (PF)6	50
ILUMYA41	
imatinib oral tablet 100 mg, 400 mg 20)3
IMBRUVICA ORAL CAPSULE 140	
MG, 70 MG19	
IMBRUVICA ORAL SUSPENSION 19)9
IMBRUVICA ORAL TABLET 140	
MG, 280 MG, 420 MG 19	
IMJUDO 43	8
IMLYGIC INJECTION SUSPENSION	
10EXP6 (1 MILLION) PFU/ML 39)8

IMPAVIDO	263
INBRIJA INHALATION CAPSULE,	
W/INHALATION DEVICE	239
indomethacin oral capsule 25 mg, 50 mg	
indomethacin oral capsule, extended	
release	194
INFLECTRA	
infliximab	204
INGREZZA	456
INGREZZA INITIATION	
PK(TARDIV)	456
INGREZZA SPRINKLE	
INLYTA ORAL TABLET 1 MG, 5	
MG	33
INQOVI	. 86
INREBIC	143
IQIRVO	
itraconazole oral solution	218
IWILFIN	
JAKAFI	358
javygtor oral tablet,soluble	
JAYPIRCA ORAL TABLET 100 MG,	
50 MG	323
50 MG JEMPERLI	
50 MG JEMPERLI <i>jinteli</i>	. 98
JEMPERLI	. 98 186
JEMPERLI	. 98 186
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG,	. 98 186 245
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG	. 98 186 245
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET	. 98 186 245 426
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS,	. 98 186 245 426 426
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL	. 98 186 245 426 426 219
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO	. 98 186 245 426 426 219 433
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA	. 98 186 245 426 426 219 433 363 150
JEMPERLI. <i>jinteli</i> JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA	. 98 186 245 426 426 219 433 363 150
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA	. 98 186 245 426 426 219 433 363 150
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KESIMPTA PEN	. 98 186 245 426 426 219 433 363 150 287
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KERENDIA KESIMPTA PEN ketorolac injection solution 15 mg/ml, 30	. 98 186 245 426 426 219 433 363 150 287
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KERENDIA ketorolac injection solution 15 mg/ml, 30 mg/ml, 30 mg/ml (1 ml)	. 98 186 245 426 219 433 363 150 287 185
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KERENDIA ketorolac injection solution 15 mg/ml, 30 mg/ml, 30 mg/ml (1 ml) ketorolac injection syringe 15 mg/ml, 30	. 98 186 245 426 219 433 363 150 287 185
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KESIMPTA PEN ketorolac injection solution 15 mg/ml, 30 mg/ml, 30 mg/ml (1 ml) ketorolac injection syringe 15 mg/ml, 30 mg/ml	. 98 186 245 426 219 433 363 150 287 185 185
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KERENDIA KESIMPTA PEN ketorolac injection solution 15 mg/ml, 30 mg/ml, 30 mg/ml (1 ml) ketorolac injection syringe 15 mg/ml, 30 mg/ml ketorolac intramuscular solution	. 98 186 245 426 219 433 363 150 287 185 185 185
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KESIMPTA PEN ketorolac injection solution 15 mg/ml, 30 mg/ml, 30 mg/ml (1 ml) ketorolac injection syringe 15 mg/ml, 30 mg/ml ketorolac intramuscular solution ketorolac intramuscular solution	. 98 186 245 426 219 433 363 150 287 185 185 185 185
JEMPERLI jinteli JUXTAPID ORAL CAPSULE 10 MG, 20 MG, 30 MG, 40 MG, 5 MG, 60 MG JYNARQUE ORAL TABLET JYNARQUE ORAL TABLETS, SEQUENTIAL KALYDECO KANJINTI KANUMA KERENDIA KERENDIA KESIMPTA PEN ketorolac injection solution 15 mg/ml, 30 mg/ml ketorolac injection syringe 15 mg/ml, 30 mg/ml ketorolac intramuscular solution ketorolac oral	. 98 186 245 426 219 433 363 150 287 185 185 185 185 361 315

KINERET	19
KISQALI FEMARA CO-PACK ORA	
TABLET 200 MG/DAY(200 MG X 1)-	
2.5 MG, 400 MG/DAY(200 MG X 2)-	
2.5 MG, 600 MG/DAY(200 MG X 3)-	
2.5 MG	341
KISQALI ORAL TABLET 200	
MG/DAY (200 MG X 1), 400 MG/DAY	Y
(200 MG X 2), 600 MG/DAY (200 MG	
X 3)	
KOSELUGO ORAL CAPSULE 10	
MG, 25 MG	.371
KRAZATI	
KYNMOBI SUBLINGUAL FILM 10	
MG, 10-15-20-25-30 MG, 15 MG, 20	
MG, 25 MG, 30 MG	2.2
lanreotide subcutaneous syringe 120	
mg/0.5 ml	225
lapatinib	
lenalidomide	
LENVIMA	
leuprolide (3 month)	
leuprolide subcutaneous kit	
lidocaine hcl mucous membrane solution	
% (40 mg/ml)	
lidocaine topical adhesive patch, medicate	
5 %	
lidocaine topical ointment	
lidocaine-prilocaine topical cream	
LONSURF ORAL TABLET 15-6.14	273
MG, 20-8.19 MG	ΔΔΔ
LOQTORZI	
LORBRENA ORAL TABLET 100	.720
MG, 25 MG.	248
LUMAKRAS ORAL TABLET 120	.240
MG, 320 MG.	392
LUNSUMIO	
LUPRON DEPOT	
LUPRON DEPOT (3 MONTH)	
LUPRON DEPOT (4 MONTH)	
LUPRON DEPOT (6 MONTH)	
LUPRON DEPOT (6 MONTH)	
LUPRON DEPOT-PED (3 MONTH).	
LYBALVI	
lyllana	100

LYNPARZA
LYTGOBI ORAL TABLET 12
MG/DAY (4 MG X 3), 16 MG/DAY (4
MG X 4), 20 MG/DAY (4 MG X 5)156
MARGENZA252
MAVENCLAD (10 TABLET PACK) 69
MAVENCLAD (4 TABLET PACK)69
MAVENCLAD (5 TABLET PACK)69
MAVENCLAD (6 TABLET PACK) 69
MAVENCLAD (7 TABLET PACK) 69
MAVENCLAD (8 TABLET PACK) 69
MAVENCLAD (9 TABLET PACK) 69
MAVYRET ORAL TABLET164
MAYZENT ORAL TABLET 0.25 MG,
1 MG, 2 MG
MAYZENT STARTER(FOR 1MG
MAINT)
MAYZENT STARTER(FOR 2MG
MAINT)
megestrol oral suspension 400 mg/10 ml
(40 mg/ml)195
megestrol oral tablet195
MEKINIST ORAL RECON SOLN 431
MEKINIST ORAL TABLET 0.5 MG, 2
MG
MEKTOVI
MEPSEVII
methocarbamol oral tablet 500 mg, 750
<i>mg</i>
mifepristone oral tablet 300 mg
miglustat
<i>mimvey</i>
modafinil oral tablet 100 mg, 200 mg 269
<i>morphine concentrate oral solution</i> 172 MOUNJARO168
MOUNJARO
NATPARA
NATPARA
NERLINA 275 NEULASTA ONPRO
NINLARO
nitisinone
NIVESTYM
NORDITROPIN FLEXPRO
norethindrone ac-eth estradiol oral tablet
0.5-2.5 mg-mcg, 1-5 mg-mcg
5.5 2.6 mg meg, 1 6 mg meg

NOXAFIL ORAL SUSP, DELAYED	
RELEASE FOR RECON	327
NPLATE	355
NUBEQA	84
NUCALA SUBCUTANEOUS AUTO-	-
INJECTOR	254
INJECTOR NUCALA SUBCUTANEOUS RECON	N
SOLN	
NUCALA SUBCUTANEOUS	
SYRINGE 100 MG/ML, 40 MG/0.4	
ML	.254
NUPLAZID	320
NURTEC ODT	. 343
NYVEPRIA	304
OCALIVA	. 285
OCREVUS	286
ODOMZO	388
OFEV	275
OGIVRI	.434
OGSIVEO ORAL TABLET 100 MG,	
150 MG, 50 MG	279
OJEMDA 100 MG TAB (400 MG	
DOSE)	100
DO2E)	429
OJEMDA ORAL SUSPENSION FOR	
,	
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500	.429
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION	.429
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500	.429 429
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5)	.429 429 265
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA	.429 429 265 36
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT	.429 429 265 36 .295
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS	.429 429 265 36 .295 .435
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO	.429 429 265 36 .295 .435 34 .282
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG	.429 429 265 36 .295 .435 34 .282
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO	.429 429 265 36 .295 .435 34 .282 283
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPSUMIT ORENCIA	.429 265 36 .295 .435 34 .282 283 .251 3
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG ORENCIA. ORENCIA (WITH MALTOSE)	.429 265 36 .295 .435 34 .282 283 .251 3 1
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG OPSUMIT ORENCIA (WITH MALTOSE) ORENCIA CLICKJECT	.429 265 36 .295 .435 .435 .282 283 .251 3 1 3
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG ORENCIA. ORENCIA (WITH MALTOSE)	.429 265 36 .295 .435 .435 .282 283 .251 3 1 3
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG OPSUMIT ORENCIA. ORENCIA (WITH MALTOSE) ORENCIA CLICKJECT ORFADIN ORAL SUSPENSION ORGOVYX	.429 265 36 .295 .435 .435 .282 283 .251 3 1 3 281
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG OPSUMIT ORENCIA (WITH MALTOSE) ORENCIA CLICKJECT ORFADIN ORAL SUSPENSION ORGOVYX ORILISSA ORAL TABLET 150 MG,	.429 265 36 .295 .435 34 .282 283 .251 3 1 3 281 .335
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG OPSUMIT ORENCIA ORENCIA ORENCIA (WITH MALTOSE) ORENCIA CLICKJECT ORFADIN ORAL SUSPENSION ORGOVYX ORILISSA ORAL TABLET 150 MG, 200 MG	.429 265 36 .295 .435 34 .282 283 .251 3 1 3 281 .335
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG OPSUMIT ORENCIA (WITH MALTOSE) ORENCIA CLICKJECT ORFADIN ORAL SUSPENSION ORGOVYX ORILISSA ORAL TABLET 150 MG,	.429 265 36 .295 .435 34 .282 283 .251 3 1 3 281 .335
OJEMDA ORAL SUSPENSION FOR RECONSTITUTION OJEMDA ORAL TABLET 500 MG/WEEK (100 MG X 5) OJJAARA OLUMIANT ONGENTYS ONTRUZANT ONUREG OPDIVO OPDUALAG OPDUALAG OPSUMIT ORENCIA ORENCIA ORENCIA (WITH MALTOSE) ORENCIA CLICKJECT ORFADIN ORAL SUSPENSION ORGOVYX ORILISSA ORAL TABLET 150 MG, 200 MG	.429 265 36 .295 .435 .435 .282 283 .251 3 1 3 281 .335 .112 .249

ORSERDU ORAL TABLET 345 MG,	
86 MG	109
OTEZLA ORAL TABLET 30 MG	.23
OTEZLA STARTER ORAL	
TABLETS, DOSE PACK 10 MG (4)-20	
MG (4)-30 MG (47), 10 MG (4)-20 MG	
(4)-30 MG(19)	23
oxandrolone	
OXLUMO	
oxycodone oral concentrate	
OZEMPIC SUBCUTANEOUS PEN	
INJECTOR 0.25 MG OR 0.5 MG (2	
MG/3 ML), 0.25 MG OR 0.5 MG(2	
MG/1.5 ML), 1 MG/DOSE (4 MG/3	
ML), 2 MG/DOSE (8 MG/3 ML)	167
PALYNZIQ.	
paroxetine hcl oral suspension	
paroxetine hcl oral tablet	
pazopanib	
PEGASYS	
PEMAZYRE	
penicillamine oral tablet	
phenobarbital	
PIQRAY ORAL TABLET 200	107
MG/DAY (200 MG X 1), 250 MG/DAY	,
(200 MG X1-50 MG X1), 300 MG/DAY	
(150 MG X 2)	
	16
pirfenidone oral capsule	
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg,	321
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg	321
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN	321
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63	321 321
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML	321 321
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS	321 321
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63	321321215
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML	321321215215
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST	 321 321 215 215 324
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST posaconazole oral	 321 321 215 215 324 326
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST posaconazole oral PREMARIN ORAL	 321 321 321 215 215 324 326 177
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST posaconazole oral PREMARIN ORAL PREMPHASE	 321 321 321 215 215 324 326 177 183
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST posaconazole oral PREMARIN ORAL PREMPHASE PREMPRO	 321 321 321 215 215 324 326 177 183
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST posaconazole oral PREMARIN ORAL PREMPHASE PREMPHASE PREMPRO PREVYMIS INTRAVENOUS	 321 321 321 215 215 324 326 177 183
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST posaconazole oral PREMARIN ORAL PREMPHASE PREMPHASE PREMPRO PREVYMIS INTRAVENOUS SOLUTION 240 MG/12 ML, 480	 321 321 321 215 215 324 326 177 183 183
pirfenidone oral capsule pirfenidone oral tablet 267 mg, 534 mg, 801 mg PLEGRIDY SUBCUTANEOUS PEN INJECTOR 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML PLEGRIDY SUBCUTANEOUS SYRINGE 125 MCG/0.5 ML, 63 MCG/0.5 ML- 94 MCG/0.5 ML POMALYST posaconazole oral PREMARIN ORAL PREMPHASE PREMPHASE PREMPRO PREVYMIS INTRAVENOUS	 321 321 321 215 215 324 326 177 183 183 231

PROMACTA ORAL POWDER IN	
PACKET 12.5 MG, 25 MG	.119
PROMACTA ORAL TABLET 12.5	
MG, 25 MG, 50 MG, 75 MG	. 119
promethazine injection solution	. 188
promethazine oral	. 188
promethazine rectal	188
promethegan	. 188
pyrimethamine	. 330
QINLOCK	. 347
quinine sulfate	
QULIPTA	
RADICAVA	105
RAVICTI	. 169
REGRANEX	38
RELEUKO	
RELISTOR ORAL	. 258
RELISTOR SUBCUTANEOUS	
SOLUTION	257
RELISTOR SUBCUTANEOUS	
SYRINGE 12 MG/0.6 ML, 8 MG/0.4	
ML	
RENFLEXIS	201
RETACRIT INJECTION SOLUTION	
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML,	1
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML,	1
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML,	, 1
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML.	
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG	
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG	
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI	
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI REZLIDHIA	131
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI REZLIDHIA REZUROCK	131 , . 370 . 114 290 41
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI REZLIDHIA REZUROCK RIABNI	131 , . 370 114 290 41 353
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI REZLIDHIA REZLIDHIA REZUROCK RIABNI RINVOQ	131 ,
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG. REVCOVI. REZLIDHIA. REZUROCK. RIABNI. RINVOQ. RITUXAN HYCELA.	131 ,
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI REZLIDHIA REZLIDHIA REZUROCK RIABNI RINVOQ RITUXAN HYCELA RIVFLOZA	131 ,
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI REZLIDHIA REZLIDHIA REZUROCK RIABNI RINVOQ RITUXAN HYCELA RIVFLOZA ROLVEDON	131 ,
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML. RETEVMO ORAL CAPSULE 40 MG 80 MG. REVCOVI. REZLIDHIA. REZUROCK. RIABNI. RINVOQ. RITUXAN HYCELA. RIVFLOZA. ROLVEDON. ROZLYTREK ORAL CAPSULE 100	131 ,
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML RETEVMO ORAL CAPSULE 40 MG 80 MG RETEVMO ORAL CAPSULE 40 MG 80 MG REVCOVI REZLIDHIA REZLIDHIA REZUROCK RIABNI RINVOQ RITUXAN HYCELA RIVFLOZA ROLVEDON ROZLYTREK ORAL CAPSULE 100 MG, 200 MG	131 ,
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML. RETEVMO ORAL CAPSULE 40 MG 80 MG. REVCOVI. REZLIDHIA. REZLIDHIA. REZUROCK. RIABNI. RINVOQ. RITUXAN HYCELA. RIVFLOZA. ROLVEDON. ROZLYTREK ORAL CAPSULE 100 MG, 200 MG. ROZLYTREK ORAL PELLETS IN	131 , , 370 114 290 41 353 450 351 272 107 126
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML. RETEVMO ORAL CAPSULE 40 MG 80 MG. REVCOVI. REZLIDHIA. REZLIDHIA. REZUROCK. RIABNI. RINVOQ. RITUXAN HYCELA. RIVFLOZA. ROLVEDON. ROZLYTREK ORAL CAPSULE 100 MG, 200 MG. ROZLYTREK ORAL PELLETS IN PACKET.	131 ,
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML. RETEVMO ORAL CAPSULE 40 MG 80 MG. REVCOVI. REZLIDHIA. REZLIDHIA. REZUROCK. RIABNI. RINVOQ. RITUXAN HYCELA. RIVFLOZA. ROLVEDON. ROZLYTREK ORAL CAPSULE 100 MG, 200 MG. ROZLYTREK ORAL PELLETS IN PACKET. RUBRACA.	131 , 370 41 353 450 351 272 107 126 127 357
RETACRIT INJECTION SOLUTION 10,000 UNIT/ML, 2,000 UNIT/ML, 20,000 UNIT/2 ML, 20,000 UNIT/ML, 3,000 UNIT/ML, 4,000 UNIT/ML, 40,000 UNIT/ML. RETEVMO ORAL CAPSULE 40 MG 80 MG. REVCOVI. REZLIDHIA. REZLIDHIA. REZUROCK. RIABNI. RINVOQ. RITUXAN HYCELA. RIVFLOZA. ROLVEDON. ROZLYTREK ORAL CAPSULE 100 MG, 200 MG. ROZLYTREK ORAL PELLETS IN PACKET.	131 ,

RYBREVANT	. 18
RYDAPT	259
sajazir	201
sapropterin oral tablet, soluble	360
SCEMBLIX ORAL TABLET 100 MG,	
20 MG, 40 MG	25
scopolamine base	190
SEROSTIM SUBCUTANEOUS	
RECON SOLN 4 MG, 5 MG, 6 MG	386
SIGNIFOR	302
sildenafil (pulm.hypertension)	
intravenous	372
sildenafil (pulm.hypertension) oral tablet	373
SIRTURO	39
SKYRIZI	348
sodium oxybate	377
sodium phenylbutyrate oral tablet	379
SOMATULINE DEPOT	
SUBCUTANEOUS SYRINGE 120	
MG/0.5 ML, 60 MG/0.2 ML, 90 MG/0.2	3
ML	225
SOMAVERT	314
sorafenib	389
SPRAVATO	136
SPRYCEL ORAL TABLET 100 MG,	
140 MG, 20 MG, 50 MG, 70 MG, 80	
MG	
STELARA 453,	
STIMUFEND	
STIVARGA	
STRENSIQ	
sunitinib malate	
SUNOSI	
SUPPRELIN LA	
SYMDEKO	
SYMLINPEN 120	
SYMLINPEN 60	
SYMPAZAN	
SYNAGIS	
SYNAREL	
SYNRIBO	
TABRECTA	
tadalafil (pulm. hypertension)	
tadalafil oral tablet 2.5 mg, 5 mg	
TAFINLAR ORAL CAPSULE	78

3	TAFINLAR ORAL TABLET FOR
•	SUSPENSION
)	TAKHZYRO SUBCUTANEOUS
_	SOLUTION
)	TAKHZYRO SUBCUTANEOUS
)	SYRINGE 150 MG/ML, 300 MG/2 ML
_	(150 MG/ML)
5	TALTZ AUTOINJECTOR
2	TALTZ SYRINGE SUBCUTANEOUS
	SYRINGE 80 MG/ML
2	TALVEY
3	TALZENNA
)	TASCENSO ODT152
3	TASIGNA ORAL CAPSULE 150 MG,
7	200 MG, 50 MG 274
)	tasimelteon
	TAVALISSE153
	TAZVERIK401
	TECENTRIQ29
5	TECVAYLI404
ł	<i>tencon</i> 175
)	TEPEZZA 408
5	ТЕРМЕТКО 407
	teriflunomide 409
	testosterone cypionate
5	testosterone enanthate
5	testosterone transdermal gel in metered-
3	<i>dose pump 12.5 mg/ 1.25 gram (1%),</i>
ł	20.25 mg/1.25 gram (1.62 %)
5	testosterone transdermal gel in packet 1 %
ł	(25 mg/2.5gram), 1 % (50 mg/5 gram)411
3	testosterone transdermal solution in
7	metered pump w/app
5	tetrabenazine
)	THALOMID
)	TIBSOVO
)	TIVDAK
)	torpenz oral tablet 10 mg, 2.5 mg, 5 mg,
7	7.5 mg
	TRACLEER ORAL TABLET FOR
5	SUSPENSION
5	TRAZIMERA
-	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT

TRELSTAR INTRAMUSCULAR
SUSPENSION FOR
RECONSTITUTION
TREMFYA
treprostinil sodium
tretinoin
tridacaine
trientine oral capsule 250 mg
TRIKAFTA ORAL GRANULES IN
PACKET, SEQUENTIAL
TRIKAFTA ORAL TABLETS,
SEQUENTIAL
TRIPTODUR
TRULICITY
TRUQAP
TRUXIMA
TUKYSA ORAL TABLET 150 MG, 50
MG
TURALIO
TYSABRI
TYVASO
UBRELVY
UDENYCA
UDENYCA AUTOINJECTOR
UDENYCA ONBODY
UPTRAVI INTRAVENOUS
UPTRAVI ORAL TABLET 1,000
MCG, 1,200 MCG, 1,400 MCG, 1,600
MCG, 200 MCG, 400 MCG, 600 MCG,
800 MCG
UPTRAVI ORAL TABLETS, DOSE
PACK
VALCHLOR
VANFLYTA
VEGZELMA47
VELCADE
VENCLEXTA ORAL TABLET 10
MG, 100 MG, 50 MG459
VENCLEXTA STARTING PACK 459
VERQUVO
VERZENIO
vigabatrin
vigadrone
vigpoder463
VIMIZIM

VITRAKVI ORAL CAPSULE 100	
MG, 25 MG	227
VITRAKVI ORAL SOLUTION	
VIZIMPRO	80
VONJO	
voriconazole oral suspension for	
reconstitution	465
VOSEVI	381
VOWST	142
VUMERITY	97
WELIREG	42
WINREVAIR	390
XADAGO	359
XALKORI ORAL CAPSULE	75
XALKORI ORAL PELLET 150 MG,	
20 MG, 50 MG	76
XELJANZ	424
XELJANZ XR	424
XERMELO	406
XGEVA	92
XIFAXAN ORAL TABLET 200 MG,	
550 MG	342
XOLAIR	.292
XOSPATA	160
XPOVIO ORAL TABLET 100	
MG/WEEK (50 MG X 2), 40	
MG/WEEK (40 MG X 1), 40MG	
TWICE WEEK (40 MG X 2), 60	
MG/WEEK (60 MG X 1), 60MG	
TWICE WEEK (120 MG/WEEK), 80	
MG/WEEK (40 MG X 2), 80MG	
TWICE WEEK (160 MG/WEEK)	369
XTANDI ORAL CAPSULE	.128
XTANDI ORAL TABLET 40 MG, 80	
MG	128
XYOSTED	413
yargesa	262
YERVOY	217
YONSA	7
ZARXIO	.148
zebutal	
ZEJULA ORAL CAPSULE	
	277
ZEJULA ORAL TABLET	
ZEJULA ORAL TABLET ZELBORAF ZIEXTENZO	277 458

ZIRABEV	49
ZOLADEX	170
ZTALMY	158
ZTLIDO	242
ZURZUVAE ORAL CAPSULE 20	
MG, 25 MG, 30 MG	467
ZYDELIG	
ZYKADIA	67
ZYMFENTRA	212
ZYNLONTA	247
ZYNYZ	